Exterior time scaling with the stiffness-free Lanczos time propagator: Formulation and application to atoms interacting with strong midinfrared lasers

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Aiming at efficient numerical treatment of tunneling ionization of atoms and molecules by mid-infrared (IR) lasers, exterior time-scaling (ETS) theory is formulated as a generalization of the time-scaled coordinate approach. The key idea of ETS is the division of the spatial volume into a small region around the nucleus and its outside; the radial coordinates are time scaled only in the outer region. The continuum components of photoelectron wave packets are prevented from reaching the edge of the spatial simulation volume, enabling the long-time evolution of wave packets with a relatively small number of basis functions without concerns of electron reflections. On the other hand, the bound-state components are free from shrinking toward the origin because of non-time scaling in the inner region. Hence, the equations of motion in ETS are less stiff than the ones in the original time-scaled coordinate approach in which the shrinking bound states make the equations of motion seriously stiff. For numerical implementation of ETS, the working equations are derived in terms of finite-element discrete-variable-representation functions. Furthermore, the stiffness-free Lanczos time propagator is introduced to remove any persistent stiffness in the treatment of mid-IR lasers due to the involvement of hundreds of angular momentum states. The test calculations for atomic hydrogen interacting with linearly polarized mid-IR pulses demonstrate the accuracy and numerical efficiency of this scheme, and exhibit its special capability if there is no recollision with the parent ion. Hence, ETS will show its true potential for the detailed analysis of photoelectron wave-packet dynamics in circularly or near-circularly polarized mid-IR fields.

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I. INTRODUCTION

The recent advance of intense few-cycle light sources in the midinfrared (IR) region (wavelength: $\lambda \geq 3 \mu$m) is leading strong-field physics to a new direction [1]. Its extreme nonlinear effect on matter generates high-order harmonic radiation covering the x-ray region [2,3] with potentially narrowing the pulse width down to the zettosecond [4], and is hence of very practical importance. Longer wavelength also ensures more detailed analysis and interpretation of experimental and numerical results, based on semiclassical (see, e.g., Refs. [5–8]) and adiabatic [9,10] theories. Circularly or near-circularly polarized mid-IR pulses are ideal for precise attoclock measurements to elucidate tunneling dynamics [11,12], and for obtaining the information of molecular orbital structure from the photoelectron momentum distribution perpendicular to the polarization plane [13]. Combining mid-IR lasers having different polarizations and colors may realize new means to reveal and control electron dynamics (see, e.g., recent references [14–17]). Numerical treatment of atoms and molecules in the mid-IR region, however, remains very challenging even within the single-active-electron (SAE) approximation [18]. The reason comes not only from the involvement of many angular momentum states in the wave function, but also from the need to set the size of spatial volume proportional to $\lambda^2$, and also from the ponderomotive energy $U_p \propto \lambda^2$ which requires the use of denser grids or many basis functions as $\lambda$ increases [19].

To manage the increasing numerical difficulty for large $\lambda$, the spatial volume is usually set as small as possible by employing a complex absorbing potential (CAP) [20] or exterior complex scaling (ECS) [21,22], which prevents the high-energy continuum part of electron wave packets from reflection. This strategy is well suited for the analysis of low-energy photoelectrons and for computing dipole acceleration to high-order-harmonic generation (HHG). An alternative strategy is to enlarge the spatial volume as time grows to prevent the reflection without losing the norm of wave function. Treatment of the time-dependent Schrödinger equation (TDSE) by periodic von Neumann basis with bi-orthogonal exchange [22] enables the extension of volume, which, however, results in a gradual increase of numerical cost. On the other hand, the time-scaled coordinate approach causes no such cost increase, and has, after its introduction in collision physics [24,25], been developed aiming at efficient treatment of ionization by photoabsorption and electron impact [26,27]. After a long-time propagation under field-free conditions, the photoelectron wave packet in the scaled coordinate becomes stationary, from which the ionization cross section is extracted without projection onto the scattering wave function. This is advantageous in the treatment of many-electron systems, in particular, for the computation of the ndouble or multiple ionization cross section [28–30]. However, the time-scaled coordinate approach has a very serious shortcoming: The bound-state part of the electron wave packet shrinks toward the origin as time goes. Using dense grids or many basis functions around the origin to account for this shrinking makes the equations of motion stiff (see, e.g., Ref. [31] for a discussion of stiffness of differential
In description of tunneling ionization at large λ, the equations become much stiffer and numerically untractable due to the involvement of hundreds of angular momentum states and the increase of the centrifugal potential barrier in the Hamiltonian. Although employing, e.g., Fatunla’s method [31 32 33 52] or other elaborate time propagators may manage the problem, it is unfavorable to deploy many basis functions for the bound states since the research interest is in the description and analysis of the outgoing continuum part of the wave packet.

The aim of this paper is twofold: (i) formulation of exterior time-scaling (ETS) theory for extending numerical exploration toward the mid-IR region, and (ii) establishment of its stiffness-free numerical implementation. The ETS theory is a generalization of the original time-scaled coordinate approach which is hereafter, for comparison, referred to as global time scaling (GTS). The idea of ETS is to divide the spatial volume into two parts: a small domain around the nucleus, and its outside. The time scaling is carried out only to radial coordinates in the outer region. The continuum components in the outer region are hence prevented from reaching the edge of the spatial simulation volume, enabling the long-time evolution of the wave packet with a relatively small number of basis functions. In the inner region, on the other hand, the bound states are free from shrinking because of non-time scaling, and the equations of motion are expected to be less stiff than the ones in GTS. As many angular momentum states are involved, however, the equations in ETS inevitably become stiff. To address this problem, we propose a stiffness removal procedure which is in particular suitable for wave functions expanded in terms of finite-element discrete-variable-representation (FEDVR) functions (see, e.g., Ref. [36]) and time propagated by the Lanczos algorithm [37]. This procedure is not specific to the ETS implementation but will also be applicable to a general class of equations appearing in atomic and molecular physics. Also note that, although in the following this paper aims at the treatment of one-electron atoms, the ETS method with the stiffness-free time propagation is applicable to the TD (restricted/generalized)-active-space configuration-interaction [TD-(RAS/GAS)CI] approach [38 39] and may also be so to the R-matrix theory including time dependence (RMT) [40 42], for instance, and can therefore be flexibly used to investigate many-electron atoms and molecules.

This paper is organized as follows. The ETS theory is formulated for atomic hydrogen in Sec. II. Expanding the wave function in terms of FEDVR functions, Sec. III shows the derivation of the working equations for practical ETS implementation. Then Sec. IV is devoted to the analysis of stiffness and the discussion of its removal; the stiffness-free FEDVR-based Lanczos algorithm proposed in this section has a wide range of applicability. Based on the ETS theory with the stiffness-free procedure, Sec. V demonstrates the tunneling ionization of atomic hydrogen in linearly polarized mid-IR lasers, and shows the accuracy and numerical efficiency of ETS. Section VI concludes this work and provides an outlook. Atomic units are used throughout unless otherwise stated.

II. FORMULATION

We consider atomic hydrogen prepared in the ground state which then starts interacting with a light pulse linearly polarized along the z axis. This simple case is considered just for notational simplicity in the formulation; the generalization to many-electron atoms and molecules in arbitrarily polarized light fields is, at least formally, straightforward in any coordinate system. Expanding the wave function in terms of spherical harmonics with magnetic quantum number \( m = 0 \),

\[
\Psi(r, t) = \frac{1}{\sqrt{r}} \sum_\ell \psi_\ell(r, t) Y_\ell^0(\Omega),
\]

the TDSE leads to a set of coupled equations for the radial functions:

\[
i \left( \frac{\partial}{\partial t} \right)_r \psi_\ell(r, t) = \left[ -\frac{1}{2} \frac{\partial^2}{\partial r^2} + V_\ell(r) \right] \psi_\ell(r, t) + \sum_{\ell'} W_{\ell\ell'}(t) \psi_{\ell'}(r, t),
\]

(2)

where

\[
V_\ell(r) = \frac{\ell(\ell + 1)}{2r^2} - \frac{1}{r},
\]

(3)

and the light-atom interaction operator is treated within the dipole approximation,

\[
g_{\ell\ell'} F(t) r,
\]

(4)

with \( g_{\ell\ell'} = \sqrt{4\pi/3} \int Y_\ell^0(\Omega)Y_\ell^0(\Omega)Y_{\ell'}^0(\Omega) d\Omega \), and the vector potential \( A(t) \) and the electric field \( F(t) = -A(t)/\partial t \) of light. The gauge-specific notation,

\[
W_{\ell\ell'}(t) = \begin{cases} W_{\ell\ell'}^L(r, t) = \frac{W_{\ell\ell'}^N(r, \partial_r, t) = -ig_{\ell\ell'} A(t)}{2r} + \frac{\ell'(\ell' + 1) - \ell(\ell + 1)}{2r} \end{cases},
\]

on page 2
or \( \xi \). Such an explicit notation is not needed in Eq. (2) but in the following helps avoid unnecessary confusion.

The formulation of ETS commences with introducing a spherical surface, \( \Sigma \), the radius of which is \( r_\Sigma \). Setting the center of \( \Sigma \) at the origin, the configuration space is divided into its inner and outer regions. Let \( \Sigma \) itself belong to the inner region. We then define an ETS map by

\[
\xi(r, t) = \begin{cases} 
  r, & (0 \leq r \leq r_\Sigma), \\
  r_\Sigma + (r - r_\Sigma)/R(t), & (r_\Sigma < r < \infty),
\end{cases}
\]

(5)

where \( R(t)(\geq 1) \) is a smooth increasing function of time; its explicit form is given later [see Eq. (33) in Sec. VII]. Noting the mutual dependence between \( r \) and \( \xi \), i.e., \( r = r(\xi, r) \) and \( \xi = \xi(r, t) \), the differentiation of Eq. (5) reads \( dr(\xi, t) = d\xi(r, t) \) for \( 0 \leq r \leq r_\Sigma \), and \( dr(\xi, t) = [\xi(r, t) - \xi(r_\Sigma)]dR(t) + R(t)d\xi(r, t) \) for \( r_\Sigma < r < \infty \). Hence the differential equation, \( dr(\xi, t) = 0 \), leads to relations:

\[
\left( \frac{\partial}{\partial t} \right)_r \xi = \left( \frac{\partial}{\partial \xi} \right)_r \xi,
\]

(0 \leq r \leq r_\Sigma)

(6)

\[
\left( \frac{\partial}{\partial \xi} \right)_r \xi = \frac{\dot{R}(t)}{R(t)}(\xi - r_\Sigma) \frac{\partial}{\partial \xi} \xi,
\]

(6)

\[
(r_\Sigma < r < \infty)
\]

In \( 0 \leq r \leq r_\Sigma \), Eq. (5) is just an identity mapping. The coupled equations obeyed by the radial functions in \( 0 \leq \xi \leq r_\Sigma \) are thus obtained from Eq. (2) by simply replacing \( r \) by \( \xi \):

\[
i \left( \frac{\partial}{\partial \xi} \right) \psi_\ell(r, t) = \left[ -\frac{1}{2} \frac{\partial^2}{\partial \xi^2} + V_\ell(\xi) \right] \psi_\ell(r, t) + \sum_{\ell'} W_{\ell\ell'}(t) \psi_{\ell'}(r, t), \quad (0 \leq \xi \leq r_\Sigma).
\]

(7)

In \( r_\Sigma < r < \infty \), on the other hand, using the second lines of Eqs. (5) and (6) in Eq. (2) leads to

\[
i \left( \frac{\partial}{\partial \xi} \right) \psi_\ell[r_\Sigma + R(t)(\xi - r_\Sigma), t] = \left[ -\frac{1}{2[R(t)]^2} \frac{\partial^2}{\partial \xi^2} + V_\ell[r_\Sigma + R(t)(\xi - r_\Sigma)] + \frac{\dot{R}(t)}{R(t)}(\xi - r_\Sigma) \frac{\partial}{\partial \xi} \right] \psi_\ell[r_\Sigma + R(t)(\xi - r_\Sigma), t]
\]

\[
+ \sum_{\ell'} W_{\ell\ell'}(t) \psi_{\ell'}[r_\Sigma + R(t)(\xi - r_\Sigma), t], \quad (r_\Sigma < \xi < \infty).
\]

(8)

Defining new radial functions by

\[
\phi_\ell(\xi, t) = \sqrt{R(t)} \exp \left[ -iR(t)\dot{R}(t)(\xi - r_\Sigma)^2/2 \right] \psi_\ell(r, t), \quad (r_\Sigma < r < \infty),
\]

(9)

Eq. (8) is recast into a concise form:

\[
i \left( \frac{\partial}{\partial \xi} \right) \phi_\ell(\xi, t) = \left[ -\frac{1}{2[R(t)]^2} \frac{\partial^2}{\partial \xi^2} + V_\ell[r_\Sigma + R(t)(\xi - r_\Sigma)] + \frac{R(t)\dot{R}(t)}{2}(\xi - r_\Sigma)^2 \right] \phi_\ell(\xi, t) + \sum_{\ell'} W_{\ell\ell'}(t) \phi_{\ell'}(\xi, t),
\]

(10)

\[
(r_\Sigma < \xi < \infty),
\]

with

\[
W_{\ell\ell'}(t) = \begin{cases} 
  W_{\ell\ell}'[r_\Sigma + R(t)(\xi - r_\Sigma), t] & \text{in length gauge}, \\
  W_{\ell\ell}'[r_\Sigma + R(t)(\xi - r_\Sigma), \partial R(t)\xi, t] + g_{\ell\ell'} A(t)\dot{R}(t)(\xi - r_\Sigma) & \text{in velocity gauge},
\end{cases}
\]

(11)

where \( g_{\ell\ell'} A(t)\dot{R}(t)(\xi - r_\Sigma) \) comes from the exponent in Eq. (9) (a corresponding factor, \( A(\xi, t)\dot{R}(t)\xi \), is missing in Eq. (8) of Refs. [31, 32] where GTS is formulated for one-dimensional systems). Importantly, although Eqs. (7) and (10) are defined in non-overlapping radial intervals, \( 0 \leq \xi \leq r_\Sigma \) and \( r_\Sigma < \xi < \infty \), respectively, and may seem to be independent of each other, the equations are indeed coupled by the requirement of continuous differentiability of the wave function across \( \Sigma \):

\[
\Psi(r, t)|_{r=\xi} = \Psi(r, t)|_{r=r_\Sigma+0} \quad \text{and} \quad \partial \Psi(r, t)/\partial r|_{r=r_\Sigma+0}.
\]

\[
\partial \Psi(r, t)/\partial r|_{r=r_\Sigma+0} \quad \text{i.e., for each } \ell,
\]

\[
\psi_\ell(r_\Sigma, t) = \frac{1}{\sqrt{R(t)}} \phi_\ell(r_\Sigma + 0, t),
\]

(12a)

\[
\left. \frac{\partial}{\partial \xi} \psi_\ell(\xi, t) \right|_{\xi=r_\Sigma} = \frac{1}{\sqrt{R(t)}} \left. \frac{\partial}{\partial \xi} \phi_\ell(\xi, t) \right|_{\xi=r_\Sigma+0}.
\]

(12b)

It is readily seen that ETS is a generalized concept of GTS; setting \( r_\Sigma = 0 \) reduces the ETS map [Eq. (5)] to the original GTS map, \( \xi(r, t) = r/R(t) \); Eq. (10) becomes, e.g., Eq. (8) in Refs. [31, 32] except for the
FIG. 1. (Color online) Illustration of the shrinking of the ground-state wave function in GTS and the non-shrinking in ETS. The Coulomb potential function, $1/r(\xi,t)$ (thin lines), and the radial part of the ground-state wave function of atomic hydrogen, $\psi_{1s}[r(\xi,t)] = 2r(\xi,t) \exp[-r(\xi,t)]$ [thick (red) lines, shifted $-0.5$ downward], are plotted at zeroth, first, second, and third optical cycles (o.c.), i.e., at $t = (2\pi/\omega) \times n$ ($n = 0, 1, 2,$ and $3$). The temporal unit is measured in terms of one optical cycle of a mid-IR laser with $\lambda = 3 \mu\text{m} = 5.669 \times 10^4$. The plot is made under laser-free conditions, but, in connection with the numerical demonstration later, the top-right corner displays the electric field $F(t) = (-dA(t)/dt)$ of the same three-cycle pulse used in Sec. V A. The time-scaling factor $R(t)$ is specified by Eq. (33) with $R_\infty = 0.01$, which is also the same as in Sec. V. The value of $r_\Sigma$ is set to zero in GTS [(a1–3)], and $30$ in ETS [(b1–3)]. The shaded region in (b1–3) indicates the radial interval of the inner region ($0 \leq r \leq r_\Sigma$) where the bound states are free from shrinking because of non-time scaling. The top and bottom of each panel represent the $r(=r(\xi,t))$ and $\xi$ coordinates, respectively.

missing factor in the velocity gauge. As in the equations of motion in the conventional GTS, Eq. (10) indicates the introduction of an effective nucleus charge, $1/R(t)$, an effective electron mass, $[R(t)]^2$, and a temporal harmonic potential only while $\ddot{R}(t) > 0$, preventing the electron from escaping to infinity. An important difference of ETS from GTS is that ETS avoids the complication of the shrinking of the bound states toward the origin present in GTS because of the time scaling only in the outer region. Figure 1 gives the comparison by illustrating the Coulomb potential function, $1/r(\xi,t)$, and the radial part of the ground-state wave function of atomic hydrogen, $\psi_{1s}[r(\xi,t)] = 2r(\xi,t) \exp[-r(\xi,t)]$. The plot is made under laser-free conditions, but, supposing the application to mid-IR lasers with $\lambda = 3 \mu\text{m} = 5.669 \times 10^4$ (see Sec. V A), the temporal unit is measured in its optical cycle (o.c.), $2\pi/\omega = 413.7$, where $\omega = 2\pi/\alpha\lambda = 0.01519$ with the fine-structure constant $\alpha = 1/137.036$. The time-scaling factor $R(t)$ is specified by Eq. (33) with $R_\infty = 0.01$ (see the discussion in Sec. V). While the Coulomb potential function and the radial function in GTS shrink toward the origin in
the ξ coordinate [Figs. 1(a1–3)], they remain almost unchanged in ETS [Figs. 1(b1–3)] even across \( r_{\Sigma} = 30 \).

III. PRACTICAL FEDVR-BASED FORMALISM

The numerical implementation of ETS requires the expansion of the radial functions, \( \psi_t(\xi, t) \) and \( \phi_t(\xi, t) \), of Eqs. (7) and (10), in terms of a set of basis functions. There may be several options: grids, B splines, a variety of DVR functions, and combinations of them in the inner and outer regions. Among the various possibilities, we choose the FEDVR functions and derive the working equations for the ETS implementation.

A. FEDVR functions

Let a closed interval, \([0, \xi_{\text{max}}]\), with \( \xi_{\text{max}} > r_{\Sigma} \), be the range of the scaled-radial coordinate, \( \xi \). We then define a set of FEDVR functions over the interval as follows: Let the two intervals, \([0, r_{\Sigma}]\) and \((r_{\Sigma}, \xi_{\text{max}}] \), divided at \( \xi = r_{\Sigma} \), be further subdivided into \( N_{\text{fe}} \) and \( N_{\text{fe}} \) intervals (FEs), respectively:

\[
[0, r_{\Sigma}] = [\xi_0, \xi_1] \cup [\xi_1, \xi_2] \cup \cdots \cup [\xi_{N_{\text{fe}}-1}, \xi_{N_{\text{fe}}}]. \tag{13a}
\]

\[
(r_{\Sigma}, \xi_{\text{max}}] = [\xi_{N_{\text{fe}}} + \xi_{N_{\text{fe}}-2}, \xi_{N_{\text{fe}}-1}] \cup [\xi_{N_{\text{fe}} + \xi_{N_{\text{fe}}-1}}, \xi_{N_{\text{fe}} + \xi_{\text{max}}}], \tag{13b}
\]

where \( \xi_0 = 0, \xi_{N_{\text{fe}}} = r_{\Sigma} \), and \( \xi_{N_{\text{fe}} + \xi_{\text{fe}}} = \xi_{\text{max}} \). Let every interval have the same width \( \Delta \xi = \xi_{\text{max}}/(N_{\text{fe}} + N_{\text{fe}}) \). Let us then define the Lobatto DVR functions with \( N_{\text{dvr}} \) quadrature points. Following the standard prescription (see, e.g., Ref. [6]), a set of the FEDVR functions, \( \{ \chi_{K}(\xi) | K = 1, \cdots, (N_{\text{dvr}}-1)(N_{\text{fe}} + N_{\text{fe}})-1 \} \), is composed. In the construction, two Lobatto DVR functions having quadrature points at \( \xi_0 \) and \( \xi_{\text{max}} \) are removed; consequently, every FEDVR function is zero at both edges of the domain, i.e., \( \chi_0(0) = \chi_{\text{max}}(\xi_{\text{max}}) = 0 \).

The FEDVR functions are then classified into three groups: \( \{ \chi_{K}(\xi) | K = 1, \cdots, (N_{\text{dvr}}-1)N_{\text{fe}}-1 \}, \{ \chi_{\text{b}}(\xi) \}, \{ \chi_{\text{d}}(\xi) \} \), and \( \{ \chi_{\text{d}}(\xi) | K = (N_{\text{dvr}}-1)N_{\text{fe}} + 1, \cdots, (N_{\text{dvr}}-1)(N_{\text{fe}} + N_{\text{fe}}) \}\).

Note that the FEDVR functions belonging to the first and third groups, respectively, are non-vanishing only below and above \( r_{\Sigma} \), and are distinguished, if need be, by the accent symbols \( \hat{,} \) atop the index \( \kappa \). On the other hand, the second group consists only of \( \chi_{\text{b}}(r) \), which is the bridge function across \( r_{\Sigma} \) (see, e.g., Ref. [36] for a discussion of the bridge function). Figure 2 illustrates the setup of the FEDVR functions.

B. Exterior time scaling with FEDVR functions

Let \( t_{\text{max}} \) be the maximum value of \( t \) in the expansion of \( \Psi(r, t) \) [Eq. (11)]. The radial function for each \( \ell \) is then expanded in terms of the FEDVR functions defined in Sec. IIIA.

\[
\psi_t(\xi, t) = \sum_{\kappa} a_{\kappa\ell}(t) \chi_{\kappa}(\xi) + \sqrt{\frac{2}{1 + R(t)}} a_{\ell}(t) \chi_{\text{b}}(\xi), \tag{14a}
\]

\[
\phi_t(\xi, t) = \sum_{\kappa} b_{\kappa\ell}(t) \chi_{\kappa}(\xi) + b_{\ell}(t) \chi_{\text{b}}(\xi), \tag{14b}
\]

where the extra factor, \( \sqrt{2/[1 + R(t)]} \), attached to \( a_{\ell}(t) \) is not just cosmetic but helps construct the working equations in Hermitian matrix form (as discussed in the last paragraph in this section). The continuity condition of the wave function, Eq. (12a), now leads to

\[
\sqrt{\frac{2}{1 + R(t)}} a_{\ell}(t) = \frac{1}{\sqrt{R(t)}} b_{\ell}(t). \tag{15}
\]

We should thus derive the equations obeyed by the expansion coefficients, \( \{ a_{\kappa\ell}(t) \} \), \( \{ b_{\kappa\ell}(t) \} \), and \( \{ a_{\ell}(t) \} \) (or alternatively \( \{ a_{\kappa\ell}(t) \} \), \( \{ b_{\kappa\ell}(t) \} \), and \( \{ b_{\ell}(t) \} \); in this latter case, the factor, \( \sqrt{2/[1 + R(t)]} \), attached to \( a_{\ell}(t) \) in Eq. (14a) should be replaced by \( 1/\sqrt{R(t)} \)). For simplicity, the rest of this subsection is devoted to the derivation only in the length gauge. The Appendix lists the instructions for formulating the working equations in the velocity gauge.

It is straightforward to derive the equations for evaluating the time derivative of \( a_{\kappa\ell}(t) \) and \( b_{\kappa\ell}(t) \). Substituting Eq. (14a) into Eq. (7), and multiplying both sides by \( \chi_{\kappa}(\xi) \), we integrate them with respect to \( \xi \) over \([0, r_{\Sigma}]\) and obtain

\[
\frac{id}{dt} a_{\kappa\ell}(t) = \sum_{\kappa'} \left\{ \left[ \frac{1}{2} \int_{0}^{\xi_{\text{max}}} \frac{d\chi_{\kappa}(\xi)}{d\xi} \frac{d\chi_{\kappa'}(\xi)}{d\xi} d\xi + \delta_{\kappa\kappa'} V_{\ell}(\xi_{\kappa}) \right] + \delta_{\kappa\kappa'} W_{\ell\ell'}^{\text{Li}}(\xi_{\kappa}, t) \right\} a_{\kappa\ell'}(t) \\
+ \frac{a_{\ell}(t)}{\sqrt{2[1 + R(t)]}} \int_{0}^{\xi_{\text{max}}} \frac{d\chi_{\kappa}(\xi)}{d\xi} \frac{d\chi_{\ell}(\xi)}{d\xi} d\xi. \tag{16}
\]

Similarly, substituting Eq. (14b) into Eq. (10), and multiplying both sides by \( \chi_{\text{b}}(\xi) \), the integration of them with
Similarly, substituting Eq. (14b) into Eq. (10), multiplying both sides by 2 in obtaining Eqs. (18a) and (18b), we used the relation, 2 \int_{r_\Sigma}^{r_{\Sigma+\Delta r}} \left( \frac{d\chi_{\Sigma}(\xi)}{d\xi} \right)^2 d\xi = \int_{r_\Sigma}^{r_{\Sigma+\Delta r}} \left( \frac{d\chi_{b}(\xi)}{d\xi} \right)^2 d\xi = 2 \int_{r_\Sigma}^{\xi_{\max}} \left( \frac{d\chi_{\Sigma}(\xi)}{d\xi} \right)^2 d\xi,

as well as the DVR quadrature rule for the bridge function: For an arbitrary integrable function \( f(\xi) \) defined in
where all the almost-equal signs in Eq. (20) become exact-equal signs if $f(\xi)$ is linear in $\xi$ in $[\xi_{N_\text{ini}} - 1; \xi_{N_\text{fin}}]$ (see, e.g., Ref. 13). Differently from Eqs. (16) and (17), Eqs. (18a) and (18b) contain the surface terms at $r_\Sigma$ because of $\chi_0(r_\Sigma) \neq 0$. In dealing with the surface terms, the Bloch operator, $L = \frac{1}{2} \delta (r - r_\Sigma) \frac{\partial}{\partial \xi}$, is a useful device as in the formulation of $R$-matrix-related theories (see, e.g., Refs. 14, 15). In the interest of keeping the derivation mathematically clear, however, we do not employ the Bloch operator the delta function of which could cause an ambiguity about whether $\Sigma$ belongs to its inside or outside or, perhaps, both sides or neither side. Getting back to the derivation, canceling out the surface terms in Eqs. (18a) and (18b) using Eq. (12a), we finally obtain

\[
\int_0^{\xi_{\text{max}}} \chi_0(\xi) f(\xi) \chi_\kappa(\xi) d\xi \approx \int_0^{\xi_{\text{max}}} \chi_0(\xi) f(\xi) \chi_\kappa(\xi) d\xi \approx 2 \int_{r_\Sigma}^{\xi_{\text{max}}} \chi_0(\xi) f(\xi) \chi_\kappa(\xi) d\xi,
\]

where Eq. (15) was used in obtaining the third line. The extra factor, $\sqrt{2/ [1 + R(t)]}$, attached to $a_\ell(t)$ in Eq. (14a) serves to provide the conclusion of Eq. (22): $f \langle |\psi(t)|^2 dr = ||\Psi(t)||^2 = 1$. In the formulation without this factor, we arrive at another normalization condition, $f \langle |\psi(t)|^2 dr = \sum_{\ell} \sum_{\kappa} |a_{\kappa \ell}(t)|^2 + |1 + R(t)||a_\ell(t)|^2/2 + \sum_{\kappa} |b_{\kappa \ell}(t)|^2 = 1 \neq ||\Psi(t)||^2$, which indicates the decrease of $||\Psi(t)||^2$ as $R(t)$ grows. To fulfill the normalization condition, a complex term, $-i R(t)/[2(1 + R(t))]|a_\ell(t)|$, shows up in an equation corresponding to Eq. (21) on its right hand side. Consequently, the working equations turn out to be non-Hermitian in matrix form if the factor $\sqrt{2/ [1 + R(t)]}$ is not included. The time propagation can still be implemented by the Arnoldi algorithm 16, 47 or the Runge-Kutta method 32, for instance, but not by the simple Lanczos algorithm. The factor, $\sqrt{2/ [1 + R(t)]}$, is hence better attached to $a_\ell(t)$ in Eq. (14a) to prevent such unnecessary complications.

IV. STIFFNESS-FREE FEDVR-BASED LANCASTER ALGORITHM

As seen from the formulation in Sec. I and the explicit form of the working equations of Sec. II, ETS is a generalization of GTS aiming at the reduction of stiffness; non-time scaling in the inner region avoids shrinking of most bound states, and hence, differently from GTS, allows the use of less dense basis functions around the origin. In the treatment of strong-field ionization at mid-IR wavelengths, corresponding to the tunneling regime, the equations of motion in ETS inevitably still become stiff as many angular momentum states are involved. To resolve this remaining problem, this section provides a detailed analysis to identify the origin of the stiffness, and proposes a procedure for its removal. Note that this section holds some independence from the other sections. The discussion in the following is not only applicable to ETS implementation, but also to a wider class of equations of motion appearing in atomic and molecular physics.
A. Error and stiffness analysis on Lanczos algorithm

This subsection focuses on stiffness (see, e.g., Ref. [33]), i.e., the degree of difficulty in a numerical treatment of the working equations, to see how it poses problems, and to identify its origin for seeking the resolution. Toward this end, the discussion commences with an error analysis on the short-time iterative Lanczos algorithm. For a given vector, $\Psi(t)$, the evolution in a small time step $\Delta t$ is given by

$$\Psi(t + \Delta t) = \sum_{k=0}^{K-1} \frac{1}{k!} [-iH(t)\Delta t]^k \Psi(t),$$

(23)

with error

$$\|\Psi_{\text{exact}}(t + \Delta t) - \Psi(t + \Delta t)\|^2 \approx \left| \frac{\beta_1(t)\beta_2(t)\cdots\beta_{K-1}(t)}{(K-1)!} (\Delta t)^{K-1} \right|^2 < \epsilon,$$

(24)

FIG. 3. (a) Plot of $\max \{e_{nl}\} / [\ell(\ell + 1)]$ as a function of $\ell$ (in this case, $\max \{e_{nl}\} = e_{89,\ell}$). Every ten data is shown by open circles. The solid gray line indicates $(2\xi_1^2)^{-1} = 137.28$. (b) Eigenvectors of $h_\ell$, satisfying $e_{\ell\ell} > E_{\text{cut}} = 900 \ (\ell = 0, \cdots, 200(= \ell_{\text{max}}))$. The plot is made in terms of the FEDVR expansion, $\varphi_{\ell\ell}(\xi) \equiv \sum_{\kappa=1}^N \tilde{\Psi}_{\ell\kappa}(\xi) \chi_{\kappa}(\xi)$. Both (a) and (b) are obtained under the same numerical condition: $N = 89 \ (N_{\text{dvr}} = 10$ and $N_{\text{fe}} = 10)$ and $\Delta \xi = 1.5$.

FIG. 4. (Color online) (a) Contour plot of $K_{\text{max}}$ as a function of $\ell_{\text{max}}$ and $\Delta t$. The color code and solid black lines represent the contour based on Eq. (24) with $\epsilon = 10^{-15}$. The blank area indicates $K_{\text{max}} > 1000$. This estimation is obtained for field-free Hamiltonian, $H(0)$ under the numerical condition: $\sum_{\ell} = \xi_{\text{max}} = 60, N_{\text{dvr}} = 10, N_{\text{fe}} = 40, N_{\text{fe}} = 0$, and $\Delta \xi = 1.5$. The dashed (red) lines show the contour based on Eq. (27). (b) The same as (a) except that the stiffness is removed by setting $N_{\text{fe}} = 10$ and $E_{\text{cut}} = 900$. See the main text for details.
where \( \{\beta_i(t)\}_{i=1}^{K-1} \) is a set of subdiagonal elements in a reduced form of \( \mathbf{H}(t) \) on the \( K \)-dimensional Krylov subspace \( 37 \), and the last inequality requires the error to be less than, say, \( \epsilon = 10^{-15} \). Setting \( \Delta t \) much smaller than the characteristic time scale of external fields, Eq. \((24)\) serves as a criterion at every time step to determine the smallest integer for \( K \). If the time propagation starts with the ground state, the product \( \beta_1(t)\beta_2(t) \cdots \beta_{K-1}(t) \) is zero at \( t = 0 \), but in due course reaches its maximum, when the Krylov subspace acquires the largest dimension, \( K_{\text{max}} \); the computation faces difficulty if \( K_{\text{max}} \) is too large. Let \( B \) denote the maximum of \( \sum_{\ell} \beta_{\ell}(t) \) during time evolution. Note that, although field-free atomic states, the energy eigenvalues of which are well above the energy range of physical interest, could participate in construction of the Krylov subspace, the population of such high-energy states are virtual excitations which happen more likely for larger \( \Delta t \). Supposing the worst case such that every energy eigenstate is accessed in the virtual excitations, let the analysis in the following be based upon an assumption, \( B = \max\{E_{n\ell}\}_{n\ell} \), i.e., \( B \) reaches the maximum eigenvalue of \( \mathbf{H}(0) \). In this assumption, the working equations become stiffer as \( \max\{E_{n\ell}\}_{n\ell} \) increases.

Meanwhile, looking at the working equations \((10)\), \((17)\), and \((24)\), we notice that \( V_{\ell}(\xi_k) \) in Eq. \((16)\) takes very large values in the vicinity of the nucleus [see Eq. \((3)\)] and is the decisive factor of \( \max\{E_{n\ell}\}_{n\ell} \); hence, Eq. \((16)\) is responsible for the stiffness. To discuss more quantitatively, we define a set of field-free Hamiltonian matrices around the nucleus,

\[
\begin{align*}
(\mathbf{h}_\ell)_{\kappa\kappa'} & = \frac{1}{2} \int_0^{\xi_{\text{max}}} d\xi \frac{d\chi_\kappa(\xi)}{d\xi} \frac{d\chi_{\kappa'}(\xi)}{d\xi} d\xi + \delta_{\kappa\kappa'} V_{\ell}(\xi_k), \\
(\kappa, \kappa') & = (1, \cdots, N, \ell = 0, \cdots, \ell_{\text{max}}), \quad (25)
\end{align*}
\]

where \( \tilde{N} = (N_{\text{dvr}} - 1)\tilde{N}_{\text{fe}} - 1 \) with \( \tilde{N}_{\text{fe}} \leq \tilde{N} \), and note the boundary condition, \( \chi_\kappa(\xi(\geq \xi_N)) = 0 \) for \( \kappa = 1, \cdots, \tilde{N} \). Diagonalizing these small matrices and obtaining \( \mathbf{u}_0^T \mathbf{h}_\ell \mathbf{u}_0 = \text{diag}(\epsilon_1, \epsilon_2, \cdots, \epsilon_{\tilde{N}}) \), the eigenvalues for the first few integers of \( n \) represent the bound-state energies, \( \epsilon_{n\ell} \approx -[2(n + \ell + 1)]^{-1} \). The rest of the eigenvalues are positive and could, in particular for large \( \ell \), be too large to be of importance in the physical process of interest. We then suppose

\[
\max\{E_{n\ell}\}_{n\ell} \approx \max\{\epsilon_{n\ell}\}_{n\ell} \approx \frac{\ell_{\text{max}}(\ell_{\text{max}} + 1)}{2\xi_{\text{max}}^2}, \quad (26)
\]

i.e., \( \max\{\epsilon_{n\ell}\} \) is not sensitive to \( \tilde{N} \) and almost determined by the centrifugal part of \( V_{\text{max}}(\xi_1) \) in Eq. \((3)\).

This conjecture is verified by numerical examples: Setting \( \tilde{N} = 89 \) (\( N_{\text{dvr}} = 10 \) and \( \tilde{N}_{\text{fe}} = 10 \) and \( \Delta \xi = 1.5 \)), the greatest eigenvalue for each \( \ell \), \( \max\{\epsilon_{n\ell}\} \approx \epsilon_{n\ell}(\ell + 1) \), is, after divided by \( \ell(\ell + 1) \), plotted in Fig. \((3)\) (a). This kind of plot is, for fixed \( N_{\text{dvr}} = 10 \) and \( \Delta \xi = 1.5 \), insensitive to the change of \( \tilde{N}_{\text{fe}} \) (not shown in the figure), and, in every case, approaching \( (2\xi_{\text{max}})^{-1} = 137.25 \) at large \( \ell \). Hence, using Eq. \((26)\) in Eq. \((24)\) with assuming \( B = \max\{E_{n\ell}\}_{n\ell} \)

gives

\[
\frac{K_{\text{max}} - 1}{\Delta t} > \epsilon : [2\pi(K_{\text{max}} - 1)\epsilon]^{-1/2(2K_{\text{max}} - 1)} \times \ell_{\text{max}}(\ell_{\text{max}} + 1) \frac{2\xi_{\text{max}}^2}{2\xi_{\text{max}}^2}, \quad (27)
\]

where \( (K_{\text{max}} - 1)! > \sqrt{2\pi(K_{\text{max}} - 1)\epsilon} \cdot [(K_{\text{max}} - 1)/\epsilon]^{K_{\text{max}} - 1} \) is used (Stirling’s formula; see, e.g., Ref. \((19)\)). Noting that the numerical cost is proportional to \( (K_{\text{max}} - 1)/\Delta t \), and also \( [2\pi(K_{\text{max}} - 1)\epsilon]^{-1/2(2K_{\text{max}} - 1)} \rightarrow 1 + 0 \) as \( K_{\text{max}} \rightarrow \infty \), Eq. \((27)\) indicates the growing numerical efficiency by setting \( \Delta t \) as large as possible (as long as it is still much smaller than the characteristic time scale of external fields). Equation \((27)\) at the same time manifests the overwhelming stiffness for large \( \ell_{\text{max}} \) and/or small \( \xi_1 \). Although the assumption \( B = \max\{E_{n\ell}\}_{n\ell} \) is so naïve that Eq. \((27)\) may overestimate \( K_{\text{max}} \), a more rigorous theoretical analysis is beyond the scope of this paper. Let us verify Eq. \((27)\) numerically instead; Fig. \((4)\) (a) shows \( K_{\text{max}} \) as a function of \( \ell_{\text{max}} \) and \( \Delta t \). The color code and solid black lines represent the contour based on Eq. \((24)\) with \( \epsilon = 10^{-15} \), \( K_{\text{max}} \) is calculated for each pair of \( \ell \) and \( \Delta t \), with \( \max\{K_i\}_{i=1}^{100} \), where \( K_i \) is the smallest integer satisfying Eq. \((24)\) in obtaining \( \Psi_i(\Delta t) \) from \( \Psi_i(0) \) which is every time \( \ell = 1, \cdots, 100 \) constructed by random-number elements and normalized. The calculation is based on a field-free non-time-scaled Hamiltonian matrix \( \mathbf{H}(0) \) with the following parameters: \( r_{\Sigma} = \xi_{\text{max}} = 60 \), \( N_{\text{dvr}} = 10 \), \( \tilde{N}_{\text{fe}} = 40 \), \( \tilde{N}_{\text{fe}} = 0 \), and \( \Delta \xi = 1.5 \). The same calculation with a Hamiltonian matrix including light-atom interaction term (with fixed light intensity \( I = 10^{14} \text{ W/cm}^2 \)) in the length gauge causes no visible change. The dashed (red) lines in Fig. \((4)\) (a), representing the contour based on Eq. \((27)\), show obvious overestimation but capture the gross feature of the landscape, verifying Eq. \((27)\) despite its simplicity.

Equation \((27)\) provides explicitly the following insights: Achieving high accuracy in the description of electronic structure and tunneling dynamics requires many FEDVR functions in the fixed interval, \([0, \xi_{\text{max}}]\), as well as many angular momentum states. With improved accuracy, however, \( \ell_{\text{max}}(\ell_{\text{max}} + 1)/2\xi_{\text{max}}^2 \) rapidly increases, requiring larger \( K_{\text{max}} \), which leads to a growing stiffness in the working equations. Note that the stiffness problem is not peculiar in atomic systems with the FEDVR-based Lanczos algorithm, but rather inherently appears in a variety of differential equations in numerical treatments.

**B. Stiffness removal**

As shown in Sec. \((14)\), the stiffness is mainly caused by high-angular momentum states, and due to the large value of the centrifugal potential around the vicinity of the nucleus. This conclusion suggests a clear strategy to remove the stiffness; setting a cutoff energy \( E_{\text{cut}} \) well above the range of physical interest, the eigenvectors of
matrices by let us approximately reconstruct \( h \). Defining a set of energy states localize only around the nucleus, hence they are formed by \( h \). Figure 3 (b) depicts eigenvectors of \( e \), which are above the cutoff, \( h \). Before the stiffness removal, each angular momentum diagonal block \( (\ell = \ell') \) has block-diagonal structure with respect to \( \xi \), and each angular momentum subdiagonal block \( (\ell = |\ell' + 1|) \) is just diagonal (because of the length gauge). After the stiffness removal, \( h_{\ell} \) is replaced by \( h_{\ell} \) for \( \ell \geq 1 \) [see Eq. (29)], and every light-atom interaction block \( F(t)w_{\ell\ell'} \) is replaced by \( F(t)\tilde{w}_{\ell\ell'} \) [see Eq. (31)]. Consequently, the top-left corners of angular momentum diagonal and subdiagonal blocks become full. This figure is consistent with Fig. 2 in color code; the shaded, non-shaded (blue), and filled (red) parts, respectively, indicate the matrix elements formed by \( \{\chi_\sigma(\xi)\}_{\kappa} \), \( \{\chi_\bar{\sigma}(\xi)\} \), and \( \{\chi_\kappa(\xi)\}_{\kappa} \).

The light-atom interaction should accordingly be modified around the nucleus; canceling out the TD field factor in the light-atom interaction operator, and defining time-independent matrix by [see Eq. (4)]

\[
(w_{\ell\ell'})_{\kappa\kappa'} \equiv \frac{1}{G(t)} \int_0^{\ell_{\text{max}}} \chi_{\kappa}(\xi)W_{\ell\ell'}(t)\chi_{\kappa'}(\xi)d\xi,
\]

(30)

with \( G(t) \) denoting \( F(t) (A(t)) \) in the length (velocity) gauge, let Eq. (30) then be approximated as

\[
w_{\ell\ell'} \approx \tilde{w}_{\ell\ell'} \equiv \tilde{w}_{\ell\ell'}.\]

The set of working equations is now approximated as \( i\dot{\Psi}(t) = H(t)\Psi(t) \), where \( H(t) \) is constructed using \( h_{\ell} \) and \( G(t)\tilde{w}_{\ell\ell'} \) (see Fig. 5) and is hence expected to be

\[
h_{\ell} = u_{\ell}u_{\ell}^T h_{\ell} u_{\ell} u_{\ell}^T \approx \tilde{u}_{\ell}u_{\ell}^T h_{\ell} u_{\ell} u_{\ell}^T \equiv \tilde{h}_{\ell}.\]

(29)
stiffness free and to lead to a reduction in $K_{\text{max}}$. Look at the contour plot of $K_{\text{max}}$ in Fig. 4(b), which is computed in the same numerical condition as in Fig. 4(a) after the application of stiffness removal procedure with $N_{\text{fe}} = 10$ and $E_{\text{cut}} = 900$. Figure 4(b) confirms our expectation.

Note that $\mathbf{h}_r$ is structured block diagonal, and $\mathbf{w}_r$ is diagonal (block diagonal) in the length (velocity) gauge, whereas both $\mathbf{h}_r$ and $\mathbf{w}_{\mathbf{r}'r'}$ are full. Hence, the stiffness removal partly destroys the sparseness of $\mathbf{H}(t)$, but $\mathbf{H}(t)$ is still largely sparse and can be efficiently handled in Harwell-Boeing format (see, e.g., Ref. [32]). In most cases, $N_{\text{fe}}(\leq N_{\text{fe}})$ will be set around 10 to safely remove the stiffness. Setting $N_{\text{fe}}$ smaller makes $\mathbf{H}(t)$ sparser. However, we always need to check before starting the time propagation that the removed high-energy states do not localize near the right edge of this small interval (around $\xi = 15$ in the case of Fig. 3(b)). If not, it is safe, but if so, $N_{\text{fe}}$ and/or $E_{\text{cut}}$ must be set larger.

The stiffness removal procedure given above is not easily applicable to GTS because in GTS every element of the Hamiltonian matrix depends on time. Also note that the stiffness removal itself is not a new concept. One can find a discussion for the (without-space-partition) TD B-spline $R$-matrix approach in Ref. [50]. Similar procedures have thereafter been proposed by several researchers. Reference [51], for instance, suggests a procedure for TD many-electron calculations based on the FEDVR functions. The stiffness is removed, however, over the whole spatial simulation volume. The total Hamiltonian matrix becomes completely full, spoiling the benefit of FEDVR functions. One can see another procedure more similar to ours in Ref. [52], where the authors investigate the photoionization of Ne$^+$ by means of RMT [40–42]. Based upon the space-partition concept, the radial wave functions are expressed in terms of the $R$-matrix basis functions and grids, respectively, inside and outside a spherical sphere the radius of which is $R_\Sigma = 15$. The working equations are free from stiffness by setting $E_{\text{cut}} = 1345$ and composing the $R$-matrix basis functions of field-free eigenstates.

V. NUMERICAL DEMONSTRATION

By solving the ETS working equations [16], [17], and [21] with the stiffness-free procedure discussed in Sec. IV.B, let atomic hydrogen, prepared in the ground state at $t = 0$, time evolve under an $N$-cycle mid-IR ($\lambda = 3 \mu\text{m} = 5.669 \times 10^2$) pulse the envelop of which is defined by

$$f_N(t) = \begin{cases} \sin^2 \left( \frac{\pi t}{T} \right) & (0 \leq t \leq T) \\ 0 & (T < t) \end{cases}$$

where $T = 2\pi N/\omega = 413.7 \times N$. In due care of Eq. (32), the time-scaling factor is now defined by

$$R(t) = \begin{cases} \frac{R_{\infty}}{2T} \left( t^2 + \frac{T^2}{2\pi^2} \left[ \cos \left( \frac{2\pi t}{T} \right) - 1 \right] \right) + 1 & (0 \leq t \leq T) \\ R_{\infty}(t - T) - \frac{TR_{\infty}}{2} + 1 & (T < t) \end{cases}$$

which gives non-vanishing $\ddot{\hat{R}}(t)$ only during the presence of the light field: $\ddot{\hat{R}}(t) = (2R_{\infty}/T)f_N(t)$ for $0 \leq t \leq T$, and $\ddot{\hat{R}}(t) = 0$ for $t > T$. The original idea of time scaling appeared in search of suitable adiabatic parameters for describing atom-diatom collisions [24]. In our case, supposing some adiabatic action of the mid-IR pulse on the electron, the time-scaling factor will be better specified by the carrier envelop function. Note that Refs. [26–32] propose another form, $R(t) = [1 + (R_{\infty}(t - t_0))^n]^{1/n}$ ($n = 2, 3,$ or 4), for investigating ionization by high-energy photoabsorption (in the extreme ultraviolet region) and fast-electron impact (in the keV region); Ref. [31] also reports the insensitivity of the calculation to the starting time $t_0$. For mid-IR laser pulses, however, this definition is not well suited because $\ddot{\hat{R}}(t)$ changes so much faster than $f_N(t)$ that it can be a source of numerical instability; the calculation is very sensitive to $n$ and $t_0$. The choice of $R(t)$ in Eq. (33) gives good properties in the present case. Studies of process-dependent optimal forms of $R(t)$ could be interesting in the future.

A. Interaction with a three-cycle pulse

Let us consider a three-cycle ($N = 3$) pulse the vector potential of which is given by

$$A(t) = \frac{F_0}{\omega} f_3(t) \sin \omega t,$$

where $F_0 = \sqrt{T} = 5.338 \times 10^{-2}$ with the intensity $I = 10^{14} \text{W/cm}^2 = 2.849 \times 10^{-3}$. Figure 11 depicts the profile of the electric field in the top-right corner. Figure 6(a) displays the radial electron densities of atomic hydrogen,

$$\rho(r, t) = \sum_{\ell = 0}^{\ell_{\text{max}}} |\psi_{\ell}(\xi(r, t), t)|^2,$$

computed in the velocity gauge by the ETS method and the usual non-time-scaled method. The snapshot at each end of cycles ($t = (2\pi/\omega) \times n, (n = 1, 2,$ and 3) exhibits the extension of the spatial radius in the ETS calculation, while the radius in the non-time scaled calculation keeps 3000, constant. The inset of Fig. 6(a) displays the smooth continuity of the radial density across $r_\Sigma$, showing the stability of the ETS implementation. As a realistic observable, Fig. 6(b) displays the HHG spectrum computed as a normed square of the Fourier transformation of the
parameters except the time step: $\Delta t$[\text{c.c.}]) in three ETS calculations using the same numerical scheme. FIG. 7. (Color online) Adapted dimension of the Krylov subspace, $K$, as a function of time [in units of optical cycle (o.c.)] in three ETS calculations using the same numerical parameters except the time step: $\Delta t = 0.01$, 0.05, and 0.1. The value of $K$ is determined based on Eq. (23) to ensure the accuracy of the results: $\epsilon = 10^{-15}$ at every time step. The calculations are carried out in the velocity gauge for atomic hydrogen interacting with a three-cycle mid-IR laser pulse ($\lambda = 3$ $\mu$m, $I = 10^{14}$ W/cm$^2$, and $N = 3$). See the main text for details.

The HHG spectrum consists of three plateaus: (1) $0 < \Omega/U_p < 1$, (2) $0.2 < \Omega/U_p < 1.8$, and (3) $1.9 < \Omega/U_p < 3.2$. Based upon the simple man’s model (a simple classical simulation) [53], the electron trajectory characterized by the ionization and return times, $t_i^{(n)}$ and $t_r^{(n)}$, respectively, has the main responsibility to the nth plateau ($n = 1, 2, 3$): $2.03 < t_i^{(1)} < 2.20$ and $2.20 < t_i^{(1)} < 3.00$, $1.52 < t_i^{(2)} < 1.74$ and $1.74 < t_i^{(2)} < 2.77$, and $1.00 < t_i^{(3)} < 1.26$ and $1.26 < t_i^{(3)} < 1.98$ (in units of o.c.); also see the electric-field profile in the top-right corner of Fig. 5. Such a clear time-to-energy mapping reflects the validity of the semiclassical picture of electron dynamics under the mid-IR lasers, and becomes less clear if $\lambda = 3$ $\mu$m is replaced by $\lambda = 0.8$ $\mu$m (not shown). A similar triple plateau in HHG spectra was recently reported for a different three-cycle pulse with $\lambda = 1.6$ $\mu$m in Ref. [54], where more detailed discussions are given based on the wavelet analysis with the help of the quantum-orbit model.

In the ETS calculation for obtaining the results in Fig. 3 the wave function is parametrized as follows: $\xi_{\text{max}} = 200$, $\gamma_{\text{c}} = 30$, $\xi_{\text{max}} = 450$, $N_{\text{dvr}} = 10$, $N_{\text{ic}} = 20$, $N_{\text{ef}} = 280$, $\Delta \xi = 1.5$, and $R_{\infty} = 0.01$. The non-time scaled calculation uses the same parameters except $\xi_{\text{max}} = 5000$, $N_{\text{ic}} = 1980$, and $R_{\infty} = 0$. The parameters for the stiffness-free time propagation are common in both calculations: $\Delta t = 0.05$, $\epsilon = 10^{-15}$, $E_{\text{cut}} = 900$, and $N_{\text{ic}} = 10$ (i.e., $N = 89$, which is the same as in Figs. 4b and 5b). The dimension of the Krylov subspace is adapted at each time step based on Eq. (44). Figure 4 shows $K$ as a function of time in three ETS calculations with different time steps: $\Delta t = 0.01$, 0.05, and 0.1 (the rest of parameters are the same). In every case, $K$ increases with the start of the laser pulse, but keeps smaller than the estimated maximum in Fig. 4b. These three calculations exemplify the decrease of $(K_{\text{max}} - 1)/\Delta t$ as $\Delta t$ increases: $(9 - 1)/0.01 = 800$, $(33 - 1)/0.05 = 640$, and $(64 - 1)/0.1 = 630$, i.e., decreasing numerical cost

FIG. 6. (Color online) Radial electron densities, $\rho(r, t)$ [Eq. (35)], at first, second, and third optical cycles (o.c.), i.e., at $t = (2\pi/\omega) \times n$ ($n = 1, 2, 3$). The inset shows the radial density around the origin; shaded area indicates the radial interval of the inner region ($0 \leq \xi(= r) \leq \gamma_{\text{c}} = 30$). (b) HHG intensity, $S(\Omega)$, computed as a normed square of the Fourier transform of dipole acceleration along the $z$ axis (polarization of the incoming pulse),

$$S(\Omega) = \left| \int_0^T (\Psi(t) - F(t)) \frac{1}{\omega} \left( \frac{\partial}{\partial \omega} F(t) \right) e^{i\Omega t} dt \right|^2.$$ (36)
as a function of $\Delta t$ [see Eq. (27)]. That is, setting $\Delta t$ larger improves the numerical efficiency as long as it is much smaller than the characteristic time scale of external fields; in this case, $\Delta t \ll 2\pi/\omega = 413.7$ should be satisfied. Also note that, without the stiffness removal, the time propagation is virtually infeasible because Eq. (24) requires $K$ to be much greater than 1000 (the blank area in Fig. 3(a)).

Due to the presence of $R(t)$, the numerical implementation of the ETS working equations (16), (17), and (21) consumes an extra CPU time, which is absent in the usual non-time scaled calculation, for updating several matrix elements in the outer region [the block-diagonal part, shown by dashed (blue) lines in Fig. 5] at each of angular momentum blocks along $\ell = \ell'$. However, this operation does not cause a major problem. In the above calculations, for instance, the time propagation by the non-time scaled method takes about six times more CPU time than by the ETS method. On the other hand, the number of nonzero elements of $\mathbf{H}(t)$ in the non-time scaled calculation is 123135659, which is 5.6 times larger than in the ETS calculation, 21987359. The efficiency by ETS is almost proportional to the reduction of nonzero matrix elements. However, we also need to mention a weak point of ETS: the ETS calculation is sensitive to the numerical condition, and finding a proper set of parameters, especially a proper value of $R_\infty$ [see Eq. (33)], is not straightforward. Assuming $r \sim kt$, i.e., a classical relation between the radial position and momentum of the photoelectron at large times, Eq. (33) then indicates $k \sim R_\infty\xi$ [24–32]. Thus one may estimate $R_\infty = \sqrt{2 \cdot 10^6/\xi_{\max}} = 0.017$, where $U_p = F_0^2/(4\omega^2) = 3.088$, and $10U_p$ is the maximum kinetic energy the photoelectron acquires by recollisions with the parent ion [53, 54]. The replacement of $R_\infty = 0.01$ by 0.017 in the above ETS calculation, however, gives some numerical instability; $R_\infty = 0.017$ results in too fast growth of the $r$ coordinate, and hence requires a larger number of FEDVR functions for the accurate description of recollisions; this is numerically unfavorable. The numerical stability also depends on the value of $r_C$. Although Figs. 1(b1–3) show smooth continuity of the Coulomb potential function and $\psi_{1s}(r,t)$ across $r_C = 30$ and seemingly imply their stable numerical treatment, the accurate description of electron recollisions during the growth of the $r$ coordinate is not so simple because of the involvement of many excited states.

Finally note that, just for computing HHG, ETS is not so useful; one can use a relatively small simulation volume in the non-time scaled calculation by employing a CAP or ECS, because only the electron dynamics around the nucleus is important to the dipole acceleration. The computation finishes with the completion of the pulse and does not require a long-time evolution any further. The true potential of ETS is the capability of long-time propagation without losing the norm of wave function as illustrated in the next subsection.

### B. Interaction with a half-cycle pulse

We now consider a half-cycle ($N = 1/2$) pulse defined by the electric field,

$$F(t) = -F_0 f_{1/2}(t) \sin \omega t,$$

and track the long-time evolution of the tunnel-ejected electron after the completion of the pulse. Note that Eq. (37) does not comply with the condition for realistic light fields, $\int_0^T F(t) dt = 0$ (see, e.g., Refs. [53, 58]), and gives a non-vanishing vector potential at $t = 0$ and $t > T$. Hence this artificial pulse necessitates employing the length gauge. Figure 8 displays the time evolution of the radial electron density [Eq. (35)]. At $t = 0.5$ o.c., i.e.,
FIG. 9. (a1–3) Real parts of $\psi(\xi, t)$ (thick dark-gray lines) and $\phi(\xi, t)/\sqrt{R(t)}$ (thin black lines) for $\ell = 1, 50, \text{ and } 100$ at 0.5th optical cycles (o.c.); (b1–3) the same but at 20th o.c. The shaded area in each panel indicates the radial interval of the inner region ($0 \leq \xi = r \leq r_\Sigma = 30$). For better visibility and comparison, each pair of curves is plotted after multiplication by the inverse of max $|\text{Re}[\psi_\ell(r, t)]|$ as indicated in each panel. The non-smooth appearance of $\text{Re}[\psi_\ell(r, t)]$ around $r_\Sigma$ in (b1) is just due to the abrupt change of the scaling in the $r$ coordinate. The results in this figure and Fig. 8 are obtained from the same ETS calculation in the length gauge for atomic hydrogen interacting with a half-cycle mid-IR laser pulse ($\lambda = 3 \mu\text{m}, I = 10^{14} \text{ W/cm}^2, \text{ and } N = 1/2$). See the main text for details.
just after the completion of the pulse, the electron wave packet occupies a volume of radius about 1000. Due to its broad energy spectrum, the wave packet then spreads over a vast expanse of volume and reaches 55000 at \( t = 20 \) o.c. There is neither difficulty nor instability to continue the evolution as long time as one wishes. The insert of Fig. 5 monitors the smooth radial density across \( r_{\Sigma} = 30 \). The numerical condition in this ETS calculation is the same as the one shown in Sec. IV A except \( \xi_{\text{max}} = 300 \), \( N_{\text{fe}} = 180 \), \( R_{\infty} = 0.025 \), and the employment of the length gauge.

Based on the classical picture, one may think of \( R_{\infty} = \sqrt{2U_p/\xi_{\text{max}}} = 0.0083 \), because now there is no recollision. This estimation is, however, too little to take into account the broad energy spectrum of the wave packet, and it is safer to set \( R_{\infty} \) a few times larger. In contrast to the three-cycle pulse in Sec. IV A, the calculation is not sensitive to \( R_{\infty} \) and other parameters; the same converged result as in Fig. 5 is also obtained more easily by setting \( R_{\infty} \) larger than 0.025 and employing smaller values for \( \xi_{\text{max}} \), \( N_{\text{fe}} \), and \( N_{\text{fe}} \). Such an insensitivity to numerical condition is due to the absence of recollision in the half-cycle pulse. That is, for the analysis of long-time evolution of electron wave packets in circularly or near-circularly polarized mid-IR pulses, ETS will show its true potential without concerns of the numerical sensitivity to the parameters.

Finally analysis of the radial function for each \( \ell \) will be worthwhile for realizing how the ETS method enables keeping numerical stability for very long-time evolution. Figure 6 shows the real parts of \( \psi_{\ell}(r, \xi, t) \) and \( \phi_{\ell}(\xi, t)/\sqrt{R(t)} \) (multiplied by a constant; see the caption) for \( \ell = 1, 50 \), and 100. At \( t = 0.5 \) o.c., although the spatial volume has not much extended yet, the oscillation of \( \text{Re}[\psi_{\ell}(r, \xi, t)] \) is already so fast that it is almost invisible. After a long-time evolution, the situation becomes worse because \( \psi_{\ell}(r, \xi, t) \) spreads over a vast interval in the \( r \) coordinate with increasing its phase gradient outward from the center of the wave packet. Direct numerical treatment of \( \psi_{\ell}(r, t) \) by the usual non-time scaled method is hence virtually infeasible. On the other hand, because the phase transformation in Eq. (9) cancels out the growing phase gradient (see the discussion in Ref. [20]), \( \text{Re}[\phi_{\ell}(\xi, t)/\sqrt{R(t)}] \) exhibits not so fast oscillation in the \( \xi \) coordinate even after a very long-time evolution. Figure 6 shows the increasing difficulty of direct numerical treatment of \( \psi_{\ell}(r, t) \) as it spreads without the help of the time scaling and the phase transformation.

VI. CONCLUSION AND OUTLOOK

Aiming at an efficient numerical treatment of tunneling ionization of atoms and molecules by mid-IR lasers, the ETS theory is formulated as a generalization of GTS. The working equations for numerical implementation are derived in terms of FEDVR basis functions. The key idea of ETS is to divide the spatial volume into two regions: a small spherical sphere around the nucleus and its outside, and then to carry out the time scaling only to the radial coordinates outside. As a result, the continuum part of the photoelectron wave packet is time scaled in the outer region and prevented from reflection. On the other hand, the bound-state part in the inner region is not time scaled and does not shrink toward the origin. Hence, ETS is less stiff than GTS. Furthermore, the stiffness-free FEDVR-based Lanczos algorithm is established to completely eliminate any stiffness for the treatment of long-wavelength lasers. The test calculations for atomic hydrogen interacting with linearly polarized mid-IR pulses demonstrate the capability of ETS and the stiffness-free time propagator. The method shows its true potential for the detailed analysis of wave-packet dynamics in non-recollision situations.

The ETS method and the stiffness-free time propagator can be flexibly used in several coordinate systems, e.g., in hyperspherical coordinates, as mentioned in Ref. [32], to treat photoionization of atomic helium. Application to RMT [40–42] may also be possible, but generalization to the TD-(RAS/GAS)CI method is straightforward, enabling the extension to many-electron systems. Setting aside the many-electron problem, tunneling ionization of atoms and molecules by strong mid-IR lasers with arbitrary polarizations remains unexplored even within the SAE approximation. Toward this direction of research, in particular for circular or near-circular polarization, the potential of ETS is very promising.

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Appendix: Light-atom interaction operator in the velocity gauge

The FEDVR-based working equations are derived in the length gauge in Sec. III B. The following is a supplementary list of instructions needed to rewrite the light-atom interaction terms in Eqs. (16), (17), and (21) in the velocity gauge.
In Eq. (16),

\[
W_{\ell t}^{1}(\xi_{t}, t)a_{\xi t}(t)
\]

\[
\rightarrow \int_{-\infty}^{r_{\Sigma}} \chi_{\xi}(\xi)W_{\ell t}^{1}(\xi, \partial_{\xi}, t)\psi_{\ell}(\xi, t)d\xi
\]

\[
= -\frac{i}{2}A(\xi)g_{\ell t}\left\{ \sum_{k} a_{\xi t}(t) \int_{0}^{\xi_{\max}} \left[ \chi_{\xi}(\xi) \frac{d\chi_{\xi}(\xi)}{d\xi} - \frac{d\chi_{\xi}(\xi)}{d\xi} \chi_{\xi}(\xi) \right] d\xi
\]

\[
+ \sqrt{\frac{2}{1 + R(t)}} a_{\xi t}(t) \int_{0}^{\xi_{\max}} \left[ \chi_{\xi}(\xi) \frac{d\chi_{\xi}(\xi)}{d\xi} - \frac{d\chi_{\xi}(\xi)}{d\xi} \chi_{\xi}(\xi) \right] d\xi + \frac{\ell(\ell + 1) - \ell(\ell + 1)}{r_{\Sigma}} a_{\xi t}(t) \right\}.
\]  

(A.1)

In Eq. (17),

\[
W_{\ell t}^{1}[r_{\Sigma} + R(t)(\xi_{t} - r_{\Sigma}), t]b_{\xi t}(t)
\]

\[
\rightarrow \int_{r_{\Sigma} + 0}^{\xi_{\max}} \chi_{\xi}(\xi)\left\{ W_{\ell t}^{1}[r_{\Sigma} + R(t)(\xi - r_{\Sigma}), \partial_{R(t)}, t] + g_{\ell t}A(t)R(t)(\xi - r_{\Sigma}) \right\} \phi_{t}(\xi, t)d\xi
\]

\[
= -\frac{i}{2}A(\xi)g_{\ell t}\left\{ \sum_{k} b_{\xi t}(t) \int_{0}^{\xi_{\max}} \left[ \chi_{\xi}(\xi) \frac{d\chi_{\xi}(\xi)}{d\xi} - \frac{d\chi_{\xi}(\xi)}{d\xi} \chi_{\xi}(\xi) \right] d\xi
\]

\[
+ \sqrt{\frac{2}{1 + R(t)}} a_{\xi t}(t) \int_{0}^{\xi_{\max}} \left[ \chi_{\xi}(\xi) \frac{d\chi_{\xi}(\xi)}{d\xi} - \frac{d\chi_{\xi}(\xi)}{d\xi} \chi_{\xi}(\xi) \right] d\xi + \frac{\ell(\ell + 1) - \ell(\ell + 1)}{r_{\Sigma}} b_{\xi t}(t) \right\}
\]  

(A.2)

In Eq. (21),

\[
W_{\ell t}^{1}[r_{\Sigma}, t]a_{\ell t}(t)
\]

\[
\rightarrow \sqrt{\frac{2}{1 + R(t)}} \left[ \int_{-\infty}^{r_{\Sigma}} \chi_{b}(\xi)W_{\ell t}^{1}(\xi, \partial_{\xi}, t)\psi_{\ell}(\xi, t)d\xi
\]

\[
+ \sqrt{R(t)} \int_{r_{\Sigma} + 0}^{\xi_{\max}} \chi_{b}(\xi)\left\{ W_{\ell t}^{1}[r_{\Sigma} + R(t)(\xi - r_{\Sigma}), \partial_{R(t)}(\xi - r_{\Sigma})] + g_{\ell t}A(t)R(t)(\xi - r_{\Sigma}) \right\} \phi_{t}(\xi, t)d\xi
\]

\[
= -\frac{i}{2}A(\xi)g_{\ell t}\left\{ \sqrt{\frac{2}{1 + R(t)}} \sum_{k} a_{\ell t}(t) \int_{0}^{\xi_{\max}} \left[ \chi_{b}(\xi) \frac{d\chi_{b}(\xi)}{d\xi} - \frac{d\chi_{b}(\xi)}{d\xi} \chi_{b}(\xi) \right] d\xi
\]

\[
+ \frac{2}{R(t)(1 + R(t))} \sum_{k} b_{\ell t}(t) \int_{0}^{\xi_{\max}} \left[ \chi_{b}(\xi) \frac{d\chi_{b}(\xi)}{d\xi} - \frac{d\chi_{b}(\xi)}{d\xi} \chi_{b}(\xi) \right] d\xi + \frac{\ell(\ell + 1) - \ell(\ell + 1)}{r_{\Sigma}} a_{\ell t}(t) \right\}.
\]  

(A.3)

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For the time-independent Hamiltonian, i.e., if there is no light field so that $H_{\text{int}} = H(0)$, it can be shown that $B < \sqrt{2} \max_{e_{\text{det}}}(E_{\text{det}})_{\text{det}} \exp[\max_{e_{\text{det}}}(E_{\text{det}})_{\text{det}}/(K - 1)]$. Additionally, if $H(0)$ is positive definite, i.e., if $H(0)$ holds no bound state, it can also be shown that $B < 2^{(K-2)/(K-1)} \max_{e_{\text{det}}}(E_{\text{det}})_{\text{det}}$. The latter is not the case in our discussion, but may give better estimation to our error and stiffness assessment, because the stiffness mainly arises from very large angular momentum states supporting almost no bound state. See Refs. [44, 61] for mathematical details.

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To see a typical behavior of the phase gradient, for simplicity, let us consider the time evolution of a one-dimensional free electron prepared at $t = 0$ in a normalized Gaussian wave packet having width $\Delta x$ and central momentum $k_0$:

$$
\psi(x,t) = \frac{1}{\sqrt{2\pi}(\Delta x)^2 + it/2} \exp\left[i k_0 x - ik_0^2 t/2 - \frac{(x - k_0 t)^2}{4[(\Delta x)^2 + it/2]}\right].
$$

The derivative of the phase with respect to $x$, $d[\arg \psi(x,t)]/dx = k_0 + t(x - k_0 t)/[4(\Delta x)^2 + t^2]$, linearly increases with $x$, and is greater than $k_0$ for $x > k_0 t$.

Also see the discussion in Ref. [26].

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