CHIRALEX: theory

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I. INTRODUCTION

Essential parameters are kept in the params dictionary, which is passed throughout the program. It is used to construct all types of objects, from grids to the propagator object.

II. BASIS FUNCTIONS AND GRIDS

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A. Radial quadratures

B. Radial grid

The radial grid used to solve the TDSE is based on an underlying Gauss-Lobatto (GL) quadrature in an underlying Gauss-

$$S_k = \frac{1}{2} R_b \left(x_k + 1 \right) \tag{1}$$

Bin size must be determined empirically to ensure good convergence and stability of the results. Next, the scaled grid is copied and translated by the bin size to generate the full primitive radial grid G_{prim} :

$$r_{ik} = S_k + T_i, \qquad i = 0, 1, ..., Nb-1, k = 0, 1, ..., Nlobs-1$$

where $T_i=i\cdot R_b+\epsilon$. The constant shift ϵ (rshift keyword) is optional and by default is set to 0. The resulting grid has the size $Nb\times Nlobs$ and contains boundary points $r_{00}=\epsilon$ and $r_{Nb-1,Nlobs-1}=R_{max}$. The last point from bin i and the first point from bin i+1 are identical in the primitive grid.

The coupled grid G is generated by removing boundary points, plus merging duplicate points at bin boundaries in G_{prim} . Through these operations the size of the coupled grid is $Nr = Nb \times (Nlobs - 1) - 1$. Note that all indices start from 0 for python compatibility. For example $G_{00} = r_{01}$ and $G_{0,Nl-3} = r_{0,Nl-2}$. The boundary of the first

bin is at $G_{0,Nl-2}=r_{0,Nl-1}=r_{10}$. The last point in the coupled grid is $G_{Nb-1,Nl-3}=r_{Nb-1,Nl-2}$.

C. Radial basis functions

1. Interpolated radial functions

D. Angular basis functions

III. INDEX MAPPING

A natural choice for mapping the basis set indices is to 44 set the grid points as major dimension (changing last). 45 Such a choice offers a simple way of cutting the Hamilto-46 nian matrix at a given grid point by slicing the basis. 47 We name this convention as 'DVR' mapping. Bridge 48 functions, as discussed in IIB, are defined for indices 50 quence of such a choice is a near block-diagonal structure 51 of the Hamiltonian. Another common choice is to choose bridge functions as first in the bin (n=0), which results in 'arms' reaching out from bin i into bin i+1, as shown in ??. Another option given in the code is to select mapping in which the angular quantum numbers l, m are the ma-56 jor dimension. We name this convention 'SPECTRAL'. 57 The choice for mapping is determined by the 'map type' = 'DVR,SPECT' keyword. With the 'DVR' convention 58 59 the map is given as follows:

$$\begin{array}{l} p(i,n,l,m) = \left(i\cdot(Nl-1) + n\right)\cdot\left(l_{max} + 1\right)^2 + l\cdot(l+1) + m \\ \text{60 with } p(0,0,0,0) = 0. \quad \text{For } l_{max} = 0 \text{ we have } p(Nb-61,Nl-3,0,0) = Nb\times(Nl-1) - 2, \text{ which retrieves the} \end{array}$$

with p(0,0,0,0) = 0. For $l_{max} = 0$ we have $p(Nb - 1, Nl - 3, 0, 0) = Nb \times (Nl - 1) - 2$, which retrieves the size of the coupled radial grid/basis set. The map is stored in a numpy array ('maparray') and is saved in file. The ranges for the radial indices i and n are the same as for the coupled grid, i.e. i = 0, 1, 2, ..., Nb - 1 and n = 0, 1, 2, ..., Nl - 2 for i < Nb - 1 and n = 0, 1, 2, ..., Nl - 3 for i = Nb - 1.

59 IV. ROTATED ELECTROSTATIC POTENTIAL

Each molecular-frame orientation gives different elecratrostatic potential as seen from the laboratory frame, in which the basis set is defined. This means that for each orientation one must generate generally a different Lebedev quadrature grid. Even if a global scheme is used,

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TABLE I. Example 'DVR' coupled basis set index mapping for Nb = 3, Nl = 3, lmax = 1

•	max = 1.						
	i	$ \mathbf{n} $	$ \xi $	1	m	$ \mathbf{p} $	
	0	0	0	0	0	0	
	0	0	0	1	-1	1	
	0	0	0	1	0	2	
	0	0	0	1	+1	3	
	0	1	1	0	0	4	
	0	1	1	1	-1	5	
	0	1	1	1	0	6	
	0	1	1	1	+1	7	
	2	0	$ _4$	1	0	18	
	2	0	$ _4$	1	+1	19	

75 the values of the electrostatic potential will be different 76 at these points for each orientation. This is because the 77 lebedev grid is defined in the laboratory frame. Therefore 78 at this stage, a separate Psi4 calculation must be initial-79 ized for each orientation. I do not see a way around it so at the moment, other than performing multipole expansi sion and calculating the angular part (spherical tensor 82 form) of the potential analytically. But finding this ex-83 pansion is costly, even more than running Psi4 for every 84 orientation.

KINETIC ENERGY OPERATOR

VI. POTENTIAL ENERGY OPERATOR

HAMILTONIAN VII.

The current paradigm assumes calculation of the so sparse Hamiltonian matrix and feeding this matrix into an eigensolver or propagator routine. In future versions, 91 for better computational performance it may be useful 2 not to calculate the Hamiltonian matrix, but rather code 93 the matrix-vector product only, as part of an interface 94 routine with appropriate iterative eigensolver. Due to 125 95 high degree of sparsity of the Hamiltonians appearing in 96 our calculations, it is sufficient to keep in memory the whole sparse matrix.

We decided to keep the bound state Hamiltonian H_0 and the initial propagation Hamiltonian H_{init} as sepa-126 rate entities, calculated independently. The computa- 127 tures: 101 tional overhead related to calculating the bound-state part twice is marginal, as we assume that the full propagation basis is much bigger than the basis for the bound Hamiltonian $Nbas \gg Nbas0$. Future releases might re-105 cycle the bound state KEO and POT and build only the 128 where $(\tilde{\theta}_s, w_s)_{s=0,1,...,s_{max}-1}$ are the quadrature points 106 outside region Hamiltonian.

WAVEPACKET PROPAGATION

time grid

The time grid for the calculation is determined by the 109 110 input keywords:

TABLE II. Time grid keywords.

keyword	description	type
t 0	start time	float
$t \max$	${ m end\ time}$	float
$\mathrm{d}\mathrm{t}$	time step	float
${\rm time_units}$	units	string
$wfn_saverate$	save rate of the wavepacket	float

112 The generated grid is equidistant given by the following 113 114 formula:

$$t[i] = t_0 + i \cdot dt; \ i = 0, 1, 2, ..., Ntpts$$
 (4)

115 where

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$$Ntpts = \left\lceil \frac{t_{max} - t_0}{dt} \right\rceil \tag{5}$$

116 such that the array of time-points contains elements: 117 t[0], t[1], ..., t[Ntpts]. Note that the last point (tmax) is 118 included in the grid.

WAVEPACKET ANALYSIS

Momentum space functions

1. Legendre decomposition

The 2D momentum probability distribution can be decomposed into a Legendre polynomials series for each length of the electron's wavevector:

$$W(k,\tilde{\theta}) = \sum_{n=0}^{n_{max}} b_n(k) P_n(\cos\tilde{\theta})$$
 (6)

where the expansion coefficients are given as:

$$b_n(k) = \frac{2n+1}{2} \int_{-1}^{1} W(k, \cos \tilde{\theta}) P_n(\cos \tilde{\theta}) d\cos \tilde{\theta}$$
 (7)

The integral in eq. 7 can be calculated using quadra-

$$b_n(k) = \frac{2n+1}{2} \sum_{s=0}^{s_{max}-1} w_s W(k, \cos \tilde{\theta}_s)) P_n(\cos \tilde{\theta}_s) \quad (8)$$

129 and weights pairs.

INPUT FILE

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Molecule

- molec name: name of the molecule (chiralium, h, 174
- matelem method: method of choice for the calculation of the potential energy matrix elements. Availability of methods given below depends on the chosen molecule.
 - analytic: use analytic formulas for the ma- 181 trix elements. When the potential is given as 182 an expansion in the spherical harmonics basis, utilize 3j symbols to calculate the matrix elements.
 - lebedev: use own implementation of the spher- 186 ical Lebedev quadratures with fixed global 187 quadrature level for all grid points defined 188 in $sph\ quad\ global=$ "lebedev_119", "lebe-189 dev 131", etc.
 - lebedev adaptive: use own implementation of 191 the spherical Lebedev quadratures with adap- 192 tive quadrature level, optimized for each grid 193 point. Stopping condition is based on convergence criteria defined in sph_quad_tol tolerance (in a.u.) for the convergence of matrix elements with the maximal values of the 196 l quantum number.
 - quadpy: use the Quadpy library to calculate 199 the spherical integrals with Lebedev quadra- 200 201
 - others: to be implemented.
- 203 • esp mode: method of choice for the calculation of 204 the cationic's core electrostatic potential. 205
 - analytic: available only for the hydrogen atom and some model systems.
 - -psi4: use psi4 quantum chemistry package for calculate the ESP for all required points.
 - only available for the chiralium molecule. Read the precomputed potential 208 kindly provided by A. Artemyev.
 - psi4_interpolated: produce an interpolated ²¹⁰ LP linearly polarized field """ ESP based on psi4 points.

- others: to be implemented. Any method one can imagine, including the use of other quantum chemistry software.
- $keo\ method = klist, slices, blocks$. Method of choice for the calculation of the kinetic energy matrix elements. See V for more details. Three main methods are considered so far:
 - klist calls build keomat klist, which uses a precomputed list (klist) of all matrix indices for which the matrix elements can be nonzero. It is by far the slowest method, even when Numba jit acceleration is used.
 - slices calls build keomat slices, which utilizes the fact that identical copies of the K_D and K_C matrices are distributed over the KEO matrix. The respective function loops over FEM-DVR bins and in each bin appropriately indexed matrix elements of the KEO matrix have the respective elements of K_D and K_C assigned. The algorithm relies on slicing a scipy.sparse.lil matrix. The centrifugal energy term is added to the diagonal at the end using *scipy.sparse.diags* method.
 - blocks calls build keomat blocks, which constructs the KEO using scipy.sparse.block diag method. First a block-diagonal matrix formed from inflated K_D matrices is formed. Next the inflated K_C matrices are further inflated to the size of each K_D block and a block-diagonal matrix is formed. The resulting matrix is then stacked with appropriate empty matrices to match the shape of the KEO. Hermitian conjugate of this matrix is calculated. All three matrices are then combined. The centrifugal energy term is added to the diagonal at the end using *scipy.sparse.diags* method.

Field В.

""" Available field types: 1) RCPL - right-circularly polarized field 2) LCPL - left-circularly polarized field 3)

""" Available envelopes: 1) gaussian 2) sin2 """