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Phys. Rev. Lett. 108, 063003 — Published 9 February 2012
DOI: 10.1103/PhysRevLett.108.063003
The effect of cusps in time-dependent quantum mechanics

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(Dated: November 22, 2011)

Abstract

Within non-relativistic quantum mechanics, spatial cusps in initial wavefunctions can lead to non-analytic behavior in time. We suggest a method for calculating the short-time behavior in such situations. For these cases, the density does not match its Taylor-expansion in time, but the Runge-Gross proof of time-dependent density functional theory still holds, as it requires only the potential to be time-analytic.

PACS numbers: 31.15.ee, 31.10.+z, 03.65.Ge
Within the Born-Oppenheimer (BO) approximation and non-relativistic quantum mechanics, a vast amount of useful information (such as geometry, thermodynamics, and vibrations) of a molecule or solid can be extracted from the electronic ground state. But electronic excitations are important for many areas, from photochemistry to photoemission spectra[1]. Ground-state density functional theory (DFT)[2–4] has been very successful for the former problem, and its time-dependent analog (TDDFT)[5] has become popular for the latter[6]. Within linear response, TDDFT yields useful predictions for the excited states of many molecules[1], and extensions to solids are a keen area of research[7, 8].

All density functional theories rely on a one-to-one correspondence between the one-body potential, such as \(-Z/r\) for an atom, and the density \(n(r)\), under some restrictions. For ground-state DFT, the particle statistics and interaction are fixed. For the time-dependent case, one also specifies the initial wavefunction. The original proof in the ground-state case of Hohenberg and Kohn[2] has been refined over the decades[9–11], but its essence remains unchanged. In the time-dependent problem, after pioneering works by others[12–18], Runge and Gross[5] gave a proof assuming the one-body potential is time-analytic, i.e., equals its Taylor expansion in time around the initial time, for a finite time-interval. Despite recent attempts[19–23], no generally applicable proof has been found that avoids this expansion.

Modern DFT calculations employ the Kohn-Sham scheme[3], in which a fictitious set of non-interacting electrons reproduces the one-electron density of the real system. If we assume that time-analytic potentials yield time-analytic densities, van Leeuwen[24] gave a constructive procedure for finding the TD KS potential for a given system, a problem that has not yet been generally solved for the ground-state case. But, in the usual treatment, matter has cusps in its ground-state electronic wavefunctions at the nuclei, and the resulting spatial non-analyticities are coupled to the time-dependence in the TD Schrödinger equation[21]. Even in the most mundane example, a hydrogen atom in a suddenly switched electric field, this coupling leads to non-analytic short time dynamics, and its density is not time-analytic.

We develop a method for extracting the exact non-analytic short-time behavior of the Schrödinger equation in the presence of cusps. There are distinct spatial regions with different asymptotic behavior for short times, and by ‘asymptotic behavior’ we mean a series expansion around the initial time \(t = 0\), whose error vanishes as \(t \to 0_+\). We calculate the exact short-time behavior for a hydrogen atom in an electric field, and demonstrate agreement with linear response theory in the limit of weak fields. The constructive procedure for
the TD KS potential fails for our examples. (In DFT-speak, the \( v \)-representability question is not solved.) Nonetheless, the original proof of Runge and Gross remains valid.

Consider a single particle prepared in a field-free ground-state, \( \psi_0 \), placed in a static field which is turned on at \( t=0 \), and remains on indefinitely. The exact TD wavefunction is:

\[
\psi(r, t) = \sum_j c_j \exp(-i \epsilon_j t) \phi_j(r),
\]

where \( \epsilon_j \) and \( \phi_j(r) \) are the eigenvalues and functions of the Hamiltonian operator \( \hat{H} \), and \( c_j = \langle \phi_j | \psi_0 \rangle \). We define the time-Taylor series

\[
\psi_{\text{TE}}(r, t) = \sum_{p=0}^{\infty} c_p(r) t^p,
\]

which is the result of the usual practice of Taylor-expanding (TE) the time-evolution operator:

\[
\hat{U}(t) = 1 - i\hat{H}t - \hat{H}^2 t^2/2 + \cdots.
\]

In many cases, the solutions agree, but not necessarily when the wavefunction has non-analyticities in space. Although \( \psi_{\text{TE}} \) formally solves the TDSE, it may not be a valid solution. Holstein and Swift\[25\] reported failure of the TE solution in a model system, in which \( \psi_0 \) has compact support (all space-derivatives of \( \psi_0 \) vanish at the boundary). However, such cases do not occur in routine electronic structure calculations, and may not be of concern in practice. For the rest of this paper, we focus on the ubiquitous cusp in \( \psi_0(r) \) at a nucleus.

Begin with a simple example. Start from the ground-state wavefunction of the hydrogen atom,

\[
\psi_0(r) = \frac{Z^{3/2}}{\sqrt{\pi}} \exp(-Zr).
\]

Our potential for \( t \geq 0 \) is simply zero: The nucleus has been instantaneously vaporized. The exact TD wavefunction can be found by applying the free-particle TD Green’s function to \( \psi_0(r) \), yielding

\[
\psi(r, t) = \frac{Z^{3/2} e^{iZ^2t/2}}{2\sqrt{\pi}r} [f(r, t) - f(-r, t)],
\]

where \( f(r, t) = (r + iZt) \exp(Zr) \text{erfc}[(r + iZt)/\sqrt{2it}] \). Fig. 1 shows the exact solution and the power-series solution as in Eq. (2), which can be summed to all orders:

\[
\psi_{\text{TE}}(r, t) = \frac{Z^{3/2}}{\sqrt{\pi}} e^{-Zr+iZ^2t/2} (1 - i \frac{Zt}{r}), \quad r > 0.
\]
The TE density diverges as \( r \to 0 \) and so is invalid.

FIG. 1. TD exact density and TE density of the ground-state wavefunction of hydrogen atom under free-propagation.

Our method begins with introducing reduced variables:

\[
s = Z \sqrt{t}, \quad \bar{r} = \frac{r}{\sqrt{2t}},
\]

where the powers are motivated by the previous example, and constants chosen for convenience. The form of these reduced variables is closely connected to the Schrödinger equation being 1st order in time but 2nd order in space.[30] These reduced variables are appropriate if the system contains only one nucleus placed at the origin. In reduced variables, the TDSE becomes

\[
\{ \mathcal{L} - is\partial/\partial s + 2s^2v/Z^2 \} \psi = 0,
\]

where

\[
\mathcal{L} = -\nabla^2/2 + i\bar{r} \cdot \nabla,
\]
$Z$ is the nuclear charge, and $v$ is the external potential. In the disappearing-nucleus case, $\psi$ is equal to its Taylor-expansion in powers of $s$ for fixed $\bar{r}$, and thus we try:

$$\psi(\bar{r}, s) = \sum_{p=0}^{\infty} \psi_p(\bar{r}) s^p, \quad (10)$$

yielding:

$$\{\mathcal{L} - m\mathbf{i}\} \psi_m(\bar{r}) = -\frac{2}{Z^2} \sum_{p=0}^{m} v_{p-2}(\bar{r}) \psi_{m-p}(\bar{r}), \quad (11)$$

where $v(\mathbf{r}, t) = \sum_{p} v_{p}(\bar{r}) s^p$. Each power of $s$ in $\psi$ produces a second-order differential equation for a function of $\bar{r}$. The initial wavefunction in reduced variables is $\psi_0(\sqrt{2} \bar{r} s / Z)$. For finite values of $\mathbf{r}$, as $s \to 0$, $\bar{r} \to \infty$, so its expansion as $s \to 0$ determines the large $\bar{r}$ behavior of the $\psi_m(\bar{r})$.

To find the leading non-analytic behavior in $t$ due to a cusp, we rearrange the TE solution of Eq. (2) in terms of the reduced variables $s$ and $\bar{r}$, and test its validity power by power in $s$. Each $\psi_{m}^{\text{TE}}$ must satisfy the differential equation, the boundary condition derived from the initial wavefunction as $\bar{r} \to \infty$, and remain finite at the nucleus ($m$ being the order of the $s$-expansion). We do this recursively, until we find $k$, the lowest value of $m$ for which $\psi_{k}^{\text{TE}}$ fails. Define

$$\xi(\bar{r}, s) = \psi(\bar{r}, s) - \psi_{\text{TE}}(\bar{r}, s) \quad (12)$$

and solve the differential equation for $\xi_k(\mathbf{r})$ to find the leading non-analytic behavior due to the cusp.

This can be quite demanding as, in 3D, the equation is a partial differential equation. A simple approach is to perform a local analysis using the method of dominant balance[26] to generate an asymptotic series in inverse powers of $\bar{r}$. Repeated application yields the series to all orders, but this is insufficient to ensure that $\psi$ remains finite at the $r = 0$. This difficulty can be overcome by performing a Borel resummation of the asymptotic series[26] to find the exact solution. The conditions where Borel resummation works are discussed in [27].

As a second example, we consider a hydrogen atom in a suddenly switched electric field which stays on indefinitely, the initial wavefunction is that of Eq. 4, and the potential after $t = 0$ is

$$v(\mathbf{r}) = -Z/r + \mathcal{E} z, \quad (13)$$
where $E$ is the amplitude of the electric field. This situation is covered by the Runge-Gross theorem as a potential that is analytic in time (in fact, constant) with an initial wavefunction that is not its instantaneous ground state. The time-dependent dipole moment $\mu(t)$ determines the photoabsorption of the atom via

$$\sigma(\omega) = \frac{4\pi\omega}{c} \int_0^\infty d\tau \left[ \frac{\partial \mu(t)}{\partial t} \right]_{t=\tau} \sin(\omega \tau),$$  \hspace{1cm} (14)$$

known analytically\textsuperscript{[28, 29]} for the H atom as $E \rightarrow 0$.

To apply our method, we first calculate the TE wavefunction order-by-order. We find each term satisfies both the differential equation and boundary conditions until the 4th order:

$$\psi_{4}^{\text{TE}} = \frac{Z^{3/2}}{24\sqrt{\pi}} (-3 + 12i\bar{r}^2 + 4\bar{r}^4) + \mathcal{E} \frac{i\bar{z}}{12\sqrt{\pi}Z^{3/2}\bar{r}^3} (1 + 6i\bar{r}^2 + 24\bar{r}^4),$$ \hspace{1cm} (15)$$

which works except that it diverges at the nucleus. Assume a correction of the form:

$$\xi_4 = \exp(S(\bar{r})) \bar{z},$$ \hspace{1cm} (16)$$

because the applied field is linear in $z$. This yields an ordinary differential equation for $S$:

$$S'' + (S')^2 - 2i\bar{r}S' + 4S'/\bar{r} + 6i = 0,$$ \hspace{1cm} (17)$$

and applying the method of dominant balance recursively, one finds the asymptotic expansion:

$$\xi_4 = c_1 \left( \bar{r}^3 + \frac{9i\bar{r}}{2} - \frac{9}{4\bar{r}} + \frac{3i}{8\bar{r}^3} \right) + c_2 e^{i\bar{r}^2/\bar{r}^8} \left[ 1 + \frac{1}{\bar{r}^2} \sum_{m=0}^{\infty} \frac{(-i)^{m+1}(m + 4)(2m + 6)!}{(m + 1)!2^{2m+5} \cdot 9\bar{r}^{2m}} \right].$$ \hspace{1cm} (18)$$

Since $\psi_{4}^{\text{TE}}$ satisfies the boundary condition at large $\bar{r}$, $\xi_4$ must vanish here, and $c_1 = 0$. But this asymptotic series does not yield the small $\bar{r}$ limit. To find this, we perform a Borel resummation, yielding

$$\xi_4 = c_2 \frac{1 + i}{72\bar{r}^{3}} \left[ (2 + 2i)e^{i\bar{r}^2\bar{r}}(-3 + 16i\bar{r}^2 + 4\bar{r}^4) \right.$$  

$$- \sqrt{2\pi}(3i - 18i\bar{r}^2 + 36i\bar{r}^4 + 8\bar{r}^6) \text{erfc} \left( \frac{1 - i}{\sqrt{2}} \bar{r} \right) \right]$$ \hspace{1cm} (19)$$

Requiring $\psi_4$ to remain finite yields

$$c_2 = (1 - i)\mathcal{E} / (2\pi Z^{3/2}).$$ \hspace{1cm} (20)$$
One can easily check analytically that $\psi_4 = \psi_4^{\text{TE}} + \xi_4$ is then the unique solution to Eq. (11) for $m = 4$.

With $\psi_4$, we find the leading half-power of the wavefunction by changing the variables back to $(r, t)$ and taking the asymptotic expansion around the $t = 0$. The leading time-half-power is then

$$-\frac{(8 - 8i)E Z^5/2 e^{ir^2/2t} \cos \theta}{\pi r^7} t^{11/2}, \quad r \gg \sqrt{t},$$

(21)

and one can show $\psi_{m>4}$ contributes to higher order time-half-powers by the method of dominant balance. This is also the lowest order in $t$ at which rapid non-time-analytic oscillations of the phase appear.

Our derivation applies for any value of $E$. The one-body potential appears linearly in our equations for $\psi_n(\vec{r})$, and only for $n \geq 3$. Thus if $\psi_4(\vec{r})$ produces the leading non-analytic behavior in $t$, then this behavior is linear in $E$, and is determined exactly by linear response theory. (Higher-order effects in $E$ produce changes in other terms, but not the leading non-analytic short-time behavior.) To see this, take the Lehmann representation of the Green’s function of the hydrogen atom and evaluate the change in the wavefunction to first order in $E$. We did not find a closed-form, but the stationary phase approximation to the integral over the wave number yields the leading half-power term in agreement with our short-time result Eq. (21).

The leading half-power term in the induced dipole moment calculated using linear response theory yields:

$$\mu(t \to 0^+) \sim \cdots - 256 Z^5 / (2835 \sqrt{\pi}) t^{9/2} + \cdots.$$  

(22)

Its Fourier transform yields the known high-frequency decay of the photoabsorption cross-section

$$\sigma(\omega \to \infty) \sim 16 \sqrt{2} Z^5 \pi / (3c \omega^{7/2}).$$

(23)

Thus the cusp at the nucleus leads to fractional powers in time-evolution and fractional powers in frequency decay.

The leading half-power term in $\sigma(\omega \to \infty)$ calculated with $\psi_4$ differs from Eq. (23) by a factor of 2. In calculating $\sigma(\omega)$ from $\psi(\vec{r}, t)$, one must integrate over $r$, and the coupling between $r$ and $t$ in $\psi$ (as seen in $\psi_4$) allows higher order terms in the $s$-expansion to contribute to the leading half-power term in $\sigma(\omega \to \infty)$ as well. In this case, $\psi_4$ and $\psi_5$ both contribute to the $\omega^{-7/2}$ term in $\sigma$.  

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Before discussing the consequences for TDDFT, we again consider the simpler, but starker example of the disappearing nucleus case. Our method correctly shows the error appearing at $s^1$ order, and produces the correct short-time behavior:

$$\psi(|r| \gg \sqrt{t}, t \to 0^+) \sim \frac{Z^{3/2}e^{-Zr}}{\sqrt{\pi}} + \frac{iZ^{5/2}(Zr - 2)e^{-Zr}}{2\sqrt{\pi r}}t$$

$$- \frac{Z^{9/2}(Zr - 4)e^{-Zr}}{8\sqrt{\pi r}}t^2 - \frac{(2 + 2i)Z^{5/2}e^{ix^2/(2t)}}{\pi r^4}t^{5/2} + \cdots$$

while its TE contains only integer powers of $t$.

Although less relevant for TDDFT, our method applies equally well in 1d and proves that the short-time behavior for the initial state with cusp example in [21] originates from $s^1$ order and has fractional power $t^{3/2}$, again with a rapid non-time-analytic phase. Similarly, it’s $s^4$ order for a particle in a delta-well with an applied electric field, and the leading short-time behavior in $\psi$ is

$$- \frac{(4 + 4i)eZ^{3/2}\exp[ix^2/(2t)]}{\sqrt{\pi x^5}}t^{9/2}, \quad |x| \gg \sqrt{t},$$

consistent with the known polarizability and its high-frequency limit.[30]

Our results are proven only for single electrons. However, the nuclear potential dominates over the electron-electron repulsion in the region of the nucleus, so that the cusp condition on the time-dependent density near a nucleus remains valid regardless of the interelectron repulsion. Thus we expect the qualitative features (i.e., half-powers of $t$ in the short-time behavior) to remain true even in the presence of interaction.

We conclude by discussing the relevance of our results for TDDFT, whose theorems certainly apply (or not) for $N = 1$. The proof of a one-to-one correspondence[5] shows that, for two $t$-analytic potentials whose Taylor expansions first differ in the $k$-th order, the difference in the $(k + 2)$th-order $t$-derivative of the densities is non-zero. Thus the two densities differ, the Runge-Gross proof applies, and the potential is a functional of the density under the conditions stated[5]. Whether or not the TE density matches the true density is irrelevant.

On the other hand, the KS potential of Ref. [24] produces a density whose $t$-derivatives equal those of the interacting density. If we assume that the density is $v$-representable by a $t$-analytic potential (e.g., if the KS and interacting wavefunctions themselves are $t$-analytic), then that potential will yield this density. But the wavefunctions are not $t$-analytic in the examples given here, and typically are not when an initial wavefunction with a cusp
undergoes a non-trivial evolution. In general, a more sophisticated procedure is needed to ensure the constructed potential generates the desired density, and some progress has been made [21–23]. This is important as the constructive procedure has been invoked or applied as is to a variety of situations [31–33].

One might argue that real atoms have finite nuclei, or that real molecules and solids have nuclear wavefunctions that smear the cusps due to nuclei, as well as that most of the practical TDDFT calculations are done on cusp-free Gaussian basis sets. But such arguments miss the basic point. Just as in ground-state DFT, time-dependent DFT is an exact mapping of the quantum mechanics of electrons, for which there are no difficulties within BO or need for finite nuclei.

We thank Tchavdar Todorov for very useful discussions. ZY and KB acknowledge U.S. D.O.E funding (DE-FG02-08ER46496) and NTM acknowledges both NSF and the Research Corporation.

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