ADT : A Generalized Algorithm and Program for Beyond Born-Oppenheimer Equations of 'N' Dimensional Sub-Hilbert Space

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USER MANUAL

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1 Introduction

Large and complicated functional form of adiabatic-to-diabatic transformation (ADT) equations is the major bottleneck of first principle based beyond Born-Oppenheimer (BBO) treatment, specially, when the number of coupled electronic states is high $(N \ge 4)$. Though explicit expressions for the ADT equations are already formulated for N=3 to 6 (See Ref. 24-28 of the main text), the algorithm 'ADT' generalizes this process of evaluating the expressions for any number of coupled electronic states (N). This program package can be efficiently used to carry out the following operations:

- Formulation of analytic functional forms of ADT quantities, i.e., ADT matrix, ADT equations, coefficient matrix of gradient of ADT angles (see Eq. 14 of the main text), coefficient matrix of nonadiabatic coupling terms (NACTs) (see Eq. 14 of the main text) and diabatic potential energy matrix for any N.
- Numerical calculation of ADT quantities (ADT angles, residue of ADT angles, ADT matrices
 and diabatic potential energy matrices) for any number of coupled electronic states, if the
 input files [ab initio adiabatic Potential Energy Surfaces (PESs) and NACTs] are already
 available.
- Interfacing with standard *ab-initio* packages such as MOLPRO¹, Gaussian² and Gamess³ to obtain equilibrium geometry, frequency and wilson matrix for spectroscopic systems.
- Interfacing of MOLPRO quantum chemistry package¹ to compute *ab initio* data, namely, adiabatic PESs and NACTs, which will serve as the input for ADT calculation if the input files (adiabatic PESs and NACTs) are not available to the user.

Finally, the program package 'ADT' can be used for the construction of diabatic surfaces of spectroscopic as well as scattering processes. For spectroscopic systems, the nuclear configuration space (CS) is expressed in terms of normal mode coordinates. On the other hand, user can employ Jacobi or hyperspherical coordinates for scattering systems.

2 Files and folder structure

Schematic Outline of Files present in 'ADT-version0' directory

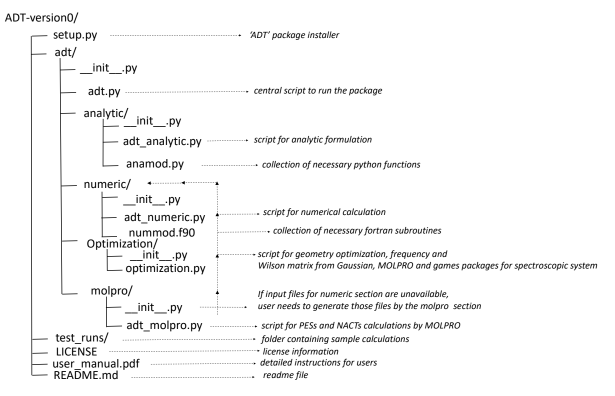
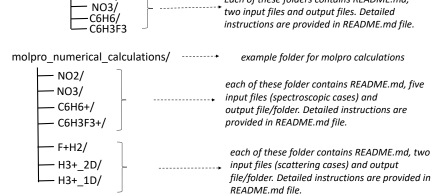


Figure 1: The above directory structure depicts the files and directories present in the parent folder, 'ADT-version0'.

Schematic Outline of Files and Folders in 'test_runs'

test runs/ analytic expressions/ ------ example folder for analytic formulation four state/ each of these folder contains README.md five state/ and one output file. Detailed instructions are six state/ provided in README.md file. eight state/ numerical calculations/ ------ example folder for numerical calculations NO2/ each of these folder contains README.md, three input files (one for adiabatic PESs and two for NO3/ two components of NACTs) and output file/folder. C6H6+/ For NO₃, 'script.py' is provided for script based C6H3F3+/ input. Detailed instructions are provided in README.md file. F+H2/ - H3+ 2D/ this folder contains README.md, two input files [adiabatic PESs and one component of NACTs along the circular coordinate (ϕ)], 'script.py' - H3+ 1D/ and output files/folders. Detailed instructions are provided in README.md file. example folder for geometry optimization optimization/ and hessian calculations for spectroscopic cases example folder for MOLPRO package .MOLPRO/ -----NO2/ Each of these folders contains README.md, NO3/ two input files and output files. Detailed C6H6. instructions are provided in README.md file. C6H3F3 example folder for Gaussian package Gaussian/ NO2/ Each of these folders contains README.md, NO3/ two input files and output files. Detailed C6H6/ C6H3F3 instructions are provided in README.md file.

Gamess/----



..... example folder for Gamess package

Each of these folders contains README.md,

Figure 2: The above directory structure carries the information about the files and subdirectories inside the folder, 'test_runs'.

3 Prerequisites

In order to install and execute the 'ADT' program package successfully, user needs a Fortran 90 compiler and Python (2.7 or 3.x), which are to be installed apriorly on the system. In addition, the following python dependencies are also required to use the package:

- 1. NumPy (v 1.13.0 or above)
- 2. setuptools
- 3. h5py

The first two modules are necessary for execution of this package, whereas the third one i.e. h5py is optional and is required only for HDF5 I/O. Alternatively, one can install these libraries through any python package manager (e.g., pip) or build them from source.

4 Installation

• Setting Fortran compiler for installation:

The 'ADT' program uses OpenMP directives in the Fortran code and to properly install the package, the code has to be compiled correctly with appropriate flags. To do so one has to let the build system know what Fortran compiler to use, during installation. The simplest way to do this is to set the F90 environment variable before installing the package. For instance to set gfortran compiler, run (in Bash):

export F90=gfortran

on the other hand to set Intel Fortran compiler:

export F90=ifort

If no F90 environment variable is specified then the program is installed using gfortran compiler by default. To use other compilers or to fine tune the installation process, modify the setup.py script.

• Installing ADT:

For installing the 'ADT' program package user has to execute the following command in the terminal:

python setup.py install

After compilation of all the necessary files, the package is installed in the system and a command line executable adt is created. By default, the installation is performed in root of the system (usually in /usr/local) and therfore, root privilege may be required. If user is unable to access the root privilege or do not wish to install in the root of system, the 'ADT' package can be installed in a local package folder for python (usually in \sim /local/) by providing the --user flag during installation:

python setup.py install --user

Alternatively, if pip is available, user can install the package by executing the following command inside the source directory:

pip install .

On the contrary, for installing the package at any other location in the system, please visit the site https://setuptools.readthedocs.io/en/latest/easy_install .html#custom-installation-locations. During installation, the Fortran file is compiled to a python extension module by numpy distutils and to further configure the compilation process, user can visit the official NumPy distutils page (https://docs.scipy.org/doc/numpy/reference/distutils.html#module-numpy.distutils).

5 Execution

After installing the 'ADT' package, one has to execute the command line utility, adt to formulate analytic expressions or to perform numerical calculation or MOLPRO interfacing followed by numerical computation of ADT quantities employing the subcommands ana, num and mol, respectively. In order to perform geometry optimization and hessian calculations, user has to execute the subcommands -opt with any one of the packages' names - -mol or -gauss or -gamess. The details of the execution of these different sections are given in the following diagram:

Working Principles and Command-line-arguments for Analytic, Numeric, geometry optimization and Molpro (adiabatic PESs and NACTs) Jobs

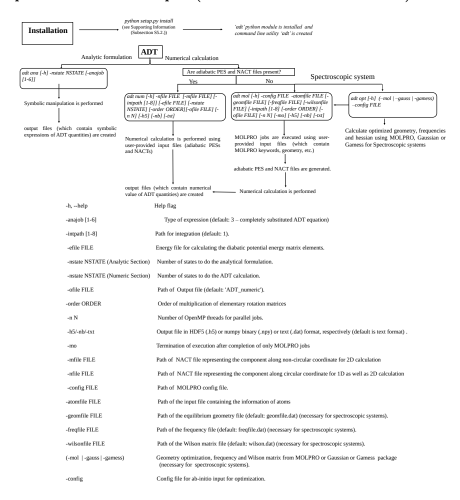


Figure 3: This diagram presents an overall schematic outline of the working principles for analytic expressions generation, numerical calculations for ADT, geometry optimization (only for spectroscopic cases) interfacing Molpro or Gaussian or Gamess and ab-initio calculations for adiabatic PESs and NACTs interfacing molpro. Detailed description of the arguments, -anajob and -intpath are presented in Tables 1 and 2.

5.1. Analytic

For analytic jobs, execute the following command line arguments,

The possible values of -anajob and their significance are presented in the following table:

Table 1: This table depicts the correlation between the possible flags of the keyword argument, -anajob and the desired output

| Keyword Argument | Analytical Quantities |
|------------------|--|
| (-anajob) | |
| 1 | elements of adiabatic to diabatic transformation (ADT) matrix |
| 2 | partially substituted forms of ADT equations [let say, for state=3, $\vec{\nabla}\theta_{12} = -\vec{\tau}_{12} - \sec\theta_{12}\tan\theta_{13}\vec{\tau}_{23} + \tan\theta_{12}\tan\theta_{13}\vec{\nabla}\theta_{13}$,] |
| 3 | complete forms of ADT equations (gradients in right hand side are replaced) [let say, for state=3, $\vec{\nabla}\theta_{12} = -\vec{\tau}_{12} - \sin\theta_{12}\tan\theta_{13}\vec{\tau}_{13} - \cos\theta_{12}\tan\theta_{13}\vec{\tau}_{23}$,] |
| 4 | elements of coefficient matrix of gradient of ADT angles (see Eq. 14 in the article for details) |
| 5 | elements of coefficient matrix of NACTs (see Eq. 14 in the article for details) |
| 6 | elements of diabatic potential energy matrix (detailed expressions) |

Examples:

1. Execute the following command to formulate completely substituted ADT equations for a five electronic state sub-Hilbert space:

```
adt ana -nstate 5 -anajob 3
```

The output file, 'ADT_EQUATIONS_COMPLETE.DAT' contains the analytic expressions of ADT equations.

2. Execute the following command to formulate diabatic potential energy matrix elements for a six electronic state sub-Hilbert space:

```
adt ana -nstate 6 -anajob 6
```

The output file is 'W_MATRIX.DAT'.

3. Execute the following command to formulate coefficient matrix of gradient of ADT angles (Eq. 14, described in the article) for a three electronic state sub-Hilbert space:

```
adt ana -nstate 3 -anajob 4
```

The output file is 'GRADCOEFF.DAT'.

4. Execute the following command to formulate coefficient matrix of NACTs (Eq. 14, described in the article) for a eight electronic state sub-Hilbert space:

```
adt ana -nstate 8 -anajob 5
```

The output file is 'TAUCOEFF.DAT'.

5.2. Numeric

User can carry out numerical calculations by using the subcommand num,

```
adt num [-h] -nfile FILE [-mfile FILE] [-intpath [1-8]]
[-efile FILE] [-nstate NSTATE] [-order ORDER] [-ofile FILE]
[-n N] [-h5] [-nb] [-txt]
optional arguments:
 -h, --help
               show the help message
 -mfile FILE
                 Specify the path of the input NACT file
                 representing the component of NACT
                 for the non-circular coordinate
                 (e.g. \{0,pi/2\} or \{0,pi\} or
                 {0,infinity})
 -intpath [1-8] Specify the path for calculation (default: 1).
 -efile FILE
                 Specify the path of the adiabatic PES file for
                 calculating the diabatic potential energy matrix
                 elements.
 -nstate NSTATE Specify the number of states to do the
                 calculation. By default it includes all
                 the data for calculation.
 -order ORDER
               Write the order of multiplication of
                 the elementary rotation matrices.
 -ofile FILE
                 Specify the path of the output file/folder
                 (w/o extension) (default: 'ADT_numeric').
```

-n N Specify the number of OpenMP threads for parallel

jobs (default: 1).

-h5 Write results in a HDF5 file (.h5).

Fast IO, smaller file size and

hierarchical filesystem-like data format, preferable for saving and sharing large

datasets in an organised way.

-nb Write results in NumPy binary file (.npy).

Preferable when working with NumPy for its

much faster IO and easy portability.

-txt Write results in a text file (default

behaviour).

Required arguments:

-nfile FILE Need to specify the path of the input

NACT file, which is required both for 1D as well as 2D calculation. This file represents the component of NACT for the circular coordinate (e.g. {0,2pi}).

The probable values of -intpath and their significance are depicted in the following table:

Table 2: This table shows the relation between the possible flags of the keyword argument <code>-intpath</code> and the integration paths

| Argument | Path of Integration (R : first coordinate, P : second coordinate) |
|------------|---|
| (-intpath) | |
| 1 | 1. At lowest value of R grid, the integration is performed for each positive step of P grid and the values of ADT angles are stored; |
| | 2. With those values of the variables at each P grid, integration is carried out for each positive increment of R grid |
| 2 | 1. At lowest value of R grid, the integration is performed for each negative step of P grid and the values of ADT angles are stored; |
| | 2. With those values of the variables at each P grid, integration is carried out for each positive increment of R grid |
| 3 | 1. At highest value of R grid, the integration is performed for each positive step of P grid and the values of ADT angles are stored; |
| | 2. With those values of the variables at each P grid, integration is carried out for each negative increment of R grid |
| 4 | 1. At highest value of R grid, the integration is performed for each negative step of P grid and the values of ADT angles are stored; |
| | 2. With those values of the variables at each P grid, integration is carried out for each negative increment of R grid |
| 5 | 1. At lowest value of P grid, the integration is performed for each positive step of R grid and the values of ADT angles are stored; |
| | 2. With those values of the variables at each R grid, integration is carried out for each positive increment of P grid |
| 6 | 1. At lowest value of P grid, the integration is performed for each negative step of R grid and the values of ADT angles are stored; |
| | 2. With those values of the variables at each R grid, integration is carried out for each positive increment of P grid |
| 7 | 1. At highest value of P grid, the integration is performed for each positive step of R grid and the values of ADT angles are stored; |
| | 2. With those values of the variables at each R grid, integration is carried out for each negative increment of P grid |
| 8 | 1. At highest value of P grid, the integration is performed for each negative step of R grid and the values of ADT angles are stored; |
| | 2. With those values of the variables at each R grid, integration is carried out for each negative increment of P grid |

User can also employ the numerical segment of this package as a python module. While carrying out 1D and 2D calculation, one can use two functions, adt1d and adt2d, respectively, which take adiabatic PESs and NACTs as inputs, and return the ADT quantities (ADT angles, residue of ADT angles, ADT matrices and diabatic PESs).

Examples

1. Using the command line executable adt

(i) Execute the following command in terminal to carry out 2D ADT calculation, where 'Tau_Rho.dat', 'Tau_Phi.dat' and 'Adiabatic_PES.dat' contain components of NACTs along non-circular coordinate, components of NACTs along circular coordinate and adiabatic PESs, respectively. The integration is done along path 6.

```
adt num -mfile Tau_Rho.dat -nfile Tau_Phi.dat
-efile Adiabatic_PES.dat -intpath 6
```

While running the command, ADT angles, residues of ADT angles, ADT matrices and diabatic potential energy matrices are computed and those are written in plain text files (default output format) inside a folder, 'ADT_numeric_6'.

(ii) Following command can be used to perform 2D ADT calculation, where 'Tau_Theta.npy', 'Tau_Phi.npy' and 'Adiabatic_PES.npy' contain components of NACTs along non-circular coordinate, components of NACTs along circular coordinate and adiabatic PESs, respectively. The integration is done along path 4.

```
adt num -mfile Tau_Theta.npy -nfile Tau_Phi.npy
-efile Adiabatic_PES.npy -intpath 4 -nb
```

Output files will be saved in NumPy binary format inside a folder, 'ADT_numeric_4'.

(iii) Following command will carry out 2D ADT calculation, where 'Tau_Rho.h5', 'Tau_Phi.h5' and 'Adiabatic_PES.h5' contain components of NACTs along non-circular coordinate, component of NACTs along circular coordinate and adiabatic PESs, respectively. The

integration is done along path 2.

```
adt num -mfile Tau_Rho.h5 -nfile Tau_Phi.h5
-efile Adiabatic_PES.h5 -intpath 2 -h5
```

Output result will be saved as a HDF5 file, 'ADT_numeric_2.h5' and the ADT quantities will be stored as different groups of dataset in that HDF5 file.

(iv) If the 'ADT' package is installed with OpenMP enabled, user can employ -n flag to run the program in parallel threads. The following command executes the program package using 4 OpenMP threads.

```
adt num -mfile Tau_Rho.dat -nfile Tau_Phi.dat -efile Adiabatic_PES.dat -intpath 6 -n 4
```

The input and output file structures are similar as the serial ones.

(v) If one wishes to perform 1D ADT, the following command can be executed with usual I/O files as the previous cases,

```
adt num -nfile Tau_Phi.dat -efile Adiabatic_PES.dat
```

The command will generate the ADT angle, residues of ADT angles, ADT matrices and diabatic potential energy matrices, and store the output in plain text files (default output format) inside a folder, 'ADT_numeric_1D'.

(vi) By default calculations are done with ADT matrix formed using the default order of multiplication of the elementary rotaion matrices $(A^{ij}(\Theta_{ij}))$ where the matrices are multiplied in increasing order of the second index (j) and then, for a fixed second index, in increasing order of the first index (i). If user wants to use any other order of multiplication, the order can be provided using the -order keyword. For example, for a 3 state sub-Hilbert space to form the ADT matrix using the order '12,23,13' i.e.

$$A(\Theta_{12}, \Theta_{23}, \Theta_{13}) = A^{12}(\Theta_{12}) \cdot A^{23}(\Theta_{23}) \cdot A^{13}(\Theta_{13})$$

execute the following command:

```
adt num -nfile Tau_Phi.dat -efile Adiabatic_PES.dat -order 12,23,13
```

The structure of input and output files are same as above.

2. Using python script

(i) **1D:** The following sample python script shows the usage of the 'ADT' package as a python module to calculate 1D ADT. The files 'Tau_Phi.dat' and 'Adiabatic_PES.dat' are the NACT (along the circular coordinate) and adiabatic PES files. As for example, those two files can be taken from the 'ADT_version0/test_runs/numerical_calculation/H3+_1D'.

```
import numpy as np
from adt.numeric import adt_numeric as an

nact = np.loadtxt('Tau_Phi.dat')
enr = np.loadtxt('Adiabatic_PES.dat')

angle, res, amat, db = an.adt1d(nact[:,0], nact[:,1:],
enr[:,1:])
```

(ii) **2D:** This python script shows 2D calculation of ADT quantities along path 6, where the files 'Tau_Rho.npy', 'Tau_Phi.npy', 'Adiabatic_PES.npy' contain components of NACTs along non-circular coordinate, components of NACTs along circular coordinate and adiabatic PESs, respectively. The input files contain *ab initio* data for five electronic manifold spanning 50 × 180 grid points and can be taken from the 'ADT_version0/test_runs /numerical_calculation/NO3'.

```
import numpy as np
from adt.numeric import adt_numeric as an
nact1 = np.load('Tau_Rho.npy')
nact2 = np.load('Tau_Phi.npy')
enr = np.load('Adiabatic_PES.npy')
```

```
grid1 = np.unique(nact1[:,0])
grid2 = np.unique(nact1[:,1])

#If the number of grid points along the first and second
#coordinates are 50 and 180, respectively over 5
#electronic states with 10 NACT elements

enr = enr[:,2:].reshape(50, 180, 5)
nact1 = nact1[:,2:].reshape(50, 180, 10)
nact2 = nact2[:,2:].reshape(50, 180, 10)

#The integration is performed along path 6.

angle, res, amat,db = an.adt2d(grid1, grid2, nact1, nact2, 6, enr)
```

5.3. MOLPRO/ Gaussian/ Gamess: Optimization

In spectroscopic cases, inputs files such as, equilibrium, frequency and wilson files can be directly obtained by optimization segment of ADT code, where geometry optimization and subsequent frequency calculations are preformed by quantum chemistry packages, namely, MOLPRO, Gaussian and Gamess. The detailed discussion is given in Section 7 and then, the output files (equigeom.dat, wilson.dat, frequency.dat) can be further utilized for computing adiabatic PESs and NACTs in Molpro section of ADT package(see Figure 3).

```
adt opt [-h] (-mol | -gauss | -gamess) -config FILE

optional arguments:

-h, --help show the help message

-mol ab-initio calculations are performed by MOLPRO package
```

| -gauss | ab-initio calculations are performed by Gaussian package |
|--------------|--|
| -gamess | ab-initio calculations are performed by Gamess package |
| -config FILE | the configuration files for respective quantum chemistry packages containing the necessary keywords of geometry optimization and hessian inputs. |

For optimization, user has to supply two inputs files:

- 1. configuration file: contains ab-initio keywords for optimization (like memory, basis set, method of calculation, etc.), system information (initial geometry file name)
- 2. initial geometry file: contains initial geometry in XYZ format (in Å).

Examples

1. By MOLPRO package, user can run following command in terminal

```
adt opt -mol -config mol.config
```

where, 'mol.config' is MOLPRO config file. This will execute optimization by Gaussian package to calculate the optimized geometry in 'XYZ' format, frequency and wilson file.

2. optimization by Gaussian package, one can run the following command in terminal,

```
adt opt -gauss -config gauss.config
```

where 'gauss.config' is the Gaussian configuration file.

3. Similarly, Gamess can be invoked for geometry optimization by executing following command

```
adt opt -gamess -config gms.config
```

where, gms.config is config file for ab-initio input of Gamess package.

Limitations

• The ground state geometry optimization and hessian calculations are interfaced with only state-of-the-art single reference quantum chemistry methods such as HF, UHF, ROHF, MP2, UMP2, CCSD, CCSD(T), UCCSD, UCCSD(T) and DFT method.

5.4. Molpro: Adiabatic PESs and NACTs

On the other hand, user can take the benefit of molpro section, where *ab initio* electronic structure calculations are performed according to the user provided input files (discussed in detail in Section 8) and then, the outputs (adiabatic PESs and NACTs) are used for subsequent ADT calculation (see Figure 3). User has to execute the following command,

```
adt mol [-h] -config FILE -atomfile FILE [-geomfile FILE]
[-freqfile FILE] [-wilsonfile FILE] [-intpath [1-8]] [-order ORDER]
[-ofile FILE] [-n N] [-mo] [-h5] [-nb] [-txt]
optional arguments:
 -h, --help
                 show the help message
 -geomfile FILE Specify the geometry file containing the initial
                  grid point in 'xyz' format (in Angstrom)
                  (default: geomfile.dat).
                  (Ignore for scattering system).
 -freqfile FILE Specify the frequency information file, where
                  frequencies of normal modes are written in cm-1
                  (default: frequency.dat).
                  (Ignore for scattering system)
 -wilsonfile FILE Specify the filename containing the Wilson matrix
                  of a molecular species (default: wilson.dat).
                  (Ignore for scattering system).
 -intpath [1-8] Specify the path for calculation (default: 1).
```

-order ORDER Write the order of multiplication of the elementary rotation matrices. -ofile FILE Specify the path of the output file/folder (w/o extension) (default: 'ADT_numeric'). -n N Specify the number of OpenMP threads for parallel jobs (default: 1). Terminate the execution only after completion -mo of MOLPRO jobs -h5 Write results in a HDF5 file (.h5). Fast IO, smaller file size and hierarchical filesystem-like data format, preferable for saving and sharing large datasets in an organised way. Write results in NumPy binary file (.npy). -nb Preferable when working with NumPy for its much faster IO and easy portability. Write results in a text file. (default -txt behaviour)

Required arguments:

-config FILE Specify the MOLPRO configuration file containing the necessary keywords of MOLPRO software.

-atomfile FILE Specify the information file constituting atomic symbols and atomic masses.

For spectroscopic cases, user has to supply five input files:

1. MOLPRO configuration file: contains MOLPRO keywords (like memory, basis set, method of

calculation, etc.), system information (like symmetry, participating normal modes) and grid information

- 2. atom information file: contains atomic symbols and atomic masses
- 3. equilibrium geometry file: contains equilibrium geometry in xyz format (in Å)
- 4. frequency file: contains normal mode frequencies (in cm⁻¹)
- 5. wilson matrix file: contains mass-weighted second derivative matrix (Wilson matrix).

On the other hand, if user wishes to perform *ab initio* calculation of adiabatic PESs and NACTs for scattering systems, the first two files, i.e. MOLPRO configuration and atom information files are necessary and sufficient. The details of the above files are given in Section 8.

In order to store the geometry and MOLPRO output files for every successful and unsuccessful jobs, two folders, 'CompleteJobs' and 'IncompleteJobs' are generated, respectively during the execution of molpro section. One can use -mo argument to terminate the execution after completion of MOL-PRO jobs to check the convergence and smoothness of adiabatic PESs and NACTs. Subsequently, user can remove/refine unphysical data points prior to ADT calculation (using the numeric section).

Examples

1. For scattering case, one can run the following command in terminal,

```
adt mol -config molpro.config -atomfile atomfile.dat
-intpath 6
```

where 'molpro.config' and 'atomfile.dat' are the MOLPRO configuration file and atom information file, respectively. This will execute MOLPRO to calculate the *ab initio* data (adiabatic PESs and NACTs) over the specified grid points defined in the configuration file and subsequently, carry out the ADT calculation along path 6.

2. In order to terminate the program after running the MOLPRO jobs (before calculating the ADT quantities), one can pass the -mo argument,

```
adt mol -config molpro.config -atomfile atomfile.dat -mo
```

3. For spectroscopic case, user has to run the following command in terminal,

```
adt mol -config molpro.config -atomfile atomfile.dat
-geomfile geomfile.dat -freqfile frequency.dat
-wilsonfile wilson.dat -intpath 4
```

where 'geomfile.dat', 'frequency.dat' and 'wilson.dat' are the equilibrium geometry file, frequency file and Wilson matrix file, respectively. Other two files, 'molpro.config' and 'atomfile.dat' are the same as discussed above.

Limitations

- non-convergence of *ab initio* points originating from complete active space self-consistent field (CASSCF)/multi-reference configuration interaction (MRCI) methodology: The interpolation function in the python code cannot improve the output data if a large number of nuclear geometries are unrealistic.
- The swapping of *ab initio* computed NACTs due to the interchange of electronic states at CI point(s) are not considered in this software package due to molecule-specific characteristics.
- The *ab initio* adiabatic PESs can be computed only in mrci approach while calculating NACTs in finite difference method (ddr) (see Section 8.1).

6 Instructions for Input Files of Numeric Section

One can employ the numeric section of 'ADT' program package to solve the ADT equations along one dimensional as well as two dimensional contours of nuclear CS using adiabatic PES and NACT files. Those files can be provided either in HDF5 (.h5) or NumPy binary (.npy) or in text format. Detailed instructions related to the preparation of input files are given in the following paragraphs.

User can choose any one of the following coordinate systems to carry out ADT calculation:

- Spectroscopic systems:
 - (i) 2D ADT calculation: plane polar counterparts (ρ and ϕ) of a pair of normal modes $(Q_i = \rho \cos \theta \text{ and } Q_j = \rho \sin \theta)$

- (ii) 1D ADT calculation: circular coordinate, ϕ for a fixed ρ
- Scattering systems:
 - 1. 2D ADT calculation:
 - (a) hyperspherical coordinate (ρ and ϕ for a fixed θ or θ and ϕ for a fixed ρ)
 - (b) Jacobi coordinate $(q \text{ and } \phi)$ (see Figure 4)
 - 2. 1D ADT calculation: circular coordinate, ϕ for a fixed ρ and θ (hyperspherical) or ϕ for a fixed q (Jacobi).

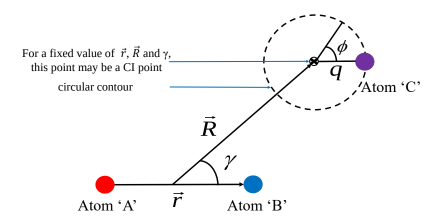


Figure 4: The above diagram briefly manifests the Jacobi coordinates of a triatomic system. The diatomic vector between Atom 'A' and Atom 'B' is represented as \vec{r} with distance r, the vector from diatomic center of mass to the third atom 'C' is symbolized as \vec{R} with distance R, and the angle between them is designated as γ . The circular contour is characterized by a radius (q) and an angle (ϕ) along which ab initio calculation is performed. In this process, two Jacobi coordinates $(R \text{ and } \gamma)$ are varying at a fixed value of r while performing such type of calculation.

6.1. Input files for 2D calculation

For the numeric section, three input files, namely, the adiabatic PES and components of NACTs are necessary.

• Input file for adiabatic PESs: The adiabatic PES input file must contain 2 + N number of columns (N is number of coupled electronic states): first two columns represent grid points

of non-circular and circular coordinates, respectively and the rest of the columns contain the adiabatic PESs in increasing order of electronic states. User can supply scaled adiabatic energies for diabatization (scaled with respect to ground state minimum energy or asymptotic energy).

Example: The following table (Table 3) represents the arrangement of input file data points for adiabatic PESs of three electronic states (N = 3).

Table 3: Sample input file for adiabatic PESs

| * * | | | | | | | |
|--------------------------------------|----------|---|-----------------------|------------------------|--|--|--|
| First Coordinate Second Coordinate | | Second Coordinate Adiabatic Energy Adiabatic Energy | | Adiabatic Energy | | | |
| | | (ground state) | (first excited state) | (second excited state) | | | |
| 0.01 | 0.000000 | 0.002 | 0.123 | 0.125 | | | |
| 0.01 | 3.141592 | 0.003 | 0.125 | 0.126 | | | |
| 0.01 | 6.283185 | 0.003 | 0.124 | 0.126 | | | |
| | | | | | | | |
| 0.02 | 0.000000 | -0.001 | 0.119 | 0.121 | | | |
| 0.02 | 3.141592 | -0.002 | 0.118 | 0.121 | | | |
| 0.02 | 6.283185 | -0.002 | 0.118 | 0.122 | | | |

• Input files for NACTs: Two NACT files (-mfile and -nfile) are required containing the components of NACTs for the non-circular and circular coordinates, respectively. There should be 2 + N * (N - 1)/2 number of columns in each file, where the first two are non-circular and circular grids, respectively and the remaining ones signify one of the component of τ values within N * (N - 1)/2 pairs of electronic states. NACT components are represented as τ_{ρ}^{ij} or τ_{θ}^{ij} or τ_{q}^{ij} and τ_{ϕ}^{ij} . The columns of NACTs in the input files should be arranged in increasing order of first index (i) for a fixed second index (j) subjecting to the condition i < j and then, by increasing the second index.

Example: For four coupled electronic states, elements of τ_{ρ} can be ordered as τ_{ρ}^{12} , τ_{ρ}^{13} , τ_{ρ}^{23} , τ_{ρ}^{14} , τ_{ρ}^{24} and τ_{ρ}^{34} as one of the input files (Table 4), where ϕ component of NACT will be given as τ_{ϕ}^{12} , τ_{ϕ}^{13} , τ_{ϕ}^{23} , τ_{ϕ}^{14} , τ_{ϕ}^{24} and τ_{ϕ}^{34} for the second input file of NACT (Table 5).

Table 4: Sample input file for ρ component of NACTs

| ρ | φ | $	au_{ ho}^{12}$ | $	au_{ ho}^{13}$ | $	au_{ ho}^{23}$ | $	au_{ ho}^{14}$ | $	au_{ ho}^{24}$ | $	au_{ ho}^{34}$ |
|------|----------|------------------|------------------|------------------|------------------|------------------|------------------|
| 0.01 | 0.000000 | 0.002 | 0.123 | 0.125 | 0.020 | 0.103 | 0.105 |
| 0.01 | 3.141592 | 0.003 | 0.125 | 0.126 | 0.021 | 0.105 | 0.101 |
| 0.01 | 6.283185 | 0.003 | 0.124 | 0.126 | 0.023 | 0.107 | 0.100 |
| | | | | | | | |
| 0.02 | 0.000000 | -0.001 | 0.119 | 0.121 | 0.019 | 0.109 | 0.100 |
| 0.02 | 3.141592 | -0.002 | 0.118 | 0.121 | 0.023 | 0.108 | 0.101 |
| 0.02 | 6.283185 | -0.002 | 0.118 | 0.122 | 0.025 | 0.108 | 0.102 |

Table 5: Sample input file for ϕ component of NACTs

| ρ | ϕ | $	au_\phi^{12}$ | $	au_\phi^{13}$ | $	au_\phi^{23}$ | $	au_\phi^{14}$ | $	au_\phi^{24}$ | $	au_{\phi}^{34}$ |
|------|----------|-----------------|-----------------|-----------------|-----------------|-----------------|-------------------|
| 0.01 | 0.000000 | 0.005 | 0.103 | 0.150 | 0.020 | 0.130 | 0.121 |
| 0.01 | 3.141592 | 0.007 | 0.105 | 0.156 | 0.030 | 0.133 | 0.128 |
| 0.01 | 6.283185 | 0.009 | 0.107 | 0.156 | 0.030 | 0.137 | 0.130 |
| | | | | | | | |
| 0.02 | 0.000000 | -0.006 | 0.105 | 0.155 | 0.021 | 0.133 | 0.121 |
| 0.02 | 3.141592 | -0.009 | 0.107 | 0.159 | 0.027 | 0.137 | 0.125 |
| 0.02 | 6.283185 | -0.011 | 0.108 | 0.160 | 0.029 | 0.139 | 0.127 |

6.2. Input Files for 1D Calculation

For 1D ADT calculation, one should supply two input files, namely, the adiabatic PESs and angular component of NACT (τ_{ϕ}) forming closed contour. There must be 1 + N and 1 + N * (N - 1)/2 number of columns (N is number of coupled electronic states) in the adiabatic PES and NACT input files, respectively. The first column contains the grid points of the circular coordinate and the rest of the columns are exactly similar as the 2D input files for ADT calculation.

7 Instructions for Input Files of optimization

In ADT, geometry optimization and hessian calculations are interfaced with three quantum chemistry packages, MOLPRO, Gaussian and Gamess. User has to provide two input files with the flags -opt and -mol or -gauss or -gamess (quantum chemistry package's name). The details of config and geometry files are depicted below:

7.1. MOLPRO Configuration File

The MOLPRO configuration file, namely, 'mol.config' is standard INI configuration file, which contains two sections, namely, optInfo, gInfo to specify all necessary keywords to run MOLPRO.

- optInfo contains the following seven keywords:
 - memory: allotted memory for MOLPRO job
 - processor (optional): number of processors used for parallelization (default is 1)
 - method: specify electronic structure method, namely any single reference method such as
 MP2, CCSD, CCSD(t) or its unrestricted versions and DFT method.
 - basis: basis set keyword compatible with MOLPRO input.
 - spin: spin multiplicity (2s) of a molecular species
 - charge: charge of a molecular species
 - symmetry(optional): any Abelian point group of the molecular species during *ab initio* calculation (default value is nosym).
- gInfo contains the following keyword:
 - file: name of initial geometry file. File must contain the first column as atom symbol and second column as the Cartesian coordinates of the atoms (in Å) in a $n \times 3$ matrix form for a n atomic molecule.

The above description of 'mol.config' is presented in Table 6.

Table 6: Description and possible inputs of MOLPRO configuration file ('mol.config')

| Sections | Keywords | Description | | | | |
|---|---|--|--|--|--|--|
| | memory | memory required for the MOLPRO job, e.g., 10,m | | | | |
| | processor | number of processors required for parallelization of the job, e.g., 1 | | | | |
| +T | method | electronic structure method for geometry optimization and hessian calculation | | | | |
| optInfo | | e.g. UMP2, B3LYP, etc. | | | | |
| | basis | basis for geometry optimization, namely: cc-pVDZ or 6-31G** | | | | |
| | spin | spin multiplicity $(2s)$ of ground state of molecular system , namely: 1 for NO $_3$ radical | | | | |
| | charge of molecular system, namely: 1 for $C_6H_3F_3^+$ | | | | | |
| symmetry point group of molecular system (default value | | point group of molecular system (default value: nosym) | | | | |
| gInfo | file | file name of initial geometry file e.g., geom.dat | | | | |

Examples:

1. The 'mol.config' file shows important inputs for NO_3 radical, where geometry optimization and hessian calculation are performed at DFT/B3LYP with basis set cc-pVDZ (see Table 7):

#Configuration file for the optimization and frequency calculation by MOLPRO job of NO3 radical

```
#Info for running MOLPRO
[optInfo]
memory = 50,m

#for parallel MOLPRO job
processor = 1

#Info for running optimization and hessian
method = b3lyp
basis = cc-pvdz
spin = 2
charge = 0
```

```
symmetry = nosym
#Info of the molecular species
[gInfo]
file = geom.dat
```

2. The format of initial geometry file ('geom.dat') is as follows:

| N | 0.00000000 | 0.00000000 | 0.00000000 |
|---|-------------|-------------|------------|
| 0 | 1.23680091 | 0.00000000 | 0.00000000 |
| 0 | -0.61840019 | -1.07110018 | 0.00000000 |
| 0 | -0.61840019 | 1.07110018 | 0.00000000 |

Table 7: Description of 'mol.config' for NO₂ radical, NO₃ radical, C₆H₆ and 1,3,5-C₆H₃F₃ for the test runs

| Sections | Keywords | NO_2 radical NO_3 radical C_6H_6 1,3,5- C_6 | | | $1,3,5$ - $C_6H_3F_3$ | |
|----------|-----------|---|---|-------|-----------------------|--|
| | memory | 50,m | | | | |
| | processor | 2 | | | | |
| t T f . | method | method UCCSD B3LYP B3I | | B3LYP | MP2 | |
| optInfo | basis | | | | | |
| | spin | 2 | | 0 | | |
| | charge | | 0 | | | |
| gInfo | file | geom.dat | | | | |

7.2. Gaussian Configuration File

The Gaussian configuration file, namely, 'gauss.config' is a standard INI configuration file, which contains two sections, namely, optInfo, gInfo to specify all necessary keywords to run Gaussian.

- optInfo contains the following eight keywords:
 - version (optional): version of Gaussian package (default is g16)
 - memory: allotted memory for Gaussian job

- processor (optional): number of processors used for parallelization (default is 1)
- method: specify electronic structure method, namely any single reference method such as
 MP2, CCSD, CCSD(t) or its unrestricted versions and DFT method
- basis: basis set keyword compatible with Gaussian input.
- spin: spin multiplicity (2s+1) of a molecular species
- charge: charge of a molecular species
- symmetry(optional): any Abelian point group of the molecular species during *ab initio* calculation (default value is nosym).
- gInfo contains the following keyword:
 - file: name of initial geometry file. File must contain the first column as atom symbols and second column as Cartesian coordinates of the atoms (in Å) in a $n \times 3$ matrix form for a n atomic molecule.

The above description of 'gauss.config' is presented in Table 8.

Table 8: Description and possible inputs of Gaussian configuration file ('gauss.config')

| Sections | Keywords | Description | | | | |
|----------|-----------|--|--|--|--|--|
| | version | version of Gaussian package, namely: g16 or g09 | | | | |
| | memory | memory required for the Gaussian job, e.g., 1GB | | | | |
| ant Infa | processor | number of processors required for parallelization of the job, e.g., 1 | | | | |
| optInfo | method | electronic structure method for geometry optimization and hessian calculation | | | | |
| | | e.g. UMP2, B3LYP, etc. | | | | |
| | basis | basis for geometry optimization, namely: cc-pVDZ or 6-31G** | | | | |
| | spin | spin multiplicity $(2s+1)$ of ground state of molecular system , namely: 2 for NO $_2$ radical | | | | |
| | charge | charge of molecular system, namely: 1 for $C_6H_6^+$ | | | | |
| | symmetry | point group of molecular system (default value: nosymm) | | | | |
| gInfo | file | name of initial geometry file e.g., geom.dat | | | | |

Examples:

1. The following sample 'gauss.config' file presents necessary inputs for NO₂ radical, where geometry optimization and hessian calculation are performed at UCCSD with basis set cc-pVDZ (see Table 9):

```
#Configuration file for the optimization and frequency calculation by Gaussian job of NO2
#Info for running Gaussian
[optInfo]
version = g16
memory = 1GB
#for parallel Gaussian job
processor = 2
#Info for running optimization and hessian
method = uccsd
basis = cc-pvdz
spin = 2
charge = 0
symmetry = nosymm
#Info of the molecular species
[gInfo]
file = geom.dat
```

2. The format of initial geometry file ('geom.dat') is as follows:

```
N 0.00000000 0.00000000 -0.324503807

D 0.00000000 1.104583330 0.142043685

D 0.00000000 -1.104583330 0.142043685
```

Table 9: Description of 'gauss.config' for NO_2 radical, NO_3 radical, C_6H_6 and 1,3,5- $C_6H_3F_3$ for the test runs

| Sections | Keywords | NO ₂ radical | NO ₃ radical | C_6H_6 | $1,3,5-C_6H_3F_3$ | |
|----------|-----------|-------------------------|-------------------------|----------|-------------------|--|
| T. f. | version | g16 | | | | |
| | memory | 1GB | | | | |
| | processor | 2 | | | | |
| optInfo | method | UCCSD | B3LYP | B3LYP | MP2 | |
| | basis | cc-pvdz | | | | |
| | spin | 2 | | | | |
| | charge | 0 | | | | |
| | symmetry | nosymm | | | | |
| gInfo | file | geom.dat | | | | |

7.3. Gamess Configuration File

The Gamess configuration file, namely, 'gms.config' is a standard INI configuration file, which contains two sections, namely, optInfo, gInfo to specify all necessary keywords to run Gamess.

- optInfo contains the following nine keywords:
 - path (optional): Path of Gamess script (default is rungms)
 - memddi: allotted memory for distributed data interface (DDI) in Gamess run
 - memory: allotted memory for Gamess job
 - processor (optional): number of processors used for parallelization (default is 1)
 - method: specify electronic structure method, namely any single reference method such as
 MP2, CCSD, CCSD(t) or its unrestricted versions and DFT method
 - basis: basis set keyword compatible with Gamess input.
 - spin: spin multiplicity (2s+1) of a molecular species
 - charge: charge of a molecular species
 - symmetry(optional): any Abelian point group of the molecular species consistent with
 Gamess input (default value is C₁).

- gInfo contains the following keyword:
 - file: name of initial geometry file. File must contain the first column as atom symbol and second column as the atomic number of atoms and third column as Cartesian coordinates of the atoms (in Å) in a $n \times 3$ matrix form for a n atomic molecule.

The above description of 'gms.config' is presented in Table 10.

Table 10: Description and possible inputs of Gamess configuration file ('gms.config')

| Sections | Keywords | Description | | | |
|----------|-----------|--|--|--|--|
| | path | path of Gamess script, e.g., /home/user/gamess/rungms | | | |
| | memddi | memory required for DDI storage, e.g., 1 | | | |
| ontInfo | memory | memory required for the Gamess job, e.g., 10 | | | |
| optInfo | processor | number of processors required for parallelization of the job, e.g., 1 | | | |
| | method | electronic structure method for geometry optimization and hessian calculation | | | |
| | | e.g. UMP2, B3LYP, etc. | | | |
| | basis | basis for geometry optimization compatible with Gamess input, e.g., ccd or accd | | | |
| | spin | spin multiplicity $(2s+1)$ of ground state of molecular system , namely: 2 for NO ₂ | | | |
| | charge | charge of molecular system, namely: 1 for $C_6H_3F_3^+$ | | | |
| | symmetry | point group of molecular system in Gamess's input, e.g., C1 or Cnv 2 | | | |
| gInfo | file | name of initial geometry file e.g., geom.dat | | | |

Examples:

1. The 'gms.config' file shows important inputs for C₆H₆ molecule, where geometry optimization and hessian calculation are performed at DFT/B3LYP method with basis set cc-pVDZ (see Table 11):

#Configuration file for the optimization and frequency calculation by Gamess job of C6H6

```
[optInfo]
# path of Gamess script
path = /home/bijit/SATYAM/gamess/rungms
```

```
# memory required to Gamess job
memodi = 30
memory = 50

#for parallel Gamess job
processor = 1

#Info for running optimization and hessian
method = b3lyp
basis = ccd
spin = 1
charge = 0

[gInfo]
#Info for geometry file name
file = geom.dat
```

2. The format of initial geometry file ('geom.dat') is as follows:

| C | 6 | 1.3820899072 | 0.000000000 | 0.0 |
|---|---|---------------|---------------|-----|
| C | 6 | 0.6910449531 | -1.1969249688 | 0.0 |
| C | 6 | -0.6910449531 | -1.1969249688 | 0.0 |
| C | 6 | -1.3820899072 | 0.000000000 | 0.0 |
| C | 6 | -0.6910449531 | 1.1969249688 | 0.0 |
| C | 6 | 0.6910449531 | 1.1969249688 | 0.0 |
| Н | 1 | 2.4568717910 | 0.000000000 | 0.0 |
| Н | 1 | 1.2284358943 | -2.1277133837 | 0.0 |
| Н | 1 | -1.2284358943 | -2.1277133837 | 0.0 |
| Н | 1 | -2.4568717910 | 0.000000000 | 0.0 |
| Н | 1 | -1.2284358943 | 2.1277133837 | 0.0 |
| Н | 1 | 1.2284358943 | 2.1277133837 | 0.0 |

Table 11: Description of 'gms.config' for NO_2 radical, NO_3 radical, C_6H_6 and 1,3,5- $C_6H_3F_3$ for the test runs

| Sections | Keywords | NO ₂ radical | NO ₃ radical | C_6H_6 | $1,3,5$ - $C_6H_3F_3$ | |
|----------|-----------|--------------------------|-------------------------|----------|-----------------------|--|
| | path | /home/user/gamess/rungms | | | | |
| | memddi | 30 | 30 | 30 | 30 | |
| ontInfo | memory | 50 | 100 | 50 | 50 | |
| optInfo | processor | 1 | | | | |
| | method | UCCSD | B3LYP | B3LYP | MP2 | |
| | basis | ccd | | | | |
| | spin | 2 | | 1 | | |
| | charge | 0 | | | | |
| gInfo | file | geom.dat | | | | |

8 Instructions for Input Files of Molpro Section for adiabatic PESs and NACTs

In order to interface MOLPRO by the software package, 'ADT' for calculation of energy and NACTs calculations, user has to provide five input files for spectroscopic problems and two input files for scattering cases. The MOLPRO configuration and the atom information files are common for both cases. The details of such files are depicted below:

8.1. MOLPRO Configuration File

The MOLPRO configuration file, namely, 'molpro.config' is a standard INI configuration file, which contains six sections, namely, molInfo, sysInfo, eInfo, mInfo, mInfo and gInfo to specify all necessary keywords to run MOLPRO.

- molInfo contains the following three keywords:
 - scrdir (optional): path of scratch directory required during MOLPRO execution and storage of wavefunction (.wfu) files.

- memory (optional): allotted memory for MOLPRO job
- processor (optional): number of processors used for parallelization (default is 1)
- sysInfo contains the following keyword:
 - type: type of the system for electronic structure calculation (spec for spectroscopic systems in normal mode coordinates, scat_hyper for scattering systems in hyperspherical coordinates and scat_jacobi for scattering cases in Jacobi coordinates)
 - symmetry (optional): any Abelian point group of the molecular species during *ab initio* calculation (default value is nosym).
- eInfo contains the following keywords for adiabatic PESs calculation:
 - method: method of adiabatic PESs calculation, namely, multi or mrci
 - basis: basis set
 - cas: active space
 - electron: number of electrons
 - spin: spin multiplicity (2s) of a molecular species
 - charge: charge of a molecular species
 - state: number of electronic states
 - scale(optional): nuclear geometry at which the ground adiabatic PES is set to zero.
 - * spectroscopic cases: ρ, ϕ
 - * scattering hyperspherical cases: ρ, θ, ϕ
 - * scattering Jacobi cases : r,R,γ
 - restricted (optional): type of the initial hartree-fock calculation. true for rhf and false for uhf, default is false
 - uhf_extra (optional): any extra keyword for initial Hartree-Fock calculation
 - multi_extra (optional): extra keywords for multi calculation
 - mrci_extra (optional): extra keywords for mrci calculation
- nInfo contains necessary keywords to calculate NACTs:

- method: method for NACT calculation, namely, cpmcscf or ddr
- basis: basis set
- nact_extra (optional): extra keywords regarding the NACT calculation.
- mInfo (only for spectroscopic cases) contains the following keywords:
 - varying: indexes of the variable normal modes. The indexes are assigned to the normal modes in the ascending order of the corresponding frequencies provided in the frequency file. [not similar as the standard nomenclature, e.g., indexes of Q_{4x} and Q_{4y} normal modes (degenerate components of asymmetric bending mode) for NO₃ radical will be 1,2 as the corresponding frequencies acquire the lowest values.]
- gInfo contains the following keywords:
 - For normal mode coordinates:
 - ▶ rho: ρ_{min} , ρ_{max} , $d\rho$. The grid is defined from ρ_{min} to ρ_{max} in stepsize $d\rho$, where $0 \le \rho_{min} < \rho_{max}$ and $d\rho > 0$. ρ is expressed in mass-weighted dimensionless coordinates.
 - ▶ phi: ϕ_{min} , ϕ_{max} , $d\phi$. The grid is defined from ϕ_{min} to ϕ_{max} in stepsize $d\phi$, where $\phi_{min} = 0$, $\phi_{max} = 2\pi$ and $d\phi > 0$. ϕ is in degrees.
 - ▶ drho (dphi): geometry increment in ρ (ϕ) coordinate used for finite difference DDR method for NACT calculation. It is not required for NACT calculation using coupled-perturbed multi-configuration self-consistent field (CP-MCSCF) method.
 - For hyperspherical coordinate:
 - ▶ rho: ρ_{min} , ρ_{max} , $d\rho$. The grid is defined from ρ_{min} to ρ_{max} in stepsize $d\rho$, where $0 < \rho_{min} < \rho_{max}$ and $d\rho > 0$. ρ is expressed in atomic units.
 - ▶ theta: θ_{min} , θ_{max} , $d\theta$. The grid is defined from θ_{min} to θ_{max} in stepsize $d\theta$, where $0 < \theta_{min} < \theta_{max}$ and $d\theta > 0$. θ is in degrees.
 - ▶ phi: ϕ_{min} , ϕ_{max} , $d\phi$. The grid is defined from ϕ_{min} to ϕ_{max} in stepsize $d\phi$, where $\phi_{min} = 0$, $\phi_{max} = 2\pi$ and $d\phi > 0$. ϕ is in degrees.
 - ▶ drho (dtheta/dphi): geometry increment in ρ (θ/ϕ) coordinate used for finite difference DDR method for NACT calculation. It is not required for NACT calculation using CP-MCSCF method.

- For Jacobi coordinate:
 - \blacktriangleright small_r: r (diatom bond length) in atomic units.
 - ightharpoonup capital_r: R (distance of the third atom from the center of the diatom) in atomic units.
 - ▶ gamma: γ (angle between \vec{r} and \vec{R} in degrees).
 - ▶ q: q_{min} , q_{max} , dq. The grid is defined from q_{min} to q_{max} in stepsize dq, where $0 < q_{min} < q_{max}$ and dq > 0. q is expressed in atomic units. One can provide a single value of q to calculate 1D PECs and NACTs along the ϕ grid.
 - ▶ phi: ϕ_{min} , ϕ_{max} , $d\phi$. The grid is defined from ϕ_{min} to ϕ_{max} in stepsize $d\phi$, where $\phi_{min} = 0$, $\phi_{max} = 2\pi$ and $d\phi > 0$. ϕ is in degrees.
 - ▶ dq (dphi): geometry increment in q (ϕ) coordinate used for finite difference DDR method for NACT calculation. It is not required for NACT calculation using CP-MCSCF method.

The above illustration of 'molpro.config' is compactly presented in Table 12.

Examples:

1. The following sample 'molpro.config' file presents necessary inputs for NO₂ radical, where adiabatic PESs are evaluated in CASSCF approach and analytic NACTs are calculated in CP-MCSCF method (see Table 13):

```
#Configuration file for the MOLPRO job of NO2
#Info for running molpro
[molInfo]
scrdir = /tmp/adtprogram
memory = 10, m
#for parallel molpro job
processor = 2
#Info of the molecular species
[sysInfo]
type = spec
#C_s point group (yz is plane of symmetry)
symmetry = x
#Info for energy calculation
[eInfo]
method = multi
basis = {spdf,n,cc-pvtz;spdf,o,cc-pvtz}
      = occ,11,3;closed,3,0
electron = 23
spin = 1
charge = 0
state = 2,0
#defining extra keyword required to run molpro for this system
#format should be same as in MOLPRO
multi_extra = maxiter,40
```

```
#defining scaling geometry for the adiabatic PESs
#ground state minima of NO2 w.r.t normal modes
scale = 0.0, 0.0
#Info for NACT calculation
[nInfo]
method = cpmcscf
basis = {spdf,n,cc-pvtz;spdf,o,cc-pvtz}
#Info for varying modes
#put the indexes of the normal modes that will be varying
#The following two numbers represent Q_1 and Q_3 normal
#modes (bending and asymmetric stretching modes)
[mInfo]
varying = 1,3
#Put grid in format of <start>, <end>, <step>
#for spectroscopic put the 2D grid as rho and phi
[gInfo]
rho = 0.2, 12.0, 0.2
phi = 0, 360, 1
#Qx = rho*cos(phi) and Qy = rho*sin(phi)
```

2. The following sample 'molpro.config' file represents necessary inputs for H_3^+ (1D PECs and NACTs calculation in Jacobi coordinate using C_s symmetry), where adiabatic energies are calculated in MRCI level of theory and analytic NACTs are evaluated in CP-MCSCF method (see Table 14):

```
#Configuration file for H3+ in Jacobi coordinates
#Info for running molpro
[molInfo]
scrdir = /tmp/adtprogram
memory = 10,m
```

```
#for parallel molpro job
processor = 2
#Info of the molecular species
[sysInfo]
type = scat_jacobi
#C_s point group (yz is plane of symmetry)
symmetry = x
#Info for energy calculation
[eInfo]
method = mrci
basis = cc-pv5z
cas = occ,10,1;closed,0,0;
electron = 2
spin = 0
charge = 1
state = 3,0
#Here scaling of PECs is not necessary as Jacobi coordinate
#is used only for locating the conical intersections (CIs)
#small_r = 1.4, capital_r = 10.0, gamma = 90.0
#minima:H2 at the assymptote of H2+H+
\#scale = 1.4,10.0,90.0
#defining extra keyword required to run molpro for this system
#format should be same as in MOLPRO
restricted = true
uhf_extra = wf,2,1,0,1;
multi_extra = maxiter,40
#Info for NACT calculation
#method should be either ddr or cpmcscf
[nInfo]
method = cpmcscf
```

```
basis = 6-311++G**

#put a fixed value of small_r, capital_r, gamma and q
#and provide a grid of phi
[gInfo]
small_r = 2.0
capital_r = 1.732050808
gamma = 90
#q must be smaller than capital_r
q = 0.5
#Put grid in format of <start>, <end>, <step>
phi = 0, 360,3
```

3. The following sample 'molpro.config' file depicts necessary inputs for H_3^+ (2D ADT calculation in hyperspherical coordinate using C_s symmetry), where adiabatic PESs are evaluated in MRCI level of theory and NACTs are computed in DDR method (see Table 14):

```
#Configuration file for H3+ in hyperspherical coordinates
```

```
#Info for running molpro
[molInfo]
scrdir = /tmp/adtprogram
memory = 10,m

#for parallel molpro job
processor = 2

#Info of the molecular species
[sysInfo]
type = scat_hyper
#C_s point group (yz is plane of symmetry)
symmetry = x

#Info for energy calculation
[eInfo]
```

```
method = mrci
basis = cc-pv5z
cas = occ, 10, 1; closed, 0, 0
electron = 2
spin = 0
charge = 1
state = 3,0
#defining extra keyword required to run molpro for this system
#format should be same as in MOLPRO
restricted = true
uhf_extra = wf,2,1,0,1;
multi_extra = maxiter,40;ACCURACY,GRADIENT=1.d-4,
ENERGY=1.d-6,STEP=1.d-2
#defining scaling geometry for the adiabatic PESs
#ground state minima of H3+
#rho = 1.65, theta = 0.0, phi = 0.0 (D3h point)
scale = 1.65, 0.0, 0.0
#Info for NACT calculation
#method should be either ddr or cpmcscf
[nInfo]
method = ddr
basis = cc-pv5z
#Put grid in format of <start>, <end>, <step>
#for scattering put theta, phi as the 2d grid, with a given
#fixed value for rho
[gInfo]
rho = 4.5
theta = 3,90,3
phi = 0, 360, 3
#following options are only required for ddr nact calculation
dtheta = 0.03
```

4. The following sample 'molpro.config' file contains necessary MOLPRO keywords and other inputs for F+H₂ (2D ADT calculation in hyperspherical coordinate), where adiabatic PESs are generated in MRCI level of theory and analytic NACTs are calculated using CP-MCSCF method (see Table 14):

```
#Configuration file for F+H2 in hyperspherical coordinates
#Info for running molpro
[molInfo]
scrdir = /tmp/adtprogram
memory = 10, m
#for parallel molpro job
processor = 2
#Info of the molecular species
[sysInfo]
type = scat_hyper
#Info for energy calculation
[eInfo]
method = mrci
basis = aug-cc-pvtz
      = occ,10;closed,2;core,2
electron = 11
spin = 1
charge = 0
state = 3
#defining extra keyword required to run molpro for this system
#format should be same as in MOLPRO
multi_extra = maxiter,40
```

mrci_extra = maxiter,200,200

```
#defining scaling geometry for the adiabatic PESs
#minima of H2 at assymptotic F+H2 entrance channel
\#rho = 25.0, theta = 90.0, phi = 175.4
scale = 25.0,90.0,175.4
#Info for NACT calculation
#method should be either ddr or cpmcscf
[nInfo]
method = cpmcscf
basis = 6-311++G**
#Put grid in format of <start>, <end>, <step>
#for scattering put theta, phi as the 2d grid, with a given
#fixed value for rho
[gInfo]
rho = 5.0
theta = 0,90,5
phi = 0,360,4
```

5. The following sample 'molpro.config' file depicts necessary *ab initio* information for NO₃ radical, where adiabatic PESs are evaluated in CASSCF level of theory and analytic NACTs are calculated in CP-MCSCF method (see Table 13):

```
#Configuration file for the MOLPRO job of NO3
#Info for running MOLPRO
[molInfo]
scrdir = /tmp/adtprogram
memory = 10,m
#for parallel MOLPRO job
processor = 2
```

```
#Info of the molecular species
[sysInfo]
type = spec
#If symmetry is not specified, by default it will take 'nosym'
#C_s point group (yz is plane of symmetry)
#symmetry = x
#Info for energy calculation
[eInfo]
method = multi
basis = 6-31G**
     = occ,19;closed,11
electron = 31
spin = 1
charge = 0
state = 5
#defining extra keyword required to run molpro for this system
#format should be same as in MOLPRO
uhf_extra = accu,5
multi_extra = maxiter,40
\hbox{\#defining scaling geometry for the adiabatic PESs}
#rho = 0.0, phi = 0.0
#ground state minima of NO3 PES w.r.t normal modes
scale = 0.0, 0.0
#Info for NACT calculation
[nInfo]
method = cpmcscf
basis = 6-31G**
nact_extra = accu=1.d-10
#Info for varying modes
#put the indexes of the normal modes that will be varying
```

```
#The following two numbers represent the lowest frequency
#modes, but actually they are Q_4x and Q_4y normal modes
#(degenerate asymmetric bending modes)
[mInfo]
varying = 1,2

#Put grid in format of <start>, <end>, <step>
#for spectroscopic put the 2D grid as rho and phi
[gInfo]
rho = 0.1, 0.3, 0.1
phi = 0, 360, 3
#Qx = rho*cos(phi) and Qy = rho*sin(phi)
```

6. The following sample 'molpro.config' file represents necessary inputs for C₆H₆⁺ radical cation, where adiabatic PESs are computed in CASSCF level of theory and analytic NACTs are evaluated in CP-MCSCF method (see Table 13):

```
#Configuration file for the MOLPRO job of C6H6+
#Info for running molpro
[molInfo]
scrdir = /tmp/adtprogram
memory = 100,m

#for parallel MOLPRO job
processor = 1

#Info of the molecular species
[sysInfo]
type = spec

#Info for energy calculation
[eInfo]
method = multi
```

```
basis = 6-31G**
cas = occ,21;closed,6
electron = 41
spin = 1
charge = 1
state = 5
#defining extra keyword required to run molpro for this system
#format should be same as in MOLPRO
uhf_extra = wf,41,1,1,1
multi_extra = maxiter,40;
#defining scaling geometry for the adiabatic PESs
#ground state minima of C6H6+ w.r.t normal modes
scale = 0.0, 0.0
#Info for NACT calculation
[nInfo]
method = cpmcscf
basis = 6-31G**
nact_extra = accu=1.d-10
#Info for varying modes
#put the indexes of the normal modes that will be varying
#The following two numbers represent Q_16x and Q_16y normal
#modes (degenerate asymmetric stretching modes)
[mInfo]
varying = 23,24
#Put grid in format of <start>, <end>, <step>
#for spectroscopic put the 2D grid as rho and phi
[gInfo]
rho = 0.1, 0.3, .1
phi = 0, 360, 3
\#Qx = rho*cos(phi) and Qy = rho*sin(phi)
```

7. The following sample 'molpro.config' file depicts relevant inputs for 1,3,5-C₆H₃F₃⁺ radical cation, where adiabatic PESs are computed in CASSCF level of theory and analytic NACTs are evaluated in CP-MCSCF method (see Table 13):

```
#Configuration file for the MOLPRO job of 1,3,5-C6H3F3+
#Info for running molpro
[molInfo]
scrdir = /tmp/adtprogram
memory = 100, m
#for parallel molpro job
processor = 1
#Info of the molecular species
[sysInfo]
type = spec
#Info for energy calculation
[eInfo]
method = multi
basis = cc-pvdz
    = occ, 36; closed, 27
cas
electron = 65
spin = 1
charge = 1
state = 6
#defining extra keyword required to run molpro for this system
#format should be same as in MOLPRO
uhf_extra = wf,65,1,1,1;accu,5
multi_extra = maxiter,40
#defining scaling geometry for the adiabatic PESs
#ground state minima of 1,3,5-C6H3F3+ w.r.t normal modes
scale = 0.0, 0.0
```

```
#Info for NACT calculation
[nInfo]
method = cpmcscf
basis = 6-31G**
#Info for varying modes
#put the indexes of the normal modes that will be varying
#The following two numbers represent Q_{12x} and Q_{12y} normal
#modes (degenerate C-C-C scissoring modes)
[mInfo]
varying = 16,17
#Put grid in format of <start>, <end>, <step>
#for spectroscopic put the 2D grid as rho and phi
[gInfo]
rho = 0.02, 0.06, 0.02
phi = 0, 360, 3
\#Qx = rho*cos(phi) and Qy = rho*sin(phi)
```

Table 12: Description and possible inputs of MOLPRO configuration file ('molpro.config') for adiabatic PESs and NACTs

| Sections | Keywords | Description | | | | |
|----------|-------------|---|--|--|--|--|
| | scrdir | path of scratch directory for wavefunction file, namely: | | | | |
| molInfo | | /tmp/adtprogram | | | | |
| | memory | memory required for the job, e.g., 10,m | | | | |
| | processor | number of processors required for parallelization of the job, e.g., 2 | | | | |
| T. 6 | type | spec or scat_hyper or scat_jacobi | | | | |
| sysInfo | symmetry | symmetry of the molecule (default value: nosym) | | | | |
| | | optional, e.g., \mathbf{x} for \mathbf{H}_3^+ (C_s point group: yz is the plane of symmetry) | | | | |
| | method | potential energy surface: multi or mrci | | | | |
| | basis | basis for energy calculation, namely: $6-31G**$ for NO_3 | | | | |
| | cas | complete active space, e.g., occ,19;closed,11 for NO_3 | | | | |
| | electron | number of electrons, namely: 31 for NO_3 | | | | |
| | spin | spin multiplicity $(2s)$, namely: 1 for NO ₃ | | | | |
| T 6 | charge | charge of a molecular species, namely: 1 for ${\rm H_3^+}$ | | | | |
| eInfo | state | number of electronic states, e.g., 5 for NO ₃ , | | | | |
| | | 3,0 for H_3^+ (two values due to C_s symmetry) | | | | |
| | scale | spectroscopic: Q_i and $Q_j \Rightarrow \rho$ and ϕ , namely, 0.0,0.0 for NO ₃ | | | | |
| | | scattering (hyperspherical): ρ,θ and $\phi,$ namely, 25.0,90.0,175.4 for F+H ₂ | | | | |
| | | scattering (Jacobi): r, R and γ , namely, 1.0,24.0,0.0 for F+H ₂ | | | | |
| | restricted | true for rhf and false for uhf (default is false) | | | | |
| | uhf_extra | extra specification for uhf, namely: $wf,2,1,0,1$; for H_3^+ | | | | |
| | multi_extra | extra specification for multi jobs, | | | | |
| | | namely: maxiter,40 for NO_3 | | | | |
| | mrci_extra | extra specification for \mathtt{mrci} jobs, namely: $\mathtt{maxiter,200,200}$ for $F+H_2$ | | | | |
| | method | NACT calculation: cpmcscf for analytic gradient | | | | |
| nInfo | | and ddr for numerical finite difference method. | | | | |
| | basis | basis set for NACT, e.g., $6-31G**$ for NO ₃ | | | | |
| | nact_extra | extra specification for NACT jobs, namely: accu=1.d-10 for NO_3 | | | | |
| mInfo | varying | specification for pairwise normal modes (Q_i, Q_j) , e.g., 1,2 | | | | |
| | | (only required for spectroscopic calculation) | | | | |
| | rho | ρ grid for spectroscopic or scattering calculation in {start, end, step} format, | | | | |
| | | namely: 0.1, 0.3, 0.1 or 1.0, 20.0, 0.25, respectively | | | | |
| | theta | θ grid (hyperspherical) for scattering calculation, e.g., 3, 90, 3 | | | | |
| | phi | ϕ grid for spectroscopic/scattering calculation, e.g., 0, 360, 3 | | | | |
| | small_r | diatomic bond length, e.g, 2.0 | | | | |
| gInfo | capital_r | distance of third atom from diatomic center of mass, e.g, 1.73205 | | | | |
| | gamma | angle between small_r and capital_r, e.g., 90.0 | | | | |
| | q | radius of the circular path (see Figure 4) e.g., 0.5 | | | | |
| | drho* | ρ spacing in ddr method, namely: 0.01 | | | | |
| | dtheta* | θ spacing in ddr method, namely: 0.03 | | | | |
| | dphi* | ϕ spacing in ddr method, namely: 0.03 | | | | |
| | dq* | q spacing in ddr method, namely: 0.02 | | | | |

^{*}The values of drho, dtheta, dphi and dq require adjustment based on the convergence of the calculated result.

Table 13: Description of 'molpro.config' for NO_2 , NO_3 , $C_6H_6^+$ and 1,3,5- $C_6H_3F_3^+$ for the test runs

| G4: | 17 1 | NO | NO | O 11+ | 1 2 5 C H E+ | |
|----------|-------------|-----------------|-----------------------|-------------|-------------------------|--|
| Sections | Keywords | NO_2 | NO_3 | $C_6H_6^+$ | $1,3,5$ - $C_6H_3F_3^+$ | |
| | scrdir | | /tmp/a | dtprogram | | |
| molInfo | memory | 10,m | 1 | | 100,m | |
| | processor | 2 | | 1 | 1 | |
| avaInfo | type | | \$ | spec | | |
| sysInfo | symmetry | x | — | — | _ | |
| | method | multi | multi | multi | multi | |
| | basis | spdf,n,cc-pvtz; | 6-31G** | 6-31G** | cc-pvdz | |
| | | spdf,o,cc-pvtz | | | | |
| | cas | occ,11,3; | occ,19; | occ,21; | occ,36; | |
| . T., f. | | closed,3,0; | closed,3,0; closed,11 | | closed,27 | |
| eInfo | electron | 23 | 31 | 41 | 65 | |
| | spin | | | 1 | | |
| | charge | 0 | 0 | 1 | 1 | |
| | state | 2,0 | 5 | 5 | 6 | |
| | scale | | 0. | 0,0.0 | | |
| | uhf_extra | | accu,5 | wf,41,1,1,1 | wf,65,1,1,1;accu,5 | |
| | multi_extra | | max | iter,40 | | |
| | method | cpmcscf | | | | |
| nInfo | basis | spdf,n,cc-pvtz; | 6-31G** | 6-31G** | 6-31G** | |
| | | spdf,o,cc-pvtz | | | | |
| | nact_extra | | accu=1.d-10 | accu=1.d-10 | _ | |
| mInfo | varying | 1,3 | 1,2 | 23,24 | 16,17 | |
| T- C | rho | 0.2,12.0,0.2 | 0.1,0.3,0.1 | 0.1,0.3,0.1 | 0.02,0.06,0.02 | |
| gInfo | phi | 0, 360, 1 | | 0, 360, | 3 | |

Table 14: Description of 'molpro.config' for H_3^+ and $\mathrm{F} + \mathrm{H}_2$ for the test runs

| Sections Keywords H_3^+ (1D calculation) H_3^+ (2D calculation) scrdir /tmp/adtprogram molInfo memory 10,m | F+H ₂ | |
|--|------------------|--|
| | | |
| molInfo memory 10.m | | |
| | | |
| processor 2 2 | 2 | |
| type scat_jacobi scat_hyper | er | |
| sysInfo symmetry x | _ | |
| method mrci | | |
| basis cc-pv5z | aug-cc-pvtz | |
| cas occ,10,1;closed,0,0 oc | occ,10;closed,2; | |
| | core,2 | |
| electron 2 | 11 | |
| spin 0 | 1 | |
| eInfo charge 1 | 0 | |
| state 3,0 | 3 | |
| scale — 1.65,0.0,0.0 2 | 25.0,90.0,175.4 | |
| restricted true true | _ | |
| uhf_extra wf,2,1,0,1; | _ | |
| multi_extra maxiter,40 maxiter,40;ACCURACY, | maxiter,40 | |
| GRADIENT=1.d-4, | | |
| ENERGY=1.d-6, | | |
| STEP=1.d-2 | | |
| mrci_extra — m | maxiter,200,200 | |
| method cpmcscf ddr | cpmcscf | |
| nInfo basis 6-311++G** cc-pv5z | 6-311++G** | |
| rho — 4.5 | 5.0 | |
| theta — 3, 90, 3 | 0, 90, 5 | |
| phi 0, 360, 3 | 0, 360, 4 | |
| small_r 2.0 — | _ | |
| gInfo capital_r 1.73205 — | _ | |
| gamma 90.0 — | | |
| q 0.5 — | | |
| dtheta — 0.03 | _ | |
| dphi — 0.03 | _ | |

8.2. Atom Information File

• necessary file both for spectroscopic and scattering calculations

• First column: atomic symbols

• Second column: atomic masses in amu

Example: Sample file for NO₃ radical, 'atomfile.dat':

```
N 14.006700

O 15.999400

O 15.999400

O 15.999400
```

8.3. Equilibrium Geometry File

- necessary file for spectroscopic calculations
- contains the equilibrium geometry of the system
- Cartesian coordinates of the constituent atoms (in Angstroms) must be in a $n \times 3$ matrix form for a n atomic molecule
- order of atoms must be same as in atom information file

Example: Sample file for NO₃ radical, 'geomfile.dat':

| 0.00000000 | 0.00000000 | 0.00000000 |
|-------------|-------------|------------|
| 1.23680091 | 0.00000000 | 0.00000000 |
| -0.61840019 | -1.07110018 | 0.00000000 |
| -0.61840019 | 1.07110018 | 0.00000000 |

8.4. Frequency File

• necessary file for spectroscopic calculations

• contains frequencies of normal modes (in cm⁻¹) of a species arranged in ascending order (in a single row)

Example: Sample file for NO₃ radical, 'frequency.dat':

| 476.88 476.91 745.13 1047.52 1185.60 1185.8 |
|---|
|---|

8.5. Wilson Matrix File

- necessary file for spectroscopic systems
- provides the Wilson matrix of a molecular species
- the dimension of Wilson matrix will be $3n \times 3n 5$ for linear molecules (vibrational DOFs: 3n 5) and $3n \times 3n 6$ for non-linear ones (vibrational DOFs: 3n 6)
- columns are written as the increasing order of normal mode frequencies
- rows are sorted as per the order of atoms in atom information file

Example: Sample file for NO₃ radical, 'wilson.dat':

| 0.4681243 | 0.0000000 | 0.0000000 | 0.0000102 | 0.0000000 | 0.7449584 |
|------------|------------|------------|------------|------------|------------|
| 0.0000000 | -0.4682375 | 0.0000000 | 0.0000000 | 0.7448872 | 0.0000000 |
| 0.0000000 | 0.0000000 | 0.8798314 | 0.0000000 | 0.0000000 | 0.0000000 |
| 0.3427462 | 0.0000000 | 0.0000000 | 0.5774593 | 0.0000000 | -0.5394732 |
| 0.0000000 | 0.6348331 | 0.0000000 | 0.0000000 | 0.0749380 | 0.0000000 |
| 0.0000000 | 0.0000000 | -0.2744083 | 0.0000000 | 0.0000000 | 0.0000000 |
| -0.3903747 | -0.4233146 | 0.0000000 | -0.2887344 | -0.2660973 | -0.0787757 |
| 0.4234383 | -0.0983620 | 0.0000000 | -0.4999028 | -0.3859480 | -0.2660775 |
| 0.0000000 | 0.0000000 | -0.2744055 | 0.0000000 | 0.0000000 | 0.0000000 |
| -0.3903747 | 0.4233146 | 0.0000000 | -0.2887344 | 0.2660973 | -0.0787757 |
| -0.4234383 | -0.0983620 | 0.0000000 | 0.4999028 | -0.3859480 | 0.2660775 |
| 0.0000000 | 0.0000000 | -0.2744055 | 0.0000000 | 0.0000000 | 0.0000000 |
| | | | | | |

The first three (3) rows signify 'x, y, z' coordinates of one (1) nitrogen atom and the following nine (9) rows represent Cartesian coordinates of three (3) oxygen atoms.

References

- [1] Werner, H.-J.; et. al., MOLPRO, version 2010.1, a package of ab initio programs. 2010; http://www.molpro.net.
- [2] Frisch, M. J. et al. Gaussian 16 Revision C.01. 2016; Gaussian Inc. Wallingford CT.
- [3] Gordon, M. S.; Schmidt, M. W. Theory and applications of computational chemistry; Elsevier, 2005; pp 1167–1189.