**Marquette University**

**e-Publications@Marquette**

***Chemistry Faculty Research and Publications/College of Arts and Sciences***

***This paper is NOT THE PUBLISHED VERSION*.**

Access the published version via the link in the citation below.

*Computer Physics Communications*, Vol. 267 (October 2021): 108084. [DOI](https://doi.org/10.1016/j.cpc.2021.108084). This article is © Elsevier and permission has been granted for this version to appear in [e-Publications@Marquette](http://epublications.marquette.edu/). Elsevier does not grant permission for this article to be further copied/distributed or hosted elsewhere without the express permission from Elsevier.

SpectrumSDT: A Program for Parallel Calculation of Coupled Rotational-Vibrational Energies and Lifetimes of Bound States and Scattering Resonances in Triatomic Systems

Igor Gayday

Department of Chemistry, Wehr Chemistry Building, Marquette University, Milwaukee, WI

Alexander Teplukhin

Theoretical Division (T-1, MS B221), Los Alamos National Laboratory, Los Alamos, NM

Jonathan Moussa

Molecular Sciences Software Institute, Blacksburg, VA

Dmitri Babikov

Department of Chemistry, Wehr Chemistry Building, Marquette University, Milwaukee, WI

# Abstract

We present SpectrumSDT – a program for calculations of energies and lifetimes of bound rotational-vibrational states below and scattering resonances above the dissociation threshold on a global potential energy surface of a triatomic system, which may include stable molecules, weekly-bound van-der-Waals complexes, and unbound atom + diatom scattering systems. Large-amplitude vibrational motion is treated explicitly using hyper-spherical coordinates. Three options for the rotational-vibrational interaction are supported: uncoupled (symmetric top rotor), partially coupled (to include interaction between several nearest states only) and full-coupled (vibrating asymmetric-top rotor). In addition to energies and lifetimes, SpectrumSDT is able to integrate ro-vibrational wave functions over the user-defined regions of potential energy surface, which helps to classify these states. In this release of the code, SpectrumSDT is limited to ABA-type molecules with wave functions that do not extend into the regions near Eckart singularities.

# Program summary

*Program title:* SpectrumSDT

*CPC Library link to program files:* https://doi.org/10.17632/9gftxjs4yk.1

*Developer's repository link:* https://github.com/IgorGayday/SpectrumSDT

*Licensing provisions:* GNU General Public License 3 (GPL)

*Programming language:* Fortran

*Nature of problem:* Calculations of energies and lifetimes of bound rotational-vibrational states below and scattering resonances above the dissociation threshold on a global potential energy surface of a triatomic system, which may include stable molecules, weekly-bound van-der-Waals complexes, and unbound atom + diatom scattering systems.

*Solution method:* A Hamiltonian matrix is built in APH coordinates using an optimal 2D basis set, adjusted for a given problem. A complex absorbing potential (CAP) is added to define a boundary condition for calculation of scattering resonances above the dissociation threshold. The eigenstates of the Hamiltonain matrix are found using a state-of-the-art iterative eigensolver.

*Restrictions:* The present version is restricted to ABA-molecules and wave functions that do not extend into linear and equilateral triangle configurations.

*Additional comments including restriction and unusual features:* Probabilities and lifetimes of the wave functions can be calculated in user-defined regions on the PES, which allows to analyze their localization properties (i.e. automatic isotopomer assignment) and obtain channel-specific lifetimes for scattering resonances.

# Keywords

Software, Bound states, Scattering resonances, Rovibrational, Lifetimes

# 1. Introduction

Calculation of numerically accurate energies of bound rotational-vibrational states and scattering resonances of a molecular system provides essential information for many applications. The spectra of bound states are used for predictions of spectroscopic properties of the molecules1, whereas scattering resonances play role of the metastable intermediate species in the chemical reactions and their properties are used as input for the kinetics simulations2. One particularly interesting example of this sort is the study of the mysterious mass-independent isotope fractionation effect in ozone3,4,5,6,7,8,9,10, for which SpectrumSDT was initially developed and applied11,12,13,14,15,16,17,18,19. In those calculations we were primarily focused on the kinetics of ozone recombination reaction , treated based on energies and lifetimes of scattering resonances above the dissociation threshold, as well as localization properties of their wave functions over the regions of interest on the potential energy surface (PES), such as covalent wells or Van der Waals plateaus.

In this manuscript we present the first general and user-ready version of SpectrumSDT (v1.0) that can be applied to the triatomic systems other than ozone. Several programs for the ro-vibrational calculations exist, such as ScalIT20,21 Geniush22,23,24, DVR3D25, BOUND26, MOLSCAT27, TRIATOM28 and Hibridon29 but our code offers unique features. In particular, SpectrumSDT can find states both below and above the dissociation threshold, with or without ro-vibrational coupling, with or without lifetime information. Moreover, it carries out integration of computed wave functions over user-defined regions of the PES, in order to automatically assign isomers (or isotopomers) and compute channel-specific decay rates of scattering resonances (spontaneous first order process determined by resonance widths). This information is essential for the description of reactions characterized by complicated potential energy landscape with multiple reaction channels. Finally, our code offers three options for the treatment of rotation-vibration coupling. Namely, in addition to the standard symmetric-top rotor approximation (when the coupling is neglected) and the exact full-coupled calculations (which is numerically demanding), we provide an affordable partially coupled approach. In this first release, SpectrumSDT is limited to ABA-type molecules (where two atoms are the same, and one is different, including the AAB and BAA isomers) and is applicable to wavefunctions that do not extend into the regions of Eckart singularities (equilateral and linear shapes). These limitations will be lifted in future releases.

The manuscript is structured as follows: Section 2 contains a brief overview of theoretical aspects; Section 3 describes practical aspects, such as setting up input parameters and interpreting output files; Section 4 provides some examples of applications of SpectrumSDT; Section 5 summarizes this manuscript with conclusions.

# 2. Theory

This section contains a brief overview of the most relevant theoretical aspects of SpectrumSDT for a lay user. For a more complete description the readers are encouraged to refer to other recent publications, where all theory aspects have been covered in detail11, 17, 18, 19.

## 2.1. Adiabatically adjusting principal axis hyperspherical (APH) coordinates

SpectrumSDT operates in APH coordinates30, which allow to treat all dissociation channels on equal footing. Molecular geometry in the APH coordinates is described with hyper-radius  and two hyper-angles  and . Qualitatively  serves as a measure of the overall “size” of the bounding triangle for the system (the “breathing” motion);  changes in the range from 0 to  and corresponds to translation from equilateral triangle at  to a linear molecule at  (the “bending” motion);  changes in the range from 0 to  and is responsible for isomerization, such that ,  and  correspond to AAB, ABA and BAA arrangements respectively, as shown in Fig. 1. See also Ref.31 for an interactive visualization tool of the motion along  and Ref.32 for 3D representation of APH coordinates on the example of ozone.

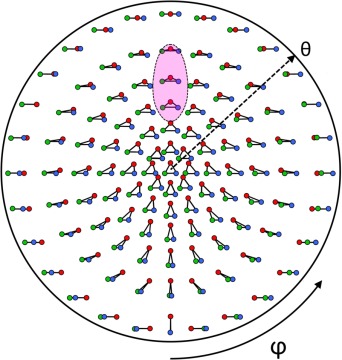


Fig. 1. Visualization of molecular arrangements corresponding to different values of and a fixed value of  in APH coordinates (adapted from Ref.52). The value of  increases radially outward from the center (where ) to the rim (where ).  is an angle in the range measured from the bottom of the circle. Symmetric obtuse ABA configuration corresponds to . The value of  defines overall size of the triangle formed by the three atoms. The pink area remarks the boundaries of energetically accessible shapes of SO2 at 8000 cm−1 (double of the maximum energy considered here). (For interpretation of the colors in the figure(s), the reader is referred to the web version of this article.)

## 2.2. Basis sets and sequential diagonalization truncation (SDT) procedure

On the lowest level, SpectrumSDT uses discrete variable representation (DVR) for  and  coordinates (basically, a grid of points) and variational basis representation (VBR) of cosines or sines functions for , which enforces either symmetric or antisymmetric property of wave functions with respect to  and , required for ABA-molecules in the APH coordinates. The two symmetries are independent, so, the calculations are split into two separate runs for each symmetry.

SpectrumSDT constructs a hierarchy of progressively more optimal basis sets through the procedure known as Sequential Diagonalization Truncation (SDT)33,34. In short, SDT starts with solving a one-dimensional Schrodinger equation, where all couplings with other coordinates are not included, in each of the 1D-slices along  at every combination of grid points in  and . The obtained 1D-solutions are used as a locally optimal basis set to solve 2D problem in  and , where the additional couplings are introduced, and the basis set is further optimized (sequential diagonalization). Finally, the 2D solutions are used as a basis set for the overall 3D problem, with all couplings taken into account. Of course keeping all the solutions from 1D and 2D levels would result in unnecessarily large basis, so only the solutions with energies less than a specified convergence parameter Ecut are retained in the basis set (truncation).

## 2.3. Hamiltonian operator in APH coordinates and Eckart singularities

The Hamiltonian operator in APH coordinates can be written as18:

(1) ,

where the individual terms are given by:

(2) ,

(3) ,

(4) ,

(5)

(6)

(7)

(8)

and  is the potential energy term.

The rotational constants *A*, *B* and *C* in Eqs. (6)–(8) are given by:

(9)

(10)

(11)

and  is the reduced mass of the triatomic system, defined as:

(12)

One can see that  as  in Eq. (4) and  as  or  in Eq. (5). These are Eckart singularities corresponding to the equilateral configuration at  and the linear configuration at . While this problem is in principle solvable within existing framework by using a special basis set35, the current release of SpectrumSDT does not implement this feature yet, therefore if a wave function for a particular molecule is allowed to have a significant probability near  or , the solution is not reliable. This issue will be addressed in future releases.

Fortunately, in many cases this issue is naturally avoided. First, many covalently bound triatomic molecules with  hybridization, such as O3, naturally have a highly repulsive PES in the regions of Eckart singularities, which prevents the vibrational wave function from reaching there. Second, the rotational potential term , given by Eq. (6), approaches in the same regions due to the *B* and *C* terms in Eqs. (10) and (11). This extra potential “shields” Eckart singularities at sufficiently high values of rotational excitation , even if the PES does not. For both of these cases our code is able to give accurate results.

## 2.4. Hamiltonian matrix structure

The rotational state of the basis functions used to build the Hamiltonian matrix can be either *fixed* or *adiabatic*19. In the *adiabatic* case, a separate basis set is used for each value of the symmetric top quantum number  (projection of total angular momentum onto -axis) that is going to be included in the Hamiltonian matrix for a given value of total angular momentum quantum number . This creates a better suited basis set, since it experiences the same rotational potential as a target wave function. This approach is relatively standard and often found in the literature13, 14, 15, but it requires computing many different basis sets and, most importantly, the corresponding overlaps for all values of  and , which can be numerically demanding.

In the alternative *fixed* option, the basis set with one (representative) combination of  and is used to describe wave functions with arbitrary values of  and (a similar idea has been utilized in the two-step approach of Ref.28). This reduces the computational cost of both *basis* and *overlaps* calculation stages, but creates a less efficient basis set, therefore a higher value of the *cut-off energy* (Ecut) may be needed. Recently we demonstrated that this approach is more efficient19. However, one has to be careful when handling a molecule with wave functions reaching the ends of -grid. In this case, using a basis set with high values of  and relative to a target wave function, will exclude basis functions near the ends of the -grid (due to the effect of rotational potential), which may result in less accurate representation of the target wave function in that area.

The rotational terms in Eqs. (6)–(8) translate to the Hamiltonian matrix structure shown in Fig. 2. In the cases when couplings between different Λ blocks can be neglected, the overall matrix can be split into the diagonal Λ blocks, which significantly simplifies the process of solving the eigenvalue problem. This is known as symmetric-top rotor approximation. As a compromise between speed and accuracy, when the effects of ro-vibrational coupling cannot be neglected, it is also possible to partially take it into account, by including only a certain number of adjacent blocks for each value of Λ. For example, including up to five blocks, within the  range, permits to capture most of the rotational-vibrational interaction.

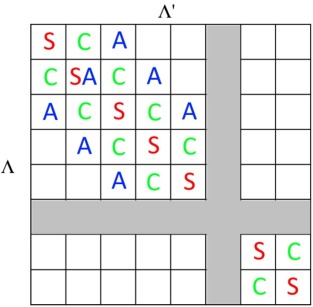


Fig. 2. Rotational block structure of the Hamiltonian matrix. Letters S, A and C indicate contributions from symmetric-top rotor, asymmetric-top rotor and Coriolis coupling terms, given in Eqs. (6)–(8) respectively. Other blocks of the matrix are zero. The blocks are labeled by and , the value of projection of total angular momentum  onto -axis.

## 2.5. Optimized -grid (optional)

Suppose we have a 1D potential  and we want to place points along  -coordinate in an optimal way to represent wave functions. The idea is to make grid spacing in  proportional to the local de Broglie wavelength . This way one can accurately capture wave function oscillations, without placing too many points in the regions where wave periods are large. To achieve this, one can define an auxiliary grid , where the distance is unitless and a given point  corresponds to  oscillations of a wave with total energy  in the original space (), as measured from a starting point . The points in *x* can be mapped to  and vice versa, therefore if one has an equidistant grid in , its mapping onto  would be an optimal grid with a fixed number of points per oscillation of the wave. Assuming the step size in  is , one can derive the following differential equation that relates the two grids11,36:

(13)

where factor of 2 in the denominator is chosen to make  correspond to 2 points per period – the smallest meaningful number. This way the values of  range from 0 to 1 and control the density of the resulting optimal grid (smaller values of *α* generate denser grids). The exact value of  for a given problem is a convergence parameter. The derivative  in Eq. (13) is further referred to as Jacobian of the grid .

In general, any function smaller than the actual potential can be used as  in Eq. (13). Using such potential “envelopes” can be helpful to increase density of points in selected regions or improve integration accuracy if the envelope function is analytical.

# 3. Practical considerations

## 3.1. General

The workflow of the program is separated into the following stages (runs): grids, potential, basis, overlaps, eigensolve and properties. Using multiple stages allows to utilize processors efficiently by separating computationally expensive sections with different degree of parallelizability. The stages have to be launched sequentially one after another from their respective folders.

The folder structure for each calculation may consist of up to 4 levels (see Fig. 3). On the first level, there are subfolders for each value of (referred to as more traditional in the code, parameters, file names and further in the text to avoid confusion), as well as separate folders for each range of  that a user wants to consider for calculation of eigenpairs. The range that includes all values of  for a given  is referred to with a keyword all. The next two levels in a  range folder identify the values of parity and wave function symmetry corresponding to the calculation within. The folders corresponding to a scalar value of  skip the level corresponding to parity since parity only has the effect on calculations involving more than one value of 18. The last level is divided into folders corresponding to stages from basis to properties. Grids and potential stages do not have a folder within a particular calculation since they can be shared across multiple calculations. The output files from each stage are stored either in the stage folder itself, or in out\_stagename subfolder within the stage folder.

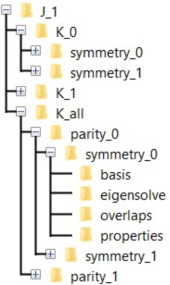


Fig. 3. An example of root folder structure for . Inner structure of the folders marked by “+” is not shown for clarity. All folders on the same level of hierarchy have identical inner structures, except -folders corresponding to a singular value of *K* skip the *parity* level.

The stage folders all need to have a copy of a configuration file with the values of , parity, symmetry, and stage corresponding to a given subfolder. A python script init\_spectrumsdt\_folders.py in scripts folder is available to generate the correct folder structure, copy a given template configuration file to all target subfolders and fill out placeholder values corresponding to each folder for a given value of . Run init\_spectrumsdt\_folders.py –help for more details. To keep the parameters consistent between stages, we do not recommend manually changing the configuration files generated by init\_spectrumsdt\_folders.py in the stage subfolders.

The configuration file has to be named spectrumsdt.config and has to be present in every stage folder. Each parameter is assigned in the format *key* = *value*. Both keys and values are case-sensitive, and each assignment has to start on a new line and finish on the same line, except for parameter groups. The order of parameters, number of spaces, empty lines, or characters after the comment mark “!” do not matter. In the case of parameter groups, the open parenthesis “(” has to be present on the same line after the assignment character “=”, followed by assignment of individual parameters of the group on the following lines and ending with the close parenthesis “)” on a separate line. Some examples of input files can be found in config\_examples folder.

All stages except grids support parallel execution with an arbitrary number of processes, implemented via message passing interface (MPI).

## 3.2. Grids

The purpose of this first stage is to generate grids for all APH coordinates. Even though the basis set for  is not a DVR, a  -grid is still generated for numerical integration of the basis functions in . The grids for  and  are described with the parameter groups grid\_rho and grid\_theta, which include at least the following parameters (determined by convergence studies):

• from – the left border of the grid,

• to – the right border of the grid,

• step – grid step,

• num\_points – the total number of points in the specified interval.

Specifying step automatically defines num\_points and vice versa, so these parameters cannot be specified together. All parameters with units are specified in Bohr for grid\_rho and in degrees for grid\_theta. The  -grid is always defined in the range from 0 to , so only the number of points is specified via num\_points\_phi key.

After this stage is completed, the three grid files corresponding to each coordinate are generated. The grid files are written in the following format: the first line specifies the values of from, to, step and num\_points that were used to generate this grid. The next num\_points lines specify the coordinate of each point and, in case of  -grid, Jacobian value at that point. Jacobians are meaningful only for optimized grids and are always equal to 1 for equidistant grids. All grid points are placed in the centers of their respective intervals. All values with units in the grid files are specified in Bohr for  -grid and radians for - and  -grids.

*Advanced parameters*  In the calculations of scattering resonances one typically needs to set up a long-range grid in  to describe the dissociation region. In these cases, using an equidistant grid leads to unnecessarily dense points in the dissociative region. One can reduce the cost of such calculations by optimizing the grid step as described in Section 2.5.

Enabling optimization procedure for  -grid requires specifying grid parameter optimal = 1, alongside with the necessary information to define Eq. (13), which includes the following grid parameters:

* envelope\_path – the path to the file defining the envelope function . The function is defined by specifying its values at the grid points. Each row contains 2 numbers:  in Bohr and  in Hartree. The grid on which  is defined does not have to be the same as the  -grid for the eigenvalue problem, which allows to re-use the same envelope function for any  -grid. As a rule of thumb, it is recommended to use minimum energy path (MEP) along  in a given PES as the envelope function .

A python script extract\_MEP.py is available in the scripts folder to automatically generate minimum energy path along . The script takes no arguments and has to be invoked in the folder with calculated pes.out and grid files. The extracted MEP is written to a file named MEP\_rho.dat. Note that the MEP does not have to be calculated precisely, since precision of MEP does not affect the precision of subsequent calculations. A rough MEP may generate slightly suboptimal grid spacing, but the final grid density is a convergence parameter, controlled by step.

In the program, the points specified in the MEP file are used to build a cubic spline for the right branch of , and fit analytical Eckart potential for the left branch . Possible oscillations of the cubic spline can easily be fixed by changing MEP grid density, which can be done for free since it does not affect complexity of the overall problem. Moreover, as outlined in the previous paragraph, even if some oscillations are present, they will not alter accuracy of calculations.

* max\_energy – the value of  in wave numbers. This specifies the maximum total energy considered for a particle moving in the specified envelope potential .  has to be larger than  for any value of . Local density of grid points is proportional to , therefore  can be considered as a tool to control balance of points between the covalent well and the dissociation region. In the limiting case when  is equal to the asymptotic value of  in the dissociation region, all the grid points would be placed in the covalent well region and little or none in the dissociation region. For larger values of , the difference between  in the well and dissociation regions becomes less pronounced and points are generated in a more uniform fashion.
* solver\_steps – optional, specifies number of steps in Runge-Kutta algorithm used to solve Eq. (13). Typically, does not need to be set explicitly. The default value of 2048 is expected to work well for majority of applications.

Note that when optimized grid is requested, step grid parameter controls the step size in the auxiliary equidistant *x*-grid, i.e. step sets the value of  in Eq. (13).

In addition to these parameters, the following config parameter has to be specified to calculate the reduced particle mass :

* mass – the description of masses of atoms in the system under consideration. The masses have to be listed as 3 comma-separated numbers corresponding to masses given in atomic unit of mass (*i.e*., in masses of an electron, ) of atoms in ABA order. Instead of providing an explicit mass value, it is possible to use a shortcut consisting of an element symbol and its mass number. For example, H1 will be replaced with the mass of hydrogen-1 and O18 will be replaced with the mass of oxygen-18. These shortcuts exist for all stable isotopes of the elements in the first three periods. The exact values of isotopic masses are taken from Ref.37 and can be found in src/base/constants.f90 file.

Note that mass is a config parameter, and not a grid parameter, therefore it is not a part of a grid setting group and should be specified as a top-level parameter. An example of a config file that generates optimized grid for ozone can be found in config\_examples folder.

## 3.3. Potential

Majority of PESs do not operate in APH coordinates directly, therefore a conversion from APH to a more common coordinate system is likely required. SpectrumSDT offers two ways to alleviate this procedure. One way is to use output\_coordinate\_system key to carry out coordinate conversion at grids stage. If used with valid values other than aph, pes.in file will be created in addition to the grid files, where each row after the header row specifies molecular geometry converted to the chosen coordinate system. In response user needs to provide a file named pes.out, with the values of the PES in Hartree at geometries in pes.in. Providing pes.out file is how SpectrumSDT obtains information about user-specified potential, i.e. there is no need to modify the code to define a potential subroutine. Note that pes.out is not expected to have a header line, therefore row numbering is shifted by 1 relative to pes.in.

The following values of output\_coordinate\_system are supported:

* aph – the default value. pes.in file is not generated since it is trivial to simply iterate the values in the grid files.
* mass jacobi – mass-scaled version of Jacobi coordinates. See Eqs. (52a)–(52c) in Ref.30.
* jacobi – regular Jacobi coordinates, where  (Bohr) is the distance between the terminal A-atoms (in ABA notation); *R* (Bohr) is the distance from the center of mass between the two A atoms (midpoint) to the B atom; and (in radians) is the angle between these two vectors.
* cartesian – molecular geometry is described with - and - coordinates of atom B and -coordinate of an atom A. The remaining 6 Cartesian coordinates are fixed to 0. All values are given in Bohr.
* all bonds – three pair-wise distances in Bohr between the atoms. Indexes 1 and 3 are assigned to the A atoms and index 2 is assigned to B.
* internal – two A-B bond lengths in Bohr and A-B-A angle in radians.

Note that mass has to be specified if a value other than aph or mass jacobi is given to output\_coordinate\_system, since coordinate conversion is mass-dependent. This makes it necessary to re-calculate potential values for different isotopomers even if underlying PES program is the same.

Another way is to directly use src/base/coordinate\_conversion.f90 module in user's PES program to convert coordinates dynamically. In this case the user can set output\_coordinate\_system to aph to avoid generating pes.in and read the APH-grids directly. The combinations of APH points should be iterated in order , ,  (*i.e.*,  changes least frequently,  changes most frequently). An example of a Fortran program that reads the grids and writes ozone potential of Dawes et al.38, in this way can be found in PES\_examples/ozone/ozone\_pes.f90.

## 3.4. Basis

This stage uses the grids and potential from the previous two stages to solve 1D and 2D problems in the given potential (see Section 2.2). The following parameters have to be specified:

* J – total angular momentum quantum number for basis functions.
* K – projection of total angular momentum onto -axis quantum number for basis functions.
* basis – parameter group that includes the following parameters:
  + num\_functions\_phi – number of VBR basis functions to include in the primitive basis set for 1D problem (convergence parameter).
  + symmetry – “0” for symmetric basis functions (cosines); “1” for antisymmetric functions (sines).
  + cutoff\_energy – maximum energy of a solution in 1D or 2D problem (convergence parameter, in wavenumbers). Solutions with energies above this are not included in the basis set.
  + min\_solutions – minimum number of solutions retained in each 1D and 2D problem, even when energies exceed cutoff\_energy, typically does not need to be set. The default value of 3 is expected to work well for majority of applications.
* grid\_path – full path to a folder with APH-grids and pes.out files.
* root\_path – full path to the top-level folder of the current calculation (from where init\_spectrumsdt\_folders.py was executed).

In addition, stage has to be set to basis and mass has to be specified. The number of 2D solutions kept in basis for each value of  is written to num\_vectors\_2d.dat file. The binary files storing 1D and 2D basis functions are written to out\_basis folder.

Note that multiple basis calculations with different values of K and symmetry are required for a single problem in cases when the Hamiltonian matrix consists of multiple -blocks and the *adiabatic* basis set is employed.

The basis set is calculated independently for each value of , which are distributed among the available processors. Therefore the number of points in  -grid limits the maximum number of processors that can do useful work at this stage.

## 3.5. Overlaps

This stage uses basis functions from the previous stage (basis) to calculate the Hamiltonian matrix elements for the row of blocks with the given value of K. In addition to the parameters specified at basis stage, the following parameters are required:

* fixed\_basis – at this stage specifying this parameter group instructs the program to compute additional overlaps necessary for the *fixed* basis mode (see Section 2.4).
* use\_rovib\_coupling – “0” is for uncoupled symmetric top; specifying “1” instructs the program to compute additional overlaps for the off-diagonal -blocks (see Fig. 2) necessary to build the Hamiltonian matrix with rotational-vibrational coupling terms.

All output files from this stage are binary and are stored in the out\_overlaps folder. Calculation of individual matrix elements is independent and distributed among available processors, therefore this stage is expected to scale efficiently with the number of processors.

## 3.6. Eigensolve

This stage uses the matrix elements computed at the previous stage (overlaps) to build the Hamiltonian matrix and find its eigenpairs (energies and wave functions). In addition to parameters specified at the overlaps stage, the following parameter are required:

* J – total angular momentum quantum number for the current problem. Note that this can be different from the value of J at the basis stage if *fixed* basis mode is employed.
* parity – the value of inversion parity of the wave functions, “0” or “1”18. Only matters if rotational-vibrational coupling is enabled (use\_rovib\_coupling = 1),  and  or 1 block is included in the Hamiltonian matrix.
* symmetry – the symmetry of the basis for  block, “0” for symmetric or “1” for antisymmetric basis. The subsequent values of  have to have alternating symmetry, therefore this choice defines symmetry for all values of  included in the Hamiltonian. Note that the symmetry is always defined with respect to  even in cases when there is no block corresponding to  in the Hamiltonian matrix.
* K – the values of  to include in the Hamiltonian matrix. Can either be a scalar number for a symmetric top rotor calculation; “all” for a fully coupled rotational-vibrational calculation with all valid values of  for given  and *parity*, from to ; or a custom -range in the form  to include only values of  for a partially coupled calculation. In the case of *adiabatic* basis, the basis and overlaps stages of appropriate symmetry have to be finished and available for all values of  specified in this parameter.
* eigensolve – parameter group with the options for an eigenvalue solver. The options include the following:
  + num\_states – number of eigenpairs to converge.
  + ncv – largest dimension of the working subspace, optional, typically does not need to be set explicitly, set mpd instead. Determined automatically by SLEPc if left unset.
  + mpd – maximum projected dimension, optional, used to further restrict the size of projected eigenproblem on certain steps of the algorithm. Determined automatically by SLEPc if left unset. Refer to SLEPc manual for more details on ncv and mpd parameters39. For small number of eigenvalues we recommend leaving mpd unset. For larger number of eigenvalues one can improve performance by setting mpd manually. Too small values of mpd lead to quickly increasing solution time, while too large values require too much memory and also worsen running time, although not as quickly as too small values. The optimal value of mpd is problem-dependent, but in our experience the values around 0.2\*num\_states worked reasonably well.
  + max\_iterations – maximum number of iterations the solver is allowed to perform, optional, typically does not need to be set explicitly. The default value of 10000 is expected to work well for majority of applications.

For calculations in the *fixed* basis mode, one has to describe what rotational state should be used for the basis set. This is done through the fixed\_basis parameter group, which includes the following parameters:

* J – total angular momentum of the basis set functions.
* K – projection of the total angular momentum onto -axis of the basis set functions.
* root\_path – full path to the top-level folder with the basis and overlaps calculations for the values of J and K specified in this parameter group.

The lifetimes of the states are computed by adding a complex absorbing potential (CAP) in the form suggested by Manolopoulos40. By default CAP is not added. One can choose to add it by specifying parameter group cap, which consists of only one parameter:

* min\_absorbed\_energy – minimum energy at which absorption is required, specified in wavenumbers relative to energy of the dissociated molecule in the lowest dissociation channel (i.e. electronic dissociation energy plus zero-point energy). Note that enabling CAP is only meaningful for calculation of scattering resonances above the dissociation threshold.

If cap is specified, cap.txt file will be created. The value on i-th line of this file is the value of CAP at  given in cm−1. The CAP is always positioned relative to the far end of the  -grid (largest value of ), where it has the largest value and then gradually decreases for smaller values of .

The computed energies and widths of resonances are printed to file states.fwc. The wavefunctions are saved in binary form in out\_eigensolve folder. This stage is parallelizable to some extent, but parallelization efficiency is decreasing with the number of processors, therefore a relatively low number of processors (around 32) is advised for efficient utilization, but higher number of processors can also be used if efficiency is not a concern.

## 3.7. Properties (optional)

At this stage one can characterize the wave functions calculated at the previous stage by analyzing their properties, such as probability distributions over specified *sections* of the overall space and splitting of resonance width between the existing dissociation channels. The boundaries of these sections typically correspond to the edges of covalent or Van der Waals wells, or other objects of interest, which are PES-dependent, therefore it is up to the user to define them for a particular problem. Each section is defined as a new subgroup of a parameter group wf\_sections, specified by an arbitrary unique key and may include the following parameters:

* name – specifies section name for the header row in the output file. Optional, the value of the key, specifying this section, will be used as name if this parameter is not explicitly given.
* K – specifies the range of the values of  for the current section, optional.
* rho – specifies the range of  -values for the current section (in Bohr), optional.
* theta – specifies the range of -values for the current section (in degrees), optional.
* phi – specifies the range of  -values for the current section (in degrees), optional.
* stat – determines which statistic is to be calculated for the current section, optional. The possible values are probability (default) or gamma (resonance width). Specifying gamma instructs the program to integrate the wave function probability in a given section, multiplied by CAP, to get a portion of the overall width associated with this section. This can be useful to calculate channel-specific decay rates of the metastable states.

All ranges are specified in the form  to include the values from *A* to *B* inclusively. The final shape of the section in the 4D (,,,) space is the intersection of the ranges specified for each individual coordinate. Omitting all range parameters defines a trivial section that spans the whole space and has to evaluate to 1 if stat = probability, or total resonance width if stat = gamma.

The method of section definition implemented here limits their shapes to hyper-rectangles, but hyper-rectangles in the APH coordinates are often what one is looking for and more complicated shapes can be approximated by adding up multiple hyper-rectangles.

The results are written to states.fwc file, where the first two columns duplicate the results of the previous stage, and the next  (number of sections) columns add the statistics corresponding to the specified sections. Each processor computes its fraction of the wave functions independently from the other processors (except for the final merge of data for printing to a file), therefore high parallelization efficiency is expected for this stage.

# 4. Results

Numerous applications of SpectrumSDT to ozone isotopomers (especially in relation to the isotope effects) can be found in recent literature, which includes examples of symmetric top rotor calculations11, 12, 13, 14, 15, 16 and fully coupled rotation-vibration calculations17, 18, 19, both for bound states and scattering resonances with their lifetimes. Calculations in partially coupled regime are a work in progress and will be reported elsewhere. Various checks and comparisons proving the validity of the computed results have been presented. For example, Ref.16 includes comparison of computed energies and splittings for  (no rotation) with the previously reported results of Ndengué et al.41, and Petty et al.42, demonstrating an excellent agreement in all cases. Refs.17 and18 include comparison of fully coupled rotation-vibration energies for the values of  up to 5 with the results of Petty et al.42, and APH3D program of Kendrick et al.35, once again showing an excellent agreement. Spectroscopical constants fitted to the computed spectrum in Ref. 18 show a precise agreement with experimental results as well43,44.

In addition to plain energies and lifetimes of scattering resonances, SpectrumSDT can compute localization properties of the wave functions and channel-specific decay rates, which can be used as input for theoretical prediction of reaction rates, as it has been demonstrated in the recent study of ozone recombination reaction  and related isotope effects13, 14, 15. The theoretical framework presented there is straightforward to generalize for application to other three-atom systems.

In this section we present a benchmark calculation to demonstrate SpectrumSDT's applicability to molecules other than ozone, on the example of sulfur dioxide (SO2). SO2 is an ABA-molecule, whose PES prevents its wave functions from reaching Eckart singularities at the edges of -coordinate, therefore it is suitable for calculations with SpectrumSDT. The calculations in this manuscript use KAKPJG MRCI-F12 PES for the singlet ground  state of SO245. In contrast to ozone, which has three covalent wells, SO2 has only one well, therefore its PES is qualitatively different from ozone.

A comparison of results obtained with SpectrumSDT in this manuscript and those obtained in Klos et al.45, with ScalIT20, 21 is presented in Fig. 4. As one can see, the difference is on the order of 10−3 cm−1, same as what we observed for the case of ozone. Therefore, we conclude that the results of SpectrumSDT are accurate and the benchmark test is successful. The exact parameters used for this calculation can be found in config\_examples/so2 folder.

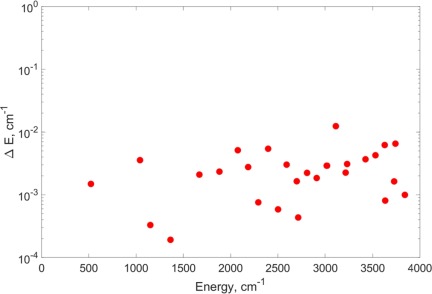


Fig. 4. Comparison between the vibrational energies of SO2 , calculated in this manuscript and those reported in Klos et al.45. The horizontal axis shows energies of states relative to the ground vibrational state. The vertical axis shows the absolute value of difference between the corresponding energy levels in the two sets of data.

# 5. Conclusions

In this manuscript we presented a first public version of SpectrumSDT: an open-source program for calculation of energies and lifetimes of bound states and scattering resonances of ABA-molecules in APH coordinates.

An extensive description of available options and practical recommendations have been provided. The accuracy of the program has been tested and verified on the examples of ozone11, 12, 13, 14, 15, 16, 17, 18, 19 and sulfur dioxide (this manuscript).

The program, building instructions and examples of running are available in the GitHub repository (see program summary) and in the Supplemental Information.

# Libraries credit

SpectrumSDT uses the following libraries:

* LAPACK for diagonalization of 1D and 2D Hamiltonians46.
* fdict for an implementation of dictionaries in Fortran47.
* SLEPc for an implementation of an iterative eigensolver48,49,50.
* PETSc as an underlying library for SLEPc51.

# Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

# Acknowledgements

This research was supported by the NSF AGS program Grant No. AGS-1920523. This research used resources of the National Energy Research Scientific Computing Center (NERSC), a U.S. Department of Energy Office of Science User Facility located at Lawrence Berkeley National Laboratory, operated under Contract No. DE-AC02-05CH11231. Igor Gayday acknowledges the support of MolSSI Investment Fellowship, funded by NSF ACI-1547580. Jacek Kłos is gratefully acknowledged for sharing the PES of SO2.

# References

1 J. Tennyson, S.N. Yurchenko, A.F. Al-Refaie, E.J. Barton, K.L. Chubb, P.A. Coles, S. Diamantopoulou, M.N. Gorman, C. Hill, A.Z. Lam, L. Lodi, L.K. McKemmish, Y. Na, A. Owens, O.L. Polyansky, T. Rivlin, C. Sousa-Silva, D.S. Underwood, A. Yachmenev, E. Zak. *J. Mol. Spectrosc*., 327 (2016), p. 73

2S.N. Elliott, K.B. Moore, A.V. Copan, M. Keçeli, C. Cavallotti, Y. Georgievskii, H.F. Schaefer, S.J. Klippenstein. *Proc. Combust. Inst*., 000 (2020), p. 1

3 K. Mauersberger. *Geophys. Res. Lett*., 8 (1981), p. 935

4 M.H. Thiemens, J.E. Heidenreich. *Science*, 219 (80) (1983), p. 1073

5 D. Babikov, B.K. Kendrick, R.B. Walker, R.T. Pack, P. Fleurat-Lessard, R. Schinke. *J. Chem. Phys*., 118 (2003), p. 6298

6 C. Janssen, J. Guenther, K. Mauersberger, D. Krankowsky. *Phys. Chem. Chem. Phys*., 3 (2001), p. 4718

7 M. Kryvohuz, R.A. Marcus. *J. Chem. Phys*., 132 (2010), Article 224305

8 M. Kryvohuz, R.A. Marcus. *J. Chem. Phys*., 132 (2010), Article 224304

9 M.H. Thiemens, S. Chakraborty, G. Dominguez. *Annu. Rev. Phys. Chem*., 63 (2012), p. 155

10 R. Schinke, S.Y. Grebenshchikov, M.V. Ivanov, P. Fleurat-Lessard. *Annu. Rev. Phys. Chem*., 57 (2006), p. 625

11 A. Teplukhin, D. Babikov. *J. Chem. Phys.,* 145 (2016), Article 114106

12 A. Teplukhin, D. Babikov. *Phys. Chem. Chem. Phys*., 18 (2016), Article 19194

13 A. Teplukhin, D. Babikov. *J. Phys. Chem. A,* 122 (2018), p. 9177

14 A. Teplukhin, I. Gayday, D. Babikov. *J. Chem. Phys.,* 149 (2018), Article 164302

15 A. Teplukhin, D. Babikov. *Faraday Discuss*., 212 (2018), p. 259

16 I. Gayday, A. Teplukhin, D. Babikov. *J. Chem. Phys*., 150 (2019), Article 101104

17 I. Gayday, A. Teplukhin, B.K. Kendrick, D. Babikov. *J. Phys. Chem. A*, 124 (2020), p. 2808

18 I. Gayday, A. Teplukhin, B.K. Kendrick, D. Babikov. *J. Chem. Phys*., 152 (2020), Article 144104

19 I. Gayday, E. Grushnikova, D. Babikov. *Phys. Chem. Chem. Phys*., 22 (2020), Article 27560

20 W. Chen, B. Poirier. *J. Comput. Phys*., 219 (2006), p. 185

21 W. Chen, B. Poirier. *J. Comput. Phys*., 219 (2006), p. 198

22 E. Mátyus, G. Czakó, A.G. Császár. *J. Chem. Phys*., 130 (2009), Article 134112

23 C. Fábri, E. Mátyus, A.G. Császár. *J. Chem. Phys*., 134 (2011), Article 074105

24 I. Simkó, T. Szidarovszky, A.G. Császár. *J. Chem. Theory* Comput., 15 (2019), p. 4156

25 J. Tennyson, M.A. Kostin, P. Barletta, G.J. Harris, O.L. Polyansky, J. Ramanlal, N.F. Zobov. *Comput. Phys. Commun*., 163 (2004), p. 85

26 J.M. Hutson, C.R. Le Sueur. *Comput. Phys. Commun*., 241 (2019), p. 1

27 J.M. Hutson, C.R. Le Sueur. *Comput. Phys. Commun*., 241 (2019), p. 9

28 J. Tennyson, S. Miller, C.R. Le Sueur. *Comput. Phys. Commun*., 75 (1993), p. 339

29 http://www2.chem.umd.edu/groups/alexander/hibridon/hib43/hibhelp.html

30 R.T. Pack, G.A. Parker. *J. Chem. Phys*., 87 (1987), p. 3888

31 A. Teplukhin, D. Babikov. *Chem. Phys. Lett.,* 614 (2014), p. 99

32 A. Teplukhin, D. Babikov. *J. Chem. Educ*., 92 (2015), p. 305

33 J.C. Light, T. Carrington. I. Prigogine, S.A. Rice (Eds.), *Adv. Chem. Phys*., John Wiley & Sons (2000), pp. 263-310

34 Z. Bačić, R.M. Whitnell, D. Brown, J.C. Light. *Comput. Phys. Commun*., 51 (1988), p. 35

35 B.K. Kendrick, R.T. Pack, R.B. Walker, E.F. Hayes. *J. Chem. Phys.,* 110 (1999), p. 6673

36 V. Kokoouline, O. Dulieu, R. Kosloff, F. Masnou-Seeuws. *J. Chem. Phys*., 110 (1999), p. 9865

37 T.J. Bruno, P.D. Svoronos. ***CRC Handbook of Chemistry and Physics.*** (93rd ed.) (2012)

38 R. Dawes, P. Lolur, A. Li, B. Jiang, H. Guo. *J. Chem. Phys.,* 139 (2013), Article 201103

39 J.E. Roman, C. Campos, E. Romero, A. Tomas. **SLEPc Users Manual.** (2020)

40 D.E. Manolopoulos. *J. Chem. Phys*., 117 (2002), p. 9552

41 S.A. Ndengué, R. Dawes, X.-G. Wang, T. Carrington, Z. Sun, H. Guo. *J. Chem. Phys*., 144 (2016), Article 074302

42 C. Petty, R.F.K. Spada, F.B.C. Machado, B. Poirier. *J. Chem. Phys*., 149 (2018), Article 024307

43 G.D. Carney, L.L. Sprandel, C.W. Kern. I. Prigogine, S.A. Rice (Eds.), *Adv. Chem. Phys*., John Wiley & Sons, New York (1978), p. 367

44 J.M. Flaud, R. Bacis. *Spectrochim. Acta, Part A, Mol. Biomol. Spectrosc*., 54 (1998), p. 3

45 J. Kłos, M.H. Alexander, P. Kumar, B. Poirier, B. Jiang, H. Guo. *J. Chem. Phys*., 144 (2016), Article 174301

46E. Anderson, Z. Bai, C. Bischof, S. Blackford, J. Demmel, J. Dongarra, J. Croz, A. Greenbaum, S. Hammarling, A. McKenney, D. Sorensen. ***LAPACK Users' Guide.*** (3rd ed.), Society for Industrial and Applied Mathematics, Philadelphia (1999)

47 https://github.com/zerothi/fdict

48 V. Hernandez, J.E. Roman, V. Vidal. *ACM Trans. Math. Softw*., 31 (2005), p. 351

49 V. Hernandez, J.E. Roman, V. Vidal. *Lect. Notes Comput. Sci*., Springer (2003), pp. 377-391

50 V. Hernandez, J.E. Roman, A. Tomas. *Parallel* *Comput*., 33 (2007), p. 521

51 https://www.mcs.anl.gov/petsc

52 V. Kokoouline, C.H. Greene. *Phys. Rev. A,* 68 (2003), Article 012703