Non-adiabatic electron dynamics in time-dependent density-functional theory

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Time-dependent density-functional theory (TDDFT) treats dynamical exchange and correlation (xc) via a single-particle potential, $V_{\text{xc}}(r, t)$, defined as a nonlocal functional of the density $n(r', t')$. The popular adiabatic local-density approximation (ALDA) for $V_{\text{xc}}(r, t)$ uses only densities at the same space-time point $(r, t)$. To go beyond the ALDA, two local approximations have been proposed based on quantum hydrodynamics and elasticity theory: (a) using the current as basic variable (C-TDDFT) [G. Vignale, C. A. Ullrich, and S. Conti, Phys. Rev. Lett. 79, 4878 (1997)], (b) working in a co-moving Lagrangian reference frame (L-TDDFT) [I. V. Tokatly, Phys. Rev. B 71, 165105 (2005)]. This paper illustrates, compares, and analyzes both non-adiabatic theories for simple time-dependent model densities in the linear and nonlinear regime, for a broad range of time and frequency scales. C- and L-TDDFT are identical in certain limits, but in general exhibit qualitative and quantitative differences in their respective treatment of elastic and dissipative electron dynamics. In situations where the electronic density rapidly undergoes large deformations, it is found that non-adiabatic effects can become significant, causing the ALDA to break down.

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\section{I. INTRODUCTION}

Time-dependent density-functional theory (TDDFT) has gained considerable popularity as a practical approach to the dynamics of many-electron systems.\textsuperscript{2,4,5} The essential idea of TDDFT is to describe $N$ interacting electrons moving in an external potential $V(r, t)$ in terms of an auxiliary non-interacting system governed by the time-dependent Kohn-Sham (TDKS) equation,

$$\left[ \frac{1}{i \hbar} \frac{\partial}{\partial t} - \frac{\nabla^2}{2} + V(r, t) + V_{\text{H}}(r, t) + V_{\text{xc}}(r, t) \right] \varphi_{\alpha}(r, t) = 0$$

(1)

(we use atomic units $\hbar = e = m = 1$ throughout). Here and in the following, we consider systems whose ground state and dynamical response are everywhere nonmagnetic, and we may therefore ignore the spin degree of freedom. Eq. (1) thus describes the time evolution of doubly occupied Kohn-Sham orbitals $\varphi_{\alpha}(r, t)$, and the time-dependent density

$$n(r, t) = \sum_{\alpha=1}^{N/2} |\varphi_{\alpha}(r, t)|^2$$

(2)

is obtained in principle exactly. In Eq. (1), $V_{\text{H}}(r, t) = \int d^3r' n(r', t)/|r - r'|$ is the time-dependent Hartree potential, and $V_{\text{xc}}(r, t)$ is the exchange-correlation (xc) potential. In practice, suitable approximations for $V_{\text{xc}}(r, t)$ are required. We assume in the following that the system evolves from its ground state at $t = t_0$, although this assumption is not strictly necessary.

The exact $V_{\text{xc}}[n](r, t)$ has a functional dependence on $n(r, t)$ that is nonlocal in space and time, i.e., contains information about the previous history of the system, including its initial state. However, almost all present applications of TDDFT employ the adiabatic approximation, ignoring all functional dependence of $V_{\text{xc}}$ on prior time-dependent densities $n(r', t')$, $t' < t$. The simplest example is the adiabatic local-density approximation (ALDA):

$$V_{\text{ALDA}}^{\text{ALDA}}(r, t) = \left. \frac{d\epsilon_{\text{xc}}(\bar{n})}{dn} \right|_{\bar{n}=n(r, t)},$$

where $\epsilon_{\text{xc}}(\bar{n})$ is the xc energy density of a homogeneous electron gas of density $\bar{n}$. The neglect of retardation in ALDA implies frequency-independent and real xc kernels in linear response. This approach has been widely used for calculating molecular excitation energies.\textsuperscript{6}

One can make the general statement that the adiabatic approximation works well for excitations of the many-body system that have a direct counterpart in the Kohn-Sham system, such as atomic and molecular single-particle excitations.\textsuperscript{7} On the other hand, for more complicated processes such as double or charge-transfer excitations the ALDA often fails dramatically.\textsuperscript{8}

Several attempts to go beyond the ALDA can be found in the literature.\textsuperscript{9-11} Vignale and Kohn showed that a non-adiabatic local approximation for exchange and correlation requires the time-dependent current $j(r, t)$ as basic variable, rather than the density $n(r, t)$ (C-TDDFT). This formalism was later cast in a physically more transparent form using the language of hydrodynamics,\textsuperscript{12,13} where non-adiabatic xc effects appear as viscoelastic stresses in the electron liquid.

To date, C-TDDFT has been applied mainly in frequency-dependent linear response. The first application was to calculate linewidths of intersubband plasmons in semiconductor quantum wells.\textsuperscript{20,21} In the absence of disorder and phonon scattering, the ALDA gives infinitely sharp plasmon lines. C-TDDFT includes damping due to electronic many-body effects, in good agreement with experimental linewidths.\textsuperscript{22} Van Faassen et al.\textsuperscript{22} calculated static axial polarizabilities in molecular chains, which are greatly overestimated with ALDA, and achieved an excellent agreement...
with ab initio quantum chemical results. Other recent studies used C-TDDFT to calculate atomic and molecular excitation energies. Beyond linear response, a wealth of interesting electron dynamics can be explored using TDKS theory. The C-TDDFT formalism has recently been applied to describe linear and nonlinear charge-density oscillations in quantum wells in the time domain. It was shown that the retardation caused by the memory dependence of the xc potential has the striking consequence of introducing decoherence and energy relaxation.

An alternative non-adiabatic theory has recently been developed by one of the authors. The idea is to relate the local stress in the electron liquid, and thus the xc potential, to the dynamics of deformations of fluid elements in the quantum many-body system. This leads to a formally exact reformulation of TDDFT from the point of view of an observer in a co-moving Lagrangian reference frame (L-TDDFT). Casting the theory in terms of Lagrangian coordinates allows one to get around the well-known problem of “ultranonlocality” in TDDFT, and to derive, in a rigorous fashion, an exact time-dependent, non-adiabatic extension of the ground-state LDA.

In the L-TDDFT formalism the xc potential appears as a local functional of the dynamic deformation tensor. At present, two limiting forms of this local functional are available. A high-frequency, “elastic” form of the non-adiabatic xc potential was derived in Ref. 12. The elastic approximation correctly accounts for all complicated nonlinear deformation effects, but completely neglects possible xc contributions to dissipation. The second available limiting form of the non-adiabatic xc potential corresponds to the regime of small deformations. As we will show, in the limit of small deformations L-TDDFT formally reduces to C-TDDFT. In this limit the xc stress (and thus the xc potential) is proportional to the linearized strain tensor which can be considered as a local linear functional of the current. In fact, C-TDDFT of Ref. 13 can be viewed as a viscoelastic linear Hooke’s law in the nonlinear quantum continuum mechanics defined by the general formulation of L-TDDFT. In contrast to the purely elastic approximation constructed in Ref. 12, C-TDDFT contains all dissipation effects, but is formally restricted to infinitesimally small deformations. Of course, the formal asymptotic criteria and the practical regimes of applicability of any approximation can be quite different. Therefore a more detailed analysis based on explicit numerical examples is required to assess the validity of the two currently available non-adiabatic xc functionals, and to analyze their relation to ALDA.

Thus, the purpose of this paper is to illustrate, compare, and analyze C-TDDFT and the elastic approximation to L-TDDFT for simple, quasi-one-dimensional model systems, in order to show the differences and common features of both approaches. We explore the performance of the two xc potentials for two kinds of analytically given time-dependent model densities, representing charge-density oscillations in the form of collective sloshing and breathing modes of varying amplitudes and frequencies. This will allow us to simulate electron dynamics over a wide range of time and frequency scales, from the linear to the nonlinear regime. In our analysis, we will focus on a detailed comparison of the time-dependent xc potentials, as well as on the instantaneous and time-averaged power absorption associated with the charge-density oscillations. This will give us insight into the inner workings of C- and L-TDDFT in different dynamical regimes, and their relation to the ALDA. In particular, we will discuss and clarify the meaning of “non-adiabatic”, the cross-over from the linear to the nonlinear domain, and the competition and coexistence of elastic and dissipative xc effects.

It turns out that, for small-amplitude deformations, C- and L-TDDFT agree in the limit of short time scales or, equivalently, in the high-frequency regime. In general, the two theories show some differences in their treatment of elastic and dissipative effects in the inhomogeneous electron liquid. However, the size of these differences strongly depends on the type of collective mode, and the associated charge-density deformations. If the deformations are large and occur on short time (high frequency) scales, both non-adiabatic theories give a clear indication of a failure of the ALDA.

This paper is organized as follows. In section II we summarize the essential formal framework of C- and L-TDDFT. Section III shows how to construct simple analytic model densities in the Lagrangian and associated laboratory reference frame. Section IV presents detailed numerical results and discussion, with a separate treatment of the linear and nonlinear regime. We give our conclusions in Section V.

II. NON-ADIABATIC THEORIES IN TDDFT

A. C-TDDFT

1. Linear-response regime

Starting point of C-TDDFT is the linear current-density response \( j_1(r, \omega) \) to an external, frequency-dependent vector potential \( A_{ext1}(r, \omega) \).

\[
  j_{1,i}(r, \omega) = \int d^3r' \chi_{KS,i,j}(r, r', \omega) [A_{ext1,j}(r', \omega) + A_{H1,j}(r', \omega) + A_{xc1,j}(r', \omega)],
\]

where \( i,j \) denote Cartesian coordinates, and \( \chi_{KS,i,j} \) is the non-interacting, Kohn-Sham current-current response tensor. Here and in the following, we use the Einstein convention for the summation over repeated indices. The Hartree vector potential is given by

\[
  A_{H1,j}(r, \omega) = \frac{\nabla_j}{(i\omega)^2} \int d^3r' \frac{\nabla' \cdot j_1(r', \omega)}{|r - r'|}. \tag{5}
\]

The simplest approximation for the linearized xc vector potential \( A_{xc1} \) is the ALDA, which is defined as

\[
  A_{xc1}(r, \omega) = \frac{\nabla_j}{(i\omega)^2} \int d^3r' f_{xc}^{ALDA}(r, r') \nabla' \cdot j_1(r', \omega), \tag{6}
\]

where

\[
  f_{xc}^{ALDA}(r, r') = \frac{d^2 \varepsilon_{xc}(\tilde{n})}{dn^2} |_{\tilde{n} = n_0(r)} \delta(r - r'). \tag{7}
\]
is the frequency-independent ALDA xc kernel \([n_0(r)\) is the ground-state density]. In contrast with the xc scalar potential, the xc vector potential admits a frequency-dependent local approximation. The resulting expression can be written as follows\(^\text{12,13,14}\)

\[
A_{\text{xc},ij}(r, \omega) = A_{\text{xc},ij}^\text{ALDA}(r, \omega) - \frac{c}{i \omega n_0(r)} \nabla_k \sigma_{\text{xc},jk}(r, \omega).
\]

Here, \(c\) denotes the speed of light, and \(\sigma_{\text{xc},jk}\) is the xc viscoelastic stress tensor:

\[
\sigma_{\text{xc},jk} = \eta_{\text{xc}} \left( \nabla_j v_{1,k} + \nabla_k v_{1,j} - \frac{2}{3} \nabla \cdot v_1 \delta_{jk} \right) + \zeta_{\text{xc}} \nabla \cdot v_1 \delta_{jk},
\]

where \(v_1(r, \omega) = j_1(r, \omega)/n_0(r)\) is the velocity field associated with the current response, and \(\eta_{\text{xc}}\) and \(\zeta_{\text{xc}}\) are complex viscosity coefficients defined as

\[
\eta_{\text{xc}}(n, \omega) = -\frac{n^2}{i \omega} f_{\text{xc}}^T(n, \omega),
\]

\[
\zeta_{\text{xc}}(n, \omega) = -\frac{n^2}{i \omega} \left( f_{\text{xc}}^L(n, \omega) - \frac{4}{3} f_{\text{xc}}^T(n, \omega) - \frac{d^2 \epsilon_{\text{xc}}}{dn^2} \right).
\]

\(f_{\text{xc}}^L(n, \omega)\) and \(f_{\text{xc}}^T(n, \omega)\) are frequency-dependent xc kernels for the homogeneous electron gas, which can be found in various parametrizations in the literature\(^\text{6,29,30}\). In Eq. (9), \(\eta_{\text{xc}}\) and \(\zeta_{\text{xc}}\) are both evaluated at the local \(n_0(r)\).

2. Nonlinear regime

The generalization\(^\text{13,37}\) of C-TDDFT into the nonlinear regime and the time domain requires solving the following TDKS equation:

\[
\frac{1}{c} \frac{\partial \phi(r, t)}{\partial t} = \left[ \frac{1}{2} \left( \nabla \frac{1}{c} + \frac{1}{c} A(r, t) + \frac{1}{c} A_{\text{xc}}(r, t) \right)^2 + V(r, t) + V_H(r, t) \right] \phi(r, t).
\]

Notice that the Hartree term can be expressed as a scalar potential, and we are free to admit external scalar as well as vector potentials.

As explained in Ref. \(^\text{13}\) the form of the nonlinear xc vector potential is dictated by a number of general requirements, such as Newton’s third law (xc force density follows from a symmetric stress tensor), and the proper limit in the linear regime, which was discussed in the subsection above. A formally exact, general expression for \(A_{\text{xc}}\) resulting from these requirements will be presented within the Lagrangian framework in Section \(\text{13,18}\). However, a straightforward expression for a nonlinear, nonadiabatic xc vector potential, valid up to second order in the spatial derivatives, follows almost immediately:

\[
\frac{1}{c} \frac{\partial A_{\text{xc},i}(r, t)}{\partial t} = -\nabla_i V_{\text{xc}}^\text{ALDA}(r, t) + \frac{\nabla_j \sigma_{\text{xc},ij}(r, t)}{n(r, t)},
\]

where the viscoelastic stress tensor \(\sigma_{\text{xc}}\) now contains the time-dependent velocity field \(v(r, t) = j(r, t)/n(r, t)\):

\[
\sigma_{\text{xc},ij}(r, t) = \int_{-\infty}^{t} dt' \left\{ \eta_{\text{xc}}(r, t') \left[ \nabla_i v_j(r, t') + \nabla_j v_i(r, t') \right] - \frac{2}{3} \nabla \cdot v(r, t') \delta_{ij} \right\} + \zeta_{\text{xc}}(r, t') \nabla \cdot v(r, t') \delta_{ij}.
\]

The viscosity coefficients in Eq. (14) are the Fourier transforms of (10) and (11):

\[
\eta(r, t', \omega) = \int \frac{d\omega}{2\pi} \eta(\bar{n}, \omega) e^{-i\omega(t-t')} \bigg|_{\bar{n}=n(r, t)}
\]

and similar for \(\zeta\). The apparent ambiguity in Eq. (15) whether the density should be evaluated at \(t\) or \(t'\) is resolved by noting that the difference involves higher gradient corrections. We emphasize again that the simple form of Eq. (13) is justified by the basic assumption that the gradients of the density and velocity are small; the velocity itself, on the other hand, need not be small. These points will be elaborated in more detail in Section \(\text{13,18}\) where we will explain how the approximate expression (13) is obtained from the Lagrangian framework in the appropriate limit.

In the following, we consider model systems where all spatial dependence is along the \(x\) direction only. One can then transform the xc vector potential, Eq. (15), into a scalar one:

\[
V_{\text{xc}}(x, t) = V_{\text{xc}}^\text{ALDA}(x, t) + V_{\text{xc}}^\text{M}(x, t)\text{ (ALDA+M)},
\]

with the memory part given by

\[
V_{\text{xc}}^\text{M}(x, t) = -\int_{-\infty}^{x} \frac{dx'}{n(x', t)} \nabla x' \sigma_{\text{xc},xx}(x', t).
\]

Assuming that the system has been in the ground state (with zero velocity field) for \(t < 0\), the \(xx\) component of the xc stress tensor becomes

\[
\sigma_{\text{xc},xx}(x', t) = \int_{0}^{t} Y(n(x', t), t-t') \nabla x' v_{xx}(x', t') dt',
\]

where the memory kernel \(Y\) is given by

\[
Y(n, t-t') = \frac{4}{3} \left[ \eta(n, t-t') + \zeta(n, t-t') \right].
\]

With the help of the Kramers-Kronig relations for \(f_{\text{xc}}^L\) we can express the memory kernel as follows:

\[
Y(n, t-t') = \frac{4}{3} \mu_{\text{xc}} \frac{n^2}{\pi} \int \frac{d\omega}{\omega} \Im f_{\text{xc}}^L(\omega) \cos(\omega(t-t')),\]

with the xc shear modulus of the electron liquid\(^\text{39}\)

\[
\mu_{\text{xc}} = \frac{3n^2}{4} \left[ \Re f_{\text{xc}}^L(0) - \frac{d^2 \epsilon_{\text{xc}}}{dn^2} \right].
\]

The short-time behavior of \(Y(n, t-t')\) is of particular interest, since it governs the high-frequency dynamics. The limit
Y(n, 0) can be expressed analytically using the Kramers-Kronig relation
\[ \int_{-\infty}^{\infty} \frac{d\omega}{\pi} \frac{\Im f_{xc}^L(\omega)}{\omega} = \Re f_{xc}^L(0) - f_{xc}^L, \]
where the high-frequency limit \( f_{xc}^L \) is known via the third-moment sum rules. The result is
\[ Y(n, 0) = -\frac{20}{3} \epsilon_{xc} + \frac{26m}{5} \frac{d\epsilon_{xc}}{dn} - \frac{2\epsilon_{xc}^2}{dn^2}. \]
It is also straightforward to see that \( \frac{dV}{dt}(n, 0) = 0 \), i.e., the memory kernel starts with zero slope.

Figure 1 shows the memory kernel \( Y(n, t - t') \) for \( r_s = 1, 2, 3, 4, 5, \) using the GK (full lines) and QV (dashed lines) parametrizations. Inset: short-time limit \( Y(n, 0) \) [Eq. (22)] versus \( r_s \).

**B. L-TDDFT**

1. General formulation

The main problem in constructing nonadiabatic approximations for xc potentials is an inherent nonlocality of the time-dependent theory. Physically, this nonlocality is related to the convective motion of the electron fluid (the particles at a given point of space retain the memory of their previous positions). The key idea of L-TDDFT is to eliminate the above source of nonlocality by reformulating the theory in the Lagrangian frame, i.e., in a local reference frame moving with the fluid. Since the convective motion in the Lagrangian frame is absent, a spatially local description of xc effects becomes possible. This possibility represents the most important outcome of L-TDDFT: it allows one to derive an exact nonadiabatic extension of LDA into the dynamic regime.

The general formulation of L-TDDFT starts with the exact relation of the xc force to the xc stress tensor \( P_{xc,i,j} \). By definition the xc potential \( A_{xc} \) ensures that the physical density and current are reproduced by an auxiliary system of noninteracting KS particles. This means that \( A_{xc} \) should produce an effective xc Lorentz force that exactly compensates for a difference of local stress forces in the real interacting system and in the noninteracting KS system. Accordingly, the xc vector potential should satisfy the following equation:
\[ -\frac{\partial A_{xc,i}}{\partial t} + v_j(\nabla_i A_{xc,j} - \nabla_j A_{xc,i}) = \frac{e}{n} \nabla_j P_{xc,i,j}, \quad (23) \]
where \( P_{xc,i,j} = P_{ij} - T_{ij}^{KS} \) is the difference of the full stress tensor for the interacting system, \( P_{ij} \), and the kinetic stress tensor for KS system, \( T_{ij}^{KS} \). Equation (23) serves as a basic definition of \( A_{xc} \), which automatically accounts for the zero force and zero torque conditions.

We note that the xc stress tensor \( P_{xc,i,j} \) is to be distinguished from the earlier introduced xc stress tensor \( \sigma_{xc,i,j} \). The connection between the two will be explained in detail in subsection B.2. The main difference lies in the fact that \( P_{xc,i,j} \), formally exactly and to all orders in the inhomogeneity, accounts for all dynamical xc effects, whereas the ALDA has been separated out in the definition of \( \sigma_{xc,i,j} \). Furthermore, \( \sigma_{xc,i,j} \) is valid only for small deformations of the electron liquid (in a sense to be defined below).

All the rest of L-TDDFT can be viewed as a calculation of the dynamic xc stress tensor \( P_{xc,i,j} \), which enters the definition of \( A_{xc} \), Eq. (23), by reformulating the problem in the co-moving Lagrangian frame. The transformation to the Lagrangian frame corresponds to a nonlinear transformation of coordinates, \( r = r(\xi, t) \), where \( r(\xi, t) \) is the trajectory of an infinitesimal fluid element that evolves from the point \( \xi \). Formally the function \( r(\xi, t) \) is defined by the following initial value problem:
\[ \frac{\partial r(\xi, t)}{\partial t} = v(r(\xi, t), t), \quad r(\xi, 0) = \xi. \]

The initial positions, \( \xi \), of the fluid elements play the role of spatial coordinates in the co-moving frame. The above transformation from the old coordinates \( r \) to the new coordinates \( \xi \) induces a change of metric. The metric tensor in the Lagrangian \( \xi \)-space coincides with the Green’s deformation tensor:
\[ g_{ij}(\xi, t) = \frac{\partial r_k(\xi, t)}{\partial \xi_i} \frac{\partial r_k(\xi, t)}{\partial \xi_j}, \quad (25) \]
which is a common characteristic of deformations in the Lagrangian formulation of continuum mechanics. A complete reformulation of TDDFT in the co-moving frame shows that a local description of xc effects is possible if one uses $g_{ij}(\xi, t)$ as a basic variable: the stress tensor in the Lagrangian frame can be consistently considered as a spatially local functional of the Green’s deformation tensor.

For most practical applications we need the xc potential in the laboratory frame, Eq. (23). Transforming the stress tensor functional from the co-moving to the laboratory frame, we find the required tensor $P_{xc, ij}$ entering Eq. (23). The locality in the Lagrangian frame translates to a local dependence of $P_{xc, ij}$ on the Cauchy’s deformation tensor,

$$g_{ij}(\mathbf{r}, t) = \frac{\partial \xi_k(\mathbf{r}, t) \partial \xi_i(\mathbf{r}, t)}{\partial r_j},$$

where the function $\xi(\mathbf{r}, t)$ is obtained by inverting the trajectory equation, $\mathbf{r} = \mathbf{r}(\xi, t)$. The Cauchy’s tensor $g_{ij}(\mathbf{r}, t)$ is commonly used to describe deformations in the Eulerian formulation of continuum mechanics. From a physical point of view, the functional dependence of the stress tensor on $g_{ij}$ thus emerges quite naturally.

Substitution of $P_{xc, ij}[g_{ij}]$ into Eq. (23) yields the xc vector potential $A_{xc}$ as a functional of the Cauchy’s deformation tensor. If gradients of the deformation tensor are small, the functional $P_{xc, ij}[g_{ij}]$ is local in space, but in general it can be nonlocal in time. The time-nonlocality of the stress tensor functional is related to the memory loss due to dissipation. In the absence of dissipation (e.g. in the exchange-only approximation) the stress tensor becomes a simple function of $g_{ij}$, which corresponds to an infinitely long memory.

Equations (24) and (26) determine the Cauchy’s deformation tensor as a functional of the velocity $\mathbf{v}(\mathbf{r}, t)$. An alternative, and practically more convenient way to compute $g_{ij}$ for a given velocity is to solve the equation of motion that governs the dynamics of $g_{ij}(\mathbf{r}, t)$ directly in the laboratory frame:

$$\frac{\partial g_{ij}}{\partial t} + \mathbf{v} \cdot \frac{\partial g_{ij}}{\partial \mathbf{r}} = -\frac{\partial g_{ij}}{\partial \mathbf{r}} \frac{\partial \mathbf{v}}{\partial \mathbf{r}} - \frac{\partial \mathbf{v}}{\partial \mathbf{r}} \delta g_{kik}, \quad g_{ij}(\mathbf{r}, 0) = \delta_{ij}. \tag{27}$$

An important property of the deformation tensor is that it allows us to relate the time-dependent density $n(\mathbf{r}, t)$ to the initial density distribution, $n_0(\mathbf{r})$: $n(\mathbf{r}, t) = \sqrt{\tilde{g}(\mathbf{r}, t)n_0(\xi(\mathbf{r}, t))}, \tag{28}$

where $\tilde{g}(\mathbf{r}, t)$ is the determinant of $g_{ij}(\mathbf{r}, t)$.

Equation (27) or, equivalently, Eqs. (24) and (26), show that in general the deformation tensor is a strongly nonlocal (both in space and in time) functional of the velocity. Therefore, in spite of the fact that the xc stress tensor and consequently the xc vector potential are local functionals of $g_{ij}$, they are nonlocal in terms of velocity or any other variable. This makes the choice of $g_{ij}(\mathbf{r}, t)$ as a basic variable much more preferable.

To obtain an explicit construction of the local functional $P_{xc, ij}[g_{ij}]$, a solution of a homogeneous time-dependent many-body problem in the Lagrangian frame is required (see Sec. V in Ref. [18]). A complete solution of this problem seems to be impossible, at least at the current level of knowledge. However, there are two practically important, exactly solvable special cases, which are described below.

2. Small deformation approximation: Recovery of C-TDDFT

The abovementioned many-body problem in the homogeneously deformed Lagrangian $\xi$-space can be solved perturbatively if the deformation tensor $g_{ij}$ only slightly deviates from the unit tensor $\delta_{ij}$:

$$\tilde{g}_{ij}(\mathbf{r}, t) = \delta_{ij} + \bar{g}_{ij}(\mathbf{r}, t). \tag{29}$$

Introducing the displacement vector, $\mathbf{u}(\mathbf{r}, t) = \mathbf{r} - \mathbf{r}(\xi, t)$, and using Eq. (26), we find that small $\delta \tilde{g}_{ij}$ corresponds to small gradients of the displacement:

$$\delta \tilde{g}_{ij}(\mathbf{r}, t) = -\left(\frac{\partial \mathbf{u}_i}{\partial \mathbf{r}_j} + \frac{\partial \mathbf{u}_j}{\partial \mathbf{r}_i}\right). \tag{30}$$

Clearly, small gradients of $\mathbf{u}(\mathbf{r}, t)$ imply that the velocity gradients are also small, since to lowest order in $\nabla \mathbf{u}$, Eq. (24) reduces to the relation $\partial_t \mathbf{u}(\mathbf{r}, t) = \mathbf{v}(\mathbf{r}, t)$. Obviously the smallness of deformations does not mean that the displacement or the velocity themselves are small (i.e., the system can be far beyond the linear response regime). A well known example is the rigid motion of a many-body system in a harmonic potential, where $g_{ij} = \delta_{ij}$ but the displacement can be arbitrarily large.

The stress tensor functional for small displacement vectors was derived in Ref. [18]. Extension of this derivation to the general regime of small deformations, i.e., to the regime of small displacement gradients, is straightforward. The resulting xc stress tensor takes the following form:

$$P_{xc, ij}(\mathbf{r}, t) = P_{xc}^{\text{ALDA}}(n(\mathbf{r}, t))\delta_{ij} + \delta P_{ij}(\mathbf{r}, t), \tag{31}$$

where $P_{xc}^{\text{ALDA}}(n)$ is the xc pressure of a homogeneous electron gas, and $\delta P_{ij}$ is a nonadiabatic correction, which is linear in $\delta \tilde{g}_{ij}$:

$$\delta P_{ij}(\mathbf{r}, t) = \int_0^t dt' \left\{ \frac{\delta_{ij}}{2} \bar{K}_{xc}(n(\mathbf{r}, t), t - t') \delta \bar{g}_{kk}(\mathbf{r}, t') + \mu_{xc}(n(\mathbf{r}, t), t - t') \left[ \delta \bar{g}_{ij}(\mathbf{r}, t') - \frac{\delta_{ij}}{3} \delta \bar{g}_{kk}(\mathbf{r}, t') \right] \right\}. \tag{32}$$

The kernels $\mu_{xc}(n, t - t')$ and $\bar{K}_{xc}(n, t - t')$ in Eq. (32) have the meaning of nonadiabatic shear and bulk moduli, respectively [the adiabatic part of the bulk modulus is included in the ALDA pressure term in Eq. (31)]. The corresponding Fourier transforms of the elastic moduli, $\mu_{xc}(\mathbf{q}, \omega)$ and $\bar{K}_{xc}(\mathbf{q}, \omega)$, are related to the complex viscosity coefficients of Eqs. (10) and (11), $\eta_{xc}(\mathbf{q}, \omega)$ and $\zeta_{xc}(\mathbf{q}, \omega)$, as follows:

$$\mu_{xc}(\omega) = -i \omega \eta_{xc}(\omega), \quad \bar{K}_{xc}(\omega) = -i \omega \zeta_{xc}(\omega). \tag{33}$$

Using Eq. (33) and the relation $\partial_t \mathbf{u} = \mathbf{v}$, we find that the nonadiabatic stress tensor $\delta P_{ij}$, Eq. (32), is identical, up to a
deformation tensor takes a diagonal form with the nonlinear elastic approximation for a one-dimensional body problem, which has been formulated in Ref. 18, admits a simple complete solution in the regime of fast dynamics when the dissipation is irrelevant. As a result, the high-frequency limit of the nonadiabatic stress tensor of Eq. (32) becomes completely local and purely elastic:

\[ \delta P_{ij}^\infty(r, t) = \frac{\delta_{ij}}{2} \tilde{K}_{\text{xc}}(n(r, t)) \delta g_{kk}(r, t) + \mu_{\text{xc}}(n(r, t)) \left[ \delta g_{ij}(r, t) - \frac{\delta_{ij}}{3} \delta g_{kk}(r, t) \right], \]

where \( \tilde{K}_{\text{xc}}(n) \) and \( \mu_{\text{xc}}(n) \) are the high-frequency limits of the bulk and shear moduli, respectively.

The structure of the small deformation approximation, Eqs. (31) and (32), clearly demonstrates that in this regime the nonadiabatic contribution appears as a small, linear in \( \delta g_{ij} \) correction to the adiabatic dynamics. If the process is strongly nonadiabatic, the deformations cannot be considered small. In fact, the deviation of the deformation tensor from \( \delta_{ij} \) can serve as a general measure of nonadiabaticity.

3. Nonlinear elastic approximation to L-TDDFT

It is very difficult to account both for the full nonlinear dependence on \( \delta g_{ij} \), and for the dissipation. C-TDDFT includes all xc dissipation effects on a level linear in \( \delta g_{ij} \). On the contrary, if we neglect the dissipation effects, a closed nonlinear local approximation for the stress tensor can be rigorously derived. The reason is that the homogeneous many-body problem, which has been formulated in Ref. 18, admits a simple complete solution in the regime of fast dynamics when the dissipation is irrelevant. In this case the xc stress tensor \( P_{\text{xc}, ij}(r, t) \) becomes a function of the time-dependent density \( n(r, t) \) and the Cauchy’s deformation tensor \( g_{ij}(r, t) \):

\[ P_{\text{xc}, ij}(r, t) = \frac{2}{3} \bar{g}_{ij} \sqrt{\bar{g}} \bar{E}_{\text{xc}}(n/\sqrt{\bar{g}}) + L_{ij}(\bar{g}_{kl}) E_{\text{pot}}(n/\sqrt{\bar{g}}), \]

where \( \bar{E}_{\text{xc}}(n) \) and \( E_{\text{pot}}(n) \) are the xc kinetic energy and the potential energy per unit volume of the homogeneous electron gas. The function \( L_{ij}(\bar{g}_{kl}) \) in Eq. (35) is explicitly defined in Appendix A of Ref. 18. In the limit of small deformations the nonlinear elastic approximation of Eq. (35) can be expanded around \( \bar{g}_{ij} = \delta_{ij} \) and reduces to the linearized form defined by Eqs. (31) and (32). In other words one recovers the high-frequency limit of C-TDDFT.

We conclude this section with the explicit formulation of the nonlinear elastic approximation for a one-dimensional motion. If all spatial variations are along the \( x \)-axis only, the deformation tensor takes a diagonal form with \( \bar{g}_{zz} = \bar{g}_{yy} = 1 \), and \( \bar{g}_{xx} = \bar{g}(x, t) \). The xc effects can then be described by an xc scalar potential that is related to the xc stress tensor as follows:

\[ V_{\text{xc}}^E(x, t) = \int_{-\infty}^{x} \frac{dx'}{n(x', t)} \frac{\partial}{\partial x'} P_{\text{xc}, xx}(n(x', t), \bar{g}(x', t)). \]

Equation (35) for the \( xx \)-component of the xc stress tensor reduces to the form

\[ P_{\text{xc}, xx}(n, \bar{g}) = \frac{2}{3} \bar{g}^{3/2} P_{\text{xc}}^{\text{kin}} \left( \frac{n}{\sqrt{\bar{g}}} \right) + L(\bar{g}) E_{\text{pot}} \left( \frac{n}{\sqrt{\bar{g}}} \right), \]

where the factor \( L(\bar{g}) \) is given by

\[ L(\bar{g}) = \frac{\bar{g}}{\bar{g} - 1} \left[ 1 - \arctan \frac{\sqrt{\bar{g} - 1}}{\sqrt{\bar{g} + 1}} \right]. \]

Finally, Eq. (27), which relates the deformation \( \bar{g}(x, t) \) to the velocity \( v(x, t) \), simplifies as follows:

\[ \frac{\partial \bar{g}}{\partial t} = -v \frac{\partial \bar{g}}{\partial x} - 2 \frac{\partial v}{\partial x} \bar{g}, \quad \bar{g}(x, 0) = 1. \]

It is worth mentioning that \( L(\bar{g} \rightarrow 1) = 1/3 \) in the limit of zero deformation. Eq. (37) then approaches the standard virial expression for the xc pressure, and \( V_{\text{xc}}^E(x, t) \) reduces to the ALDA xc potential. We define the post-ALDA contribution of the nonlinear elastic L-TDDFT xc potential as

\[ \bar{V}_{\text{xc}}^E(x, t) = V_{\text{xc}}^E(x, t) - V_{\text{ALDA}}^{\text{xc}}(x, t). \]

To shorten the notation for the rest of this paper: from now on, whenever we refer to L-TDDFT, we actually mean the nonlinear elastic approximation to L-TDDFT, defined in this section.

III. CONSTRUCTION OF ONE-DIMENSIONAL ANALYTICAL EXAMPLES

In one dimension, the trajectory of a fluid element with Lagrangian coordinate \( \xi \) is determined by the following equation:

\[ \frac{\partial x(\xi, t)}{\partial t} = v \left( x(\xi, t), t \right), \quad x(\xi, 0) = \xi. \]

\( x \) is the position, at time \( t \), of a fluid element that started out at \( \xi \), and \( v \) is its velocity. In general, this is a complicated nonlinear differential equation, with formal solution

\[ x(\xi, t) = \xi + \int_{0}^{t} v(\xi, t') dt'. \]

In other words: if we know the velocity, at time \( t \), of a fluid element that started out at \( \xi \), we can determine its trajectory by
direct integration. From this, we can then determine the time-dependent density in the laboratory frame: we first invert (42) to obtain \( \xi(x, t) \), then compute the deformation as
\[
\bar{g}(x, t) = \left( \frac{\partial \xi}{\partial x} \right)^2,
\]
and finally obtain
\[
n(x, t) = \sqrt{\bar{g}(x, t)} n_0(\xi(x, t)).
\]
In practice, of course, this procedure is not very helpful, since the functional form of \( v(\xi, t) \) is unknown. However, we can use it to construct simple analytic examples, as follows.

A. Sloshing mode

We assume that the system is confined within hard walls, \(-L/2 \leq (x, \xi) \leq L/2\), with initial density
\[
n_0(\xi) = \frac{2N}{L} \cos^2 \left( \frac{\pi \xi}{L} \right),
\]
where \( N \) is the number of electrons per unit area (sheet density) in the \( y-z \) plane. We assume a simple quadratic form of the velocity field:
\[
v(\xi, t) = A \omega \left( \frac{L}{4} - \frac{\xi^2}{L} \right) \cos \omega t.
\]
Equation (42) is then easily integrated:
\[
x(\xi, t) = \xi + A \left( \frac{L}{4} - \frac{\xi^2}{L} \right) \sin \omega t.
\]
The next step is to invert Eq. (47) to determine the trajectories of the fluid elements, which requires solving a quadratic equation, with the result
\[
\xi(x, t) = \frac{L}{2A \sin \omega t} \left( 1 - \sqrt{1 + A^2 \sin^2 \omega t - \frac{4Ax}{L} \sin \omega t} \right)
\]
which properly reduces to \( \xi = x \) for \( A \to 0 \). The range of allowed amplitudes is \( |A| \leq 1 \), which is dictated by the constraint that no fluid element can cross the hard-wall boundaries at \( \pm L/2 \).

We can now calculate the deformation using Eq. (43):
\[
\bar{g}(x, t) = \left( 1 + A^2 \sin^2 \omega t - \frac{4Ax}{L} \sin \omega t \right)^{-1}.
\]
The time-dependent density of the sloshing mode in the laboratory frame, \( n(x, t) \), then follows from Eq. (44), using (48) and (49).
B. Breathing mode

To simulate a breathing mode, we assume a linear velocity distribution of the fluid elements:

\[ v(\xi, t) = A \omega \xi \cos \omega t \, . \]  

(50)

According to Eq. (42), this gives the following trajectory:

\[ x(\xi, t) = \xi (1 + A \sin \omega t) \, , \quad |A| < 1 \, . \]  

(51)

This is easily inverted:

\[ \xi(x, t) = \frac{x}{1 + A \sin \omega t} \, , \]  

(52)

and the resulting deformation is

\[ \tilde{g}(x, t) = \frac{1}{(1 + A \sin \omega t)^2} \, . \]  

(53)

We choose the same initial density distribution \( n_0(\xi) \), Eq. (45), as for the sloshing mode, and the resulting time-dependent density of the breathing mode is

\[ n(x, t) = \frac{2N}{L} \cos^2 \left( \frac{\pi x}{L} \right) \left( 1 + A \sin \omega t \right)^{-1} \, , \]  

(54)

where \( |x| < (L/2)(1 + A \sin \omega t) \).

Figs. 2 and 3 show snapshots of \( n(x, t) \), \( v(x, t) \), and \( \tilde{g}(x, t) \) for the sloshing and the breathing mode, taken at time \( t = 0, T/4, T/2, 3T/4 \), where \( T = 2\pi/\omega \). The amplitude is \( A = 0.5 \) in both cases, length is measured in units of \( L \), and density and velocity are plotted in units of \( N/L \) and \( L \omega \), respectively. The deformation \( \tilde{g}(x, t) \) is maximal at the turning points of the oscillations (at \( t = T/4 \) and \( 3T/4 \)). We find that the breathing mode features large deformations everywhere, i.e., \( \tilde{g} \) deviates strongly from one. The sloshing mode, on the other hand, is strongly deformed only towards the edges where the density is small. We will see below how this affects the non-adiabaticity of the xc potential of the two modes.

IV. RESULTS AND DISCUSSION

In the following numerical examples, we choose a system where, in atomic units, we have sheet density \( N = 1 \) a.u. and quantum well width \( L = 10 \) a.u. We shall measure frequencies in units of the average plasmon frequency of the system, given by

\[ \tilde{\omega}_p = \frac{1}{L} \int dx \omega_p(n(x)) \, . \]  

(55)

For our initial density distribution \( n_0(x) \), Eq. (45), we find \( \tilde{\omega}_p = \sqrt{32N/\pi L} \), which for the above values of \( N \) and \( L \) comes out as \( \tilde{\omega}_p = 1.009 \) a.u.

A. Linear regime

1. Time-dependent xc potentials

We first consider the linear regime of small density fluctuations. For the time-dependent densities associated with the breathing mode and the sloshing mode, we calculate and compare three different time-dependent xc potentials: \( V_{xc}^M(x, t) \) from C-TDDFT [Eq. (16)], \( V_{xc}^{\nu}(x, t) \) from L-TDDFT [Eq. (40)], and the ALDA potential fluctuations \( V_{xc}^{ALDA}(x, t) = V_{xc}^{ALDA}(n(x, t)) - V_{xc}^{ALDA}(n_0(x)) \). Figures 4 show four snapshots of \( V_{xc}^M(x, t) \) and \( V_{xc}^{ALDA}(x, t) \) during one cycle of the sloshing/breathing modes, at \( t = 0, T \) and \( T/2 \) (density passes through equilibrium, with maximal velocity) and \( t = T/4 \) and \( 3T/4 \) (density at turn-
The results in Figs. 4–7 reveal the following features:

(i) The high-frequency limit of $V_{xc}^{M}(x, t)$ [i.e., $\tilde{V}_{E}^{ALDA}(x, t)$] is phase-shifted by $\pi$ with respect to $\tilde{V}_{E}^{ALDA}(x, t)$ in all cases considered. This is to be expected for a purely elastic potential: it reaches its maximum at the instant of largest displacement from equilibrium.

(ii) At the low-frequency limit $\omega_L$, the behavior of $V_{xc}^{M}$ depends on the parametrization used to calculate the memory kernel: in GK, $V_{xc}^{M}$ vanishes, whereas in QV, $V_{xc}^{M}$ again becomes purely elastic. This reflects the different long-time behaviors of the memory kernel: in GK, it decreases exponentially, whereas in QV it approaches a finite constant (see Fig. 1).

(iii) At intermediate frequencies, the phase shift between $\tilde{V}_{E}^{ALDA}$ and $V_{xc}^{M}$ varies between $\pi$ and $\pi/2$, where $\pi/2$ indicates a purely dissipative potential (see below).

(iv) The average strength of the forces associated with $V_{xc}^{M}$ grows with frequency, and becomes comparable to the ALDA fluctuating forces in the high-frequency limit (notice that $\tilde{V}_{E}^{ALDA}$ is scaled by 0.1 in Figs. 4 and 5). This clearly shows that non-adiabatic effects can become non-negligible in practice. We will say more about this below when we discuss the nonlinear regime.
2. Power and dissipation

For a more quantitative analysis, it is useful to consider the power associated with the charge-density oscillations of the two types of modes. We define the power in the usual way as force density times velocity:

\[
P(t) = \int dx \, v(x,t) \, n(x,t) \, \frac{\partial}{\partial x} \, V_{xc}^M(x,t) . \tag{56}
\]

Figure 8 shows \(P(t)\), scaled by \(\omega A^2\), during one cycle of the sloshing and breathing modes (calculated using the GK parametrization). In the low-frequency limit, where the currents are vanishing, the power tends to zero, but as the frequency increases, a periodic input/output of power is observed to take place during a cycle. It can be clearly seen that, on average, \(P\) is more negative than positive for intermediate frequencies, which indicates net power dissipation. In the high-frequency, elastic limit, \(P(t)\) has sizable amplitudes, but averages to zero during a cycle.

These findings are summarized in Figures 9 and 10. We plot \(-\overline{P}\), the absolute value of the time-average of \(P(t)\), scaled by \(\omega A^2\), which represents the net power absorption per cycle, for the sloshing and the breathing modes. Notice that \(-\overline{P}\) in all cases has a pronounced enhancement for frequencies of the order of \(\bar{\omega}_p\). We define the crossover frequencies \(\omega_C\) as those frequencies where the maxima of power absorption occur.

While the two modes behave qualitatively very similarly as far as their power absorption is concerned, one observes in Figs. 8 and 10 that the absorbed power of the breathing mode is about an order of magnitude higher than for the sloshing mode, for the same value of the amplitude, \(A = 0.005\). This hardly comes as a surprise: our sloshing mode can be viewed as a cousin of Kohn’s mode and the Harmonic Potential Theorem tells us that charge-density oscillations in parabolic quantum wells are undamped. The breathing mode, on the other hand, bears no resemblance at all to Kohn’s mode. The sloshing mode has a smooth hydrodynamic flow with relatively little internal compression except at the turning points. By contrast, the defining feature of the breathing mode is the periodic compression and rarefaction of its density profile, i.e., a very “un-hydrodynamic” behavior.

It turns out that there is a direct relation between the average power absorption and the imaginary part of the xc kernel, \(\Im f_{xc}^M(\omega)\). This is clear from Eq. (19), which expresses the memory kernel \(Y(n,t-t')\) via Fourier transform of \(\Im f_{xc}^M(\omega)/\omega\). The full lines in Figs. 8 and 10 plot \(-\gamma \Im f_{xc}^M(\omega)\), where \(\gamma\) is a constant scaling factor. For a best fit, \(\gamma_{sl} = 0.00525\) and \(r_{sl} = 1.41\) for the sloshing mode, and \(\gamma_{br} = 0.071\) and \(r_{br} = 1.18\) for the breathing mode (same for GK and QV). Notice that the equilibrium density \(n_0(x)\)
has a value of \( r_s = 1.06 \) in the center. It thus emerges that the dissipation is dominated by different regions of the density distribution for the two modes: around \( x = \pm 0.27L \) for the sloshing mode, i.e., halfway between center and walls, and \( x = \pm 0.18L \) for the breathing mode, i.e., much closer to the center.

The phase lag of the modes as a function of frequency was already discussed in the context of Figs. 4-7. However, in Figs. 9 and 10 one can see the difference between GK and QV most dramatically. We consider the phase lag between the instantaneous power absorption, \( \mathcal{P}(t) \), and the ALDA potential fluctuations (which are in phase with the charge-density oscillations). Fig. 9 shows that in GK one has a transition from a purely dissipative behavior in the low-frequency limit, with phase difference \( \pi/2 \), via a crossover region of mixed dissipative/elastic behavior, to the high-frequency, purely elastic regime with phase lag \( \pi \). The maximal power absorption occurs for a phase lag of \( 3\pi/4 \). On the other hand, in the QV parametrization the low-frequency regime is also purely elastic, i.e. both the low- and high-frequency limits have phase lag \( \pi \), and dissipative contributions come in only at intermediate frequencies. The resulting power loss as a function of frequency is thus a bit higher and more narrowly peaked about \( \omega_C \) in QV than in GK.

As mentioned earlier, the qualitative differences between GK and QV have their origin in the different long-time behavior of the memory kernels, see Fig. 1 and the discussion in Section II A 2. The overall result is that there is a broad range of frequencies, between about 0.1 to 10 times the characteristic average plasma frequency \( \bar{\omega}_p \), where the system exhibits a mixed elastic/dissipative behavior, which can lead to substantial dissipation.

**B. Nonlinear regime**

1. C- versus L-TDDFT in the high-frequency limit

In the following, we will extend our numerical studies of C- and L-TDDFT into the nonlinear regime. We begin by directing our attention to the high-frequency, purely elastic region, since our approximate version of L-TDDFT becomes exact in that limit and can thus be used as a benchmark to assess the accuracy of nonlinear C-TDDFT.

Fig. 11 shows the scaled power \( \mathcal{P}(t)/\omega A^2 \) [Eq. (56)] for \( \omega = 10^3\omega_p \) during one cycle of the sloshing and breathing modes, comparing small amplitudes (\( A = 0.005 \)) and large amplitudes (\( A = 0.5 \) and 0.75). In the small-amplitude regime, \( \mathcal{P}(t)/\omega A^2 \) has a sinusoidal shape for both modes, independent of the amplitude \( A \) of the oscillations (see also Fig. 8), but for large amplitudes, nonlinear deviations occur. In the sloshing mode, \( \mathcal{P}(t) \) tends to a more sawtooth-like shape. In the breathing mode, \( \mathcal{P}(t) \) becomes suppressed when the charge density spreads out in the first half of the cycle, and dramatically enhanced when the density gets squeezed in the second half of the cycle. As we will see below, these strong deformations give rise to sizable non-adiabatic effects.

For small amplitude, \( A = 0.005 \), C- and L-TDDFT are
identical, but for larger amplitudes, \( A = 0.5 \) and 0.75, some differences develop. However, the magnitude of these deviations strongly depends on the type of mode. For the sloshing mode, we find that C-TDDFT remains quite close to L-TDDFT even for large-amplitude oscillations, whereas in the breathing mode, C-TDDFT deviates from L-TDDFT by about a factor of two for large values of \( A \).

A more quantitative measure of the deviation of C-TDDFT from the exact high-frequency L-TDDFT result is shown in Fig. 12. We plot the difference in percent of the time average of the absolute value of the scaled power, \( (t_2 - t_1)^{-1} \int_{t_1}^{t_2} dt |P(t)|/\omega A^2 \), over one cycle for the sloshing mode, and over the first and second half-cycle for the breathing mode, as a function of the amplitude \( A \). As expected, the difference is seen to be increasing as the amplitude grows, but the two modes exhibit a very different behavior.

Due to its close kinship to Kohn’s mode, it comes as no surprise that the deviation is much smaller for the sloshing mode, namely at most around 2% even for very large amplitudes. The C-TDDFT error of the breathing mode is larger by at least an order of magnitude. However, the overall error of C-TDDFT in the high-frequency limit compared to L-TDDFT remains surprisingly small, as long as the amplitude is not too large: For \( A = 0.2 \), we get a deviation of about 0.2% for the sloshing mode, and about 5% for the breathing mode. For the largest amplitude considered (\( A = 0.9 \)), we obtain a deviation of 2.5% for the sloshing mode, and 20% and 100% for first and second half-cycles of the breathing mode. In the latter case, the deformations are so strong as to invalidate the basic assumptions used to derive the simple form of nonlinear C-TDDFT.

2. Non-adiabatic corrections to the ALDA

The preceding high-frequency analysis shows that the C-TDDFT non-adiabatic xc potentials remain close (to within a few percent) to the exact L-TDDFT results, except for modes with extremely strong deformations such as sloshing modes with \( A \gtrsim 0.9 \) or breathing modes with \( A \gtrsim 0.2 \). As long as the deformations remain within these approximate limits, it is reasonable to expect C-TDDFT to be accurate for finite frequencies as well.

To get an impression of the magnitude of the non-adiabatic corrections to the ALDA, we plot in Figs. 13 and 14 the full adiabatic and non-adiabatic xc potentials, \( V_{xc}^{ALDA}(x, t) \) and \( V_{xc}^{ALDA}(x, t) + V_{xc}^{M}(x, t) \), where \( V_{xc}^{M} \) is calculated within C-TDDFT using the QV parametrization. Fig. 14 shows results for a large-amplitude sloshing mode with \( A = 0.75 \), and Fig. 14 for a breathing mode with \( A = 0.5 \). The figures show four snapshots taken during one cycle of the mode, similar to what was presented in Figs. 3 and 4, except that now we are plotting the total potential. The ALDA xc potential is of course independent of the mode frequency, and we compare it with the ALDA+M potential at three different frequencies: \( \omega = 0.1 \tilde{\omega}_p \) (low-frequency regime), \( \omega = \tilde{\omega}_p \) (around cross-over), and \( \omega = 10 \tilde{\omega}_p \) (approaching the high-frequency regime).

The results in Figs. 13 confirm again that the non-adiabatic effects for the sloshing mode are relatively modest, even for

![Diagram](image-url)
large-amplitude deformations. The post-ALDA corrections become more important for the high-frequency oscillations ($\omega = 10\tilde{\omega}_p$), where one finds deviations from ALDA of the order of 10% at the turning points of the oscillation at $T/4$ and $3T/4$. For the lower-frequency modes, the non-adiabatic corrections to the ALDA stay mostly within about 1%.

On the other hand, the breathing mode exhibits much more dramatic non-adiabatic effects, see Fig. 14. Again we find that the post-ALDA corrections are moderate for the lower frequencies considered. However, for the high-frequency case ($\omega = 10\tilde{\omega}_p$), we find that at the instances of maximum deformation ($T/4$ and $3T/4$) the memory effects cause a correction to the ALDA of up to a factor of 2, which is indeed substantial.

Lastly, to illustrate a case of extreme non-adiabaticity let us consider the high-frequency limit of strongly nonlinear dynamics. Fig. 15 shows a comparison of the ALDA and L-TDDFT xc potentials for both modes with amplitude $A = 0.75$. For the instants $0$, $T/2$, and $T$, ALDA and L-TDDFT coincide, but for $T/4$ and $3T/4$, large deviations occur. The most dramatic non-adiabatic effect is observed for the breathing mode at $3T/4$, where the elastic contribution of L-TDDFT is so large that the resulting xc potential is of the same magnitude as the ALDA, but with opposite sign. This clearly shows that in situations in which the electron density is rapidly and strongly deformed the ALDA
becomes drastically wrong, with non-adiabatic corrections of the same order of magnitude as the potential itself.

V. CONCLUSION

In this paper, we have carried out a detailed comparative study of two non-adiabatic local approximations for the xc potential in TDDFT, using quasi-one-dimensional model systems (whose electronic density is uniform in two spatial directions, and nonuniform in the third). The goal was to compare the xc potentials resulting from C-TDDFT and L-TDDFT when a given time-dependent density is used as input, and to determine the magnitude of the resulting non-adiabatic effects when compared to the ALDA. These comparisons were carried out for two types of collective modes: a sloshing and a breathing mode, for a wide dynamical range of amplitudes and frequencies.

In our discussion of non-adiabatic xc effects, we were paying particular attention to dynamical regimes where the two theories, C-TDDFT and elastic L-TDDFT, are known to be exact: C-TDDFT in the linear regime of small-amplitude oscillations, regardless of the frequency, and elastic L-TDDFT in the high-frequency limit, for oscillations of an arbitrary amplitude. This helps to shed light on the so far open question concerning the validity of C-TDDFT in the nonlinear regime: it turns out that, at least in the high-frequency limit, C-TDDFT is surprisingly accurate. Our results show that for moderate deformations (up to 20% of the initial density distribution) errors in the xc potential remain within a few percent. This gives us good reason to believe that this nonlinear formalism should work well for finite frequencies, too (as long as the deformations are not too large).

Let us now address the fundamental question of the meaning and the importance of “non-adiabatic effects” in TDDFT, based on the insights resulting from our study. In analyzing these effects, the language of hydrodynamics, as it is used in C- and L-TDDFT, proves very useful. In general, non-adiabaticity manifests itself through elastic and dissipative components of the electron dynamics, which can be distinguished by their characteristic phase lag with respect to the ALDA. Both elastic and dissipative effects enter the theory via the dependence of xc potential on the deformation tensor. Since the deformation is defined relative to the initial state, this dependence reflects how much of its history the system carries in its memory. A local-in-time dependence on the deformation, which occurs in the high-frequency regime, corresponds to an extremely pronounced memory and a purely elastic xc potential. The dissipative contribution formally appears in the form of a time-nonlocality in the dependence of xc potential on deformations of the electron fluid. In general this time-nonlocality tends to shorten the characteristic memory time, and, as a result, it somewhat diminishes the elastic contribution. In addition it brings about fundamentally new effects, such as relaxation and the corresponding power absorption. Which of the two contributions (elastic or dissipative) is dominant, or whether both play an equally important role, depends on the dynamical regime in which the system under study is evolving. The results from our simple model system lead us to the following conclusions:

Linear regime. For small-amplitude oscillations, dissipation is the most important and dramatic consequence of non-adiabaticity, which leads to qualitative corrections to the adiabatic dynamics. There is a dynamical range which we call “crossover” regime in which the power absorption due to xc retardation effects is maximal. This crossover regime occurs for frequencies that are comparable to an average plasmon frequency for the system. On the other hand, for very low or for very high frequencies, dissipation vanishes. Elastic effects in the linear regime appear to be less important, in a sense that they do not qualitatively change an overall behavior of the xc potential. However, the quantitative effect of elastic corrections to the linearized ALDA can be significant, especially in the high-frequency regime. In particular, for our breathing mode the post-ALDA non-adiabatic corrections are in general of the same order of magnitude as the dynamic part of the ALDA potential itself.

Nonlinear regime. Here, elastic effects become more important, especially at high frequencies, which can lead to substantial, and in certain regimes absolutely dominant contributions to the time-dependent xc potential. As we discussed for the sloshing and breathing modes, the elasticity of the electron liquid tends to oppose attempts to subject the electron system to strong and rapid deformations. This general tendency also naturally explains why C-TDDFT has been successful for...
molecular polarizabilities, which are greatly overestimated in ALDA. Dynamic polarization of the system corresponds to a redistribution of the charge density, i.e., to the deformation of an electron subsystem, which causes a counterracting xc force. This force is an intrinsically non-adiabatic effect that is completely missing in ALDA.

Our results once again illustrate the special role of Kohn’s mode in TDDFT: If the electron dynamics sufficiently resembles Kohn’s mode, as is the case for the sloshing mode considered here, non-adiabatic effects are generally small. On the other hand, for electron dynamics involving high degrees of compression, such as our breathing mode, the non-adiabatic “corrections” can become several times larger than the ALDA potential itself, so that the ALDA completely breaks down, leading to a qualitatively wrong behavior. A striking illustration of such a situation was shown in Fig. 15.

Finally, most of the discussions in this paper, especially those related to dissipative effects, are relevant only for non-adiabaticity in extended systems, where purely electronic dissipation has a well-defined meaning. The situation is less clear when one attempts to describe non-adiabatic effects in finite systems such as atoms and molecules. Here, the elasticity of the electron liquid leads to small shifts of excitation energies, but one also obtains finite linewidths due to dissipation, which is clearly an unwanted effect. The question how non-adiabatic xc potentials for small systems should be constructed thus remains an open issue.

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