EXP-T

a relativistic multireference coupled cluster program v 1.7.0

User manual $public\ version$

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1 Introduction

The EXP-T program package is designed for the high-precision modeling of molecular electronic structure using the relativistic Fock space coupled cluster method (FS-RCC). Features and program components are listed in Sec. 1.1, 1.2 and 1.3.

The authors will be grateful for any comments or suggestions:

```
exp-t-program@googlegroups.com
https://groups.google.com/d/forum/exp-t-program
```

1.1 Electronic structure models

The EXP-T package does not include subroutines for solving the (Dirac-) Hartree-Fock equations and subsequent four-index transformation, so molecular integrals have to be imported from third party electronic structure packages. Currently EXP-T is interfaced to the DIRAC program package [1], thus getting access to the wide variety of Hamiltonians and property operators implemented therein.

Models available in EXP-T:

- single-point energy calculations with any point groups and (nearly) all Hamiltonians, implemented in DIRAC (4c-DC, X2Cmmf, 2c-ECP, non-relativistic);
- ground state energy calculations (the 0h0p sector): CCSD, CCSD(T), CCSDT-n (n=1,2,3), CCSDT models;
- the FS-CCSD method for excited states is implemented for the 0h1p, 1h0p, 1h1p, 0h2p, 2h0p, 0h3p Fock space sectors;
- FS-CC models for excited states accounting for triples (CCSDT-n (n=1,2,3), CCSDT) are implemented for the 0h1p, 1h0p, 1h1p, 0h2p, 0h3p Fock space sectors.

The summary of all electronic structure models implemented in the EXP-T program system is given below:

FS sector	CCSD	CCSD+T(3)	CCSDT-1	CCSDT-2	CCSDT-3	CCSDT
$\overline{0h0p}$	+	+	+	+	+	+
0h1p	+	t	t	t	t	t
1h0p	+	t	t	t	t	t
1h1p	+	t	t	t	t	t
0h2p	+	t	t	t	t	t
2h0p	+	_	t	t	t	t
0h3p	+	t	t	t	t	t

"+" - implemented, "-" - not implemented, "t" - currently being tested

- "dynamic" energy denominators shifts as a solution of the intruder-state problem [2];
- Padé extrapolation to the zero-shift limit [3];
- intermediate Hamiltonian technique for incomplete main model spaces (IH-IMMS) [4];

- finite-field transition moments calculations [5];
- quasidiabatization of SO-coupled states and SO extraction [2].
- «quasi-natural» spinors (NS) and natural transition spinors (NTS);
- one-electron properties via interface to the OneProp program by L. V. Skripnikov [6].

1.2 Features for high-performance calculations

- OpenMP parallelization (for shared-memory systems);
- parallel calculations on NVIDIA GPUs using the CUDA platform [7].

1.3 Program components

expt.x Input processing + all CC calculations.

heffman.x Manipulations with effective Hamiltonian matrices: Padé extrapolations, finite-field transition moments calculations, quasidiabatization of SO-coupled states.

expt_transform_natorb.x

Transformation of "quasi-natural" spinors to the AO basis.

expt_diatomic.x

Rotational-vibrational levels and properties of diatomic molecules.

expt2pam.x Tool for convenient basis set and ECP input.

1.4 Citation

We kindly ask you to acknowledge any use of the EXP-T program system that results in published material using the following citation:

A. V. Oleynichenko, A. Zaitsevskii, E. Eliav. Towards high performance relativistic electronic structure modelling: the EXP-T program package. Commun. Comput. Inf. Sci. 1331, 375 (2020). doi: 10.1007/978-3-030-64616-5 33

In addition, credit must be given to the corresponding papers which describe the implementation and the underlying methodological developments:

- Triples in the low Fock space sectors: [8]
- The 0h3p Fock space sector: [9]
- Finite-field transition matrix elements (general): [5]
- Finite-field transition matrix elements (the $0h0p \rightarrow 1h1p$ excitations): [10]
- Spin-orbit coupling calculations: [2]

- Dynamic denominator shifts (real): [2]
- Dynamic denominator shifts (imaginary, or their real simulation): [11]
- Padé extrapolation of effective Hamiltonian series: [3]
- Intermediate Hamiltonian for incomplete main model spaces (IH-IMMS): [4]

1.5 Credits

EXP-T is based on ideas and design solutions of the suite of FS-CC programs written by Ephraim Eliav and Uzi Kaldor.

2 Installation from source code

2.1 Compiling EXP-T

EXP-T is currently oriented at Unix-like operating systems.

Tools required for compiling the EXP-T package from source code:

• C and Fortran compilers. Currently supported compiler systems are

```
gcc/gfortran GNU compilers icc/ifort Intel compilers
```

• cmake version 3.0.2 or higher and make utilities:

```
https://cmake.org/
https://www.gnu.org/software/make/
```

Optional dependencies:

- Implementation of the BLAS/LAPACK linear algebra libraries (OpenBLAS or Intel MKL are recommended);
- NVIDIA drivers and CUDA Toolkit are required to perform parallel calculations on GPU: https://www.nvidia.com/Download/index.aspx?lang=en-us https://developer.nvidia.com/cuda-toolkit

Download the EXP-T source code as a *.tar.xz file and unpack it:

```
tar xvf expt-X.Y.Z.tar.xz
```

Go to the EXP-T home directory, create the build directory and make it the current working dir:

```
mkdir build
cd build
Compilation (Intel compilers):
CC=icc FC=ifort cmake ..
make
Compilation (GNU compilers):
CC=gcc FC=gfortran cmake ..
make
```

We recommend specifying compilers explicitly (since the default C compiler may be something other than gcc).

The CMake utility will try to locate BLAS/LAPACK libraries on your computer; in case of multiple implementations found you will be asked to choose one of them. If there are no pre-installed libraries on your machine, the internal OpenBLAS will be compiled and linked to EXP-T.

When using Intel MKL together with GNU compilers, you will probably need to explicitly specify the path to the MKL home directory by setting the environment variable MKLROOT. For example:

export MKLROOT=/opt/intel/mkl

In case of successful compilation, the executable file expt.x will appear in the build directory.

We strongly recommend that compilation is followed by testing (see Sect 2.4).

Do not forget to add the directory containing the binaries to the PATH environment variable!

2.1.1 Compiling on Mac OS X

The default C compiler for Mac OS X is clang; this compiler was not tested, so we strongly recommend to use gcc instead:

```
CC=gcc FC=gfortran cmake .. make
```

We strongly recommend not to use GNU compilers installed with Homebrew. Instead, download the GNU compilers suitable for your version of Mac OS X from:

```
http://hpc.sourceforge.net/
```

In order to use a BLAS/LAPACK implementation other than the one attached to the EXP-T package, you can use the following manual:

https://pheiter.wordpress.com/2012/09/04/howto-installing-lapack-and-blas-on-mac-os/However, the Netlib implementation described there can perform several times slower than the OpenBLAS implementation included in EXP-T.

2.2 Compiling with CUDA

Compilation of EXP-T does not require the user to have a CUDA-compatible GPU device. If the CUDA toolkit is missing, the CUDA code will simply be excluded from the compilation process.

EXP-T is adapted for parallel calculations on NVIDIA GPUs (the CUDA technology [7]). Notes:

- 1. At the moment, the OpenMP+CUDA hybrid model as well as multi-GPU support is not implemented. Only one GPU can be used for calculations.
- 2. non-professional GPUs for gaming are not suitable for CUDA calculations (except for Kepler cards).
- 3. CUDA code in EXP-T was tested only for CUDA 9.1 and 10.0.
- 4. Only Intel and GNU compilers were tested for compatibility with CUDA.

To compile EXP-T with CUDA:

- 1. Make sure your graphics card supports CUDA. CUDA-compatible GPUs are listed here: https://developer.nvidia.com/cuda-gpus
- 2. Install proprietary NVIDIA drivers (not the nouveau driver!). https://www.nvidia.com/Download/index.aspx?lang=en-us. For Linux you can find drivers in your distribution's repository.
- 3. Install CUDA toolkit https://developer.nvidia.com/cuda-downloads or from distribution's repository.

- 4. Make sure that all drivers and CUDA toolkit are installed and work correctly. Compile the test suite in the samples directory of the CUDA home directory and run some of them. The most important are the tests employing the CUBLAS linear algebra library.
- 5. Compile EXP-T as described in Sec. 2.1. All CUDA tools will be located automatically by CMake.
- 6. Make sure that the results produced by EXP-T do not depend on the use of CUDA.

2.3 Troubleshooting

This section describes the most common problems which can occur during the compilation process. In case of any questions or problems, please don't hesitate to contact us. Don't forget to attach the output file (in case of problems with calculations) or the file with the output of the make VERBOSE=1 command (in case of problems with compilation).

Google Groups:

```
https://groups.google.com/d/forum/exp-t-program
```

E-mail:

```
exp-t-program@googlegroups.com
```

2.3.1 Outdated version of CMake

We highly recommend using CMake version 3.0.2 or higher, otherwise, problems may occur when building the internal OpenBLAS. The following command will show what version of CMake is installed:

```
cmake --version
```

If CMake version is lower than 3.0.2, download the binary distribution from their official website https://cmake.org/download/. After installation do not forget to add the directory containing the CMake binary files to the PATH environment variable.

Installation of CMake binary files does not require root privileges.

2.4 Testing

Test suite is located in the test directory. The testing system is written in the Python programming language (must be pre-installed on your machine). Either Python 2 or Python 3 can be used.

In order to run the testing suite, run the following command:

make test

3 Running EXP-T

See Sec. 6 for a description of an additional utility programs included in the EXP-T package.

A typical calculation consists of two sequential steps:

- 1. SCF and integral transformation within DIRAC;
- 2. CC calculation within EXP-T.

3.1 Precomputing spinors and molecular integrals: DIRAC step

The interface to DIRAC was tested only for the DIRAC17 and DIRAC18 releases. DIRAC stores transformed molecular integrals in the following binary files:

MRCONEE information about SCF calculation, symmetry, occupation numbers, one-electron integrals;

MDCINT symmetry-unique nonzero two-electron integrals;

MDPROP one-electron property integrals.

Run DIRAC and save the files containing the transformed integrals to the working directory using the --get option:

```
pam --noarch --mol=<mol-file> --inp=<inp-file> --get="MRCONEE MDCINT MDPROP"
```

If DIRAC starts in parallel mode (MPI), it is necessary to choose the transformation algorithm "scheme 4". By default, DIRAC uses the "scheme 6" algorithm which produces multiple MDCINT files [12] and currently EXP-T requires that all integrals are stored in a single file. In order to accomplish this, add the following lines to the DIRAC input file:

```
**MOLTRA
.SCHEME
```

To calculate approximate transition moments and intensities without resorting to the finite-field scheme transformed dipole moment integrals are required. We recommend explicitly ask DIRAC to calculate and transform these integrals by adding the <code>.PRPTRA</code> and <code>.DIPOLE</code> keywords to DIRAC input files:

```
**MOLTRA
. PRPTRA
. . . . . . .
**PROPERTIES
. DIPOLE
```

If the molecular-mean-field X2C Hamiltonian (X2Cmmf) was used at the DIRAC stage, one must specify the x2cmmf flag in the EXP-T input file! (see also page 23).

3.2 Coupled cluster: EXP-T step

Run an EXP-T job using the following command:

```
expt.x [options] <input-file>
```

The EXP-T output (including error messages) is flushed to the UNIX standard output (stdout). Use the output redirection operator ">" in case of long time calculations:

```
nohup expt.x <input-file> > <output-file> &
```

Examples of DIRAC and EXP-T input files are given in Sec. 5.

3.3 Command-line arguments

Use the expt.x --help command to print a list of available command-line arguments:

Usage: expt.x [OPTION...] <input-file>
expt -- A Fock-Space Multireference Relativistic Coupled-Cluster Program

-n,no-clean	Do not clean scratch directory on exit			
	(use this option to keep cluster amplitudes etc)			
-s,scratch=PATH	Path to scratch directory (default: ./scratch)			
-?,help	Give this help list and exit			
usage	Give a short usage message and exit			
-V,version	Print program version and exit			

Mandatory or optional arguments to long options are also mandatory or optional for any corresponding short options.

Please report bugs to <exp-t-program@googlegroups.com>.

4 Input files syntax and keywords

See Sec. 6 for the description of additional utitily programs included in the EXP-T package.

The input file format may change in future versions.

The EXP-T input file contains a list of instructions. Each instruction is placed on a separate line and consists of a keyword and a list of arguments. Single-line comments start with '#' and continue until the end of the line. Input is case insensitive. Examples of EXP-T input files are given in Sec. 5. Molecular geometry, basis set etc have to be described only once in the DIRAC input files at the molecular integrals calculation stage (not in EXP-T input files).

The detailed description of the keywords used in the EXP-T input file is provided below. For convenience, keywords are grouped into several sections. The order of keywords in the input file can be arbitrary.

General

title optional comment.

Syntax: title <quoted-string>

Default: no title

Example: title "my first ccsd calculation"

print print level.

Syntax: print (low || medium || high || debug || <quoted-string>) where <quoted-string> can be:

"model vectors" effective configurations of spinors for each electronic state;

"model vectors" expansions of model vectors in terms of model determinants;

"model space" information about model space determinants.

One is allowed to combine several print instructions.

Default: print medium

Example:

print low

print "eff config"

degen_thresh energy levels are considered as degenerate if the energy gap between them is lower than degen_thresh.

Syntax: degen_thresh <real thresh>

Units: atomic
Default: 1e-8

spinor_labels manual assignment of symbolic names (labels) of specific spinors (orbitals). Symbolic name is printed together with spinor symmetry and energy. This option can be useful when studying composition of electronic states.

Syntax:

Model

sector

specifies the target Fock space sector. To the moment the 0h0p, 0h1p, 1h0p, 1h1p, 0h2p, 2h0p, 0h3p FS sectors are implemented (h - holes, p - particles).

Syntax: sector <integer H>h<integer P>p

Default: sector OhOp (vacuum)

model

CC model (approximation to the cluster operator).

Default: model ccsd

Notes:

- Keywords ccsd(t) and ccsd+t(3) are equivalent.
- The codes implementing the CCSD+T(3), CCSDT-n and CCSDT models in the non-trivial sectors and all CC models in the 0h3p sector are experimental (currently being tested), to be used with care.
- The current implementation of the CCSD(T) model for the 0h0p sector is not very computationally efficient.
- The CCS and CCD models are implemented simply by setting all T_2 (CCS) or T_1 (CCD) amplitudes to zero, this does not make computation faster, and exist primarily for the purposes of testing and debugging.

occ spinor occupation numbers. This keyword can be used to setup calculations with a "high-spin" vacuum state [13]. This keyword is alternative to the occ_irreps keyword (see below).

Syntax: occ <list of the 0 and 1 digits separated by spaces>

Default: occupation numbers are read from MO integrals files

Example: 5 electrons and 10 spinors:

occ 1 1 0 0 0 1 1 1 0 0

CC calculations with high-spin vacuum states were tested for the 0h0p FS sector only.

occ_irreps overall number of electrons in each irrep. Overall irrep occupations are ignored if occupation numbers of individual spinors are given (by the occ keyword, see above).

Syntax: occ_irreps <list of integers>

Default: occupation numbers are read from MO integrals files

Active space

nacth/nactp active (i.e. valence) space specification (by the overall number of active spinors, regardless of their symmetry). This method of configuring the active space is recommended.

- nacth number of highest occupied spinors (active holes)
- nactp number of lowest virtual spinors (active particles)

It is convenient to setup the active space by the nacth/nactp keywords when exploring electronic states of molecules in a range of geometries (potential energy surfaces, transition moment functions etc). Using the nacth/nactp keywords one can specify either overall number of active spinors (holes or particles) or number of active spinors in each irreducible representation.

```
Syntax: nacth <integer dim>
    nactp <integer dim>
    or
    nacth <pairs irrep:number-of-spinors>
    nactp <pairs irrep:number-of-spinors>
```

Default: nacth 0, nactp 0 (no active spinors).

Example: consider the low-lying electronic states of the La atom. Those are dominated by the $6s^n5d^m$ configurations, generating even electronic states. So it seems reasonable to include the $6s_{1/2}$, $6d_{3/2}$ and $6d_{5/2}$ atomic spinors of the La³⁺ cation into the active space and achieve target electronic states in the 0h3p Fock space sector. In order to make the input file easy to read, one can use two sequential

nactp directives, one for the "negative" projections of m_j , and one for the "positive" ones:

```
nactp [1/2g+]:3 [3/2g+]:2 [5/2g+]:1 nactp [1/2g-]:3 [3/2g-]:2 [5/2g-]:1
```

One can extend the scope of achievable electronic states by inclusion of the 6p and 4f spinors:

```
nactp [1/2u+]:4 [3/2u+]:3 [5/2u+]:2 [7/2u+]:1 nactp [1/2u-]:4 [3/2u-]:3 [5/2u-]:2 [7/2u-]:1
```

active

active (i.e. valence) space specification by energy range. Active spinors must have one-electron energies in the range $\varepsilon_i \in [\varepsilon_{min}; \varepsilon_{max}]$ (regardless of their symmetry). Setting the active space using the active keyword is convenient only for atomic calculations.

Syntax: active energy <real eps_min> <real eps_max>

Default: no active spinors

Example: active energy -1.0 5.0

Intermediate Hamiltonian for incomplete main model spaces (IH-IMMS)

Intermediate Hamiltonian for incomplete main model spaces (IH-IMMS) can be used to achieve convergence of iterative solution of CC equations [4].

The basic idea of the method is to split model space into the "main" and "intermediate" subspaces. Slater determinants $|\Phi_M\rangle$ which span the main model space must be chosen to describe the bulk of wavefunctions of target electronic states. FS-CC equations for amplitudes corresponding to excitations from main determinants remain unchanged. In contrast, equations for amplitudes from intermediate ("buffer") determinants $|\Phi_I\rangle$ are solved with shifted energy denominators to suppress the intruder state problem. For each cluster amplitude corresponding to the excitation $I \to K$ from $|\Phi_I\rangle$ a shift parameter S_{IK} is calculated as:

$$S_{IK} = \overline{E}_0 - E_I,$$

where E_I stands for the energy of the $|\Phi_I\rangle$ model-space determinant and \overline{E}_0 stands for the some "frontier" energy. This value is used to calculate a shift for energy denominator corresponding to a cluster amplitude using one of the formulas (1)–(4). All model space determinants $|\Phi_M\rangle$ with energy $E_M \leq E_0$ will be considered as main ones. Determinants $|\Phi_I\rangle$ with energy $E_I > E_0$ are intermediate.

By default, IH-IMMS is disabled. IH settings are specified using the complex directive ih_imms:

```
ih_imms
    <option-1>
     <option-2>
     <option-3>
          . . .
end
```

To set up the IH-IMMS calculation:

- 1. split the set of active spinors into several subspaces. In the current version of EXP-T only spinor energies can be used to select these subspaces;
- 2. define model Slater determinants which will be considered as the "main" ones. Sets of "main" determinants are defined via the numbers of particles (holes) in each spinor subspace;
- 3. (optional) define the "frontier" energy \overline{E}_0 . By default, frontier energy is calculated automatically.

The use of IH-IMMS for a simple case of carbon atom is considered in details in Sect. 5.4. The following keywords can be used inside the ih_imms block:

sectors list of Fock space sectors in which the IH-IMMS technique will be applied.

Syntax: sectors <list-of-sector-labels>

Default: no target FS sectors. Example: sectors 0h1p 0h2p

subspace defines subspace of active spinors with energies $\varepsilon \in [\varepsilon_{\min}, \varepsilon_{\max}]$. Subspaces are used to define composition of main model space determinants via the main_occ keyword. One can use several subspace options in the input file to define more than one subspace of active spinors.

Syntax: subspace energy <real emin> <real emax>

Default: no subspaces are defined.

Example: subspace energy -1.0 -0.5

main_occ defines subspace of main model space determinants. The argument of this option contains number of particles (holes) in the 1st, 2nd, ... subspaces of active spinors. One can use several main_occ options in the input file to define more than one subspace of main model space determinants.

Syntax: main_occ <list-of-integers>

Default: no main model spaces are defined.

Example: imagine that we have to define model space determinants with 2 electrons on active spinors belonging to the 1st spinor subspace, 1 electron on spinors from the 2nd subspace and 0 electrons on the other active spinors (such a model determinant belongs to the 0h3p sector). The proper option is as follows:

main_occ 2 1 0

shift_type which formula for denominator shift should be used for cluster amplitudes associated with "intermediate" model space determinants. See also description for the shifttype keyword 21.

Syntax: shift_type (none || real || realimag || imag || taylor)

Default: shift_type realimag

npower attenuation parameter used in the shifting formula (see Eq. (1)–(4)).

Syntax: npower <int n>

Default: npower 5

scale scaling factor for the shift parameter S:

$$S \rightarrow S' = \alpha S$$

Syntax: scale <real factor>
Default: scale 1.0 (no scaling)

frontier_energy "frontier" energy \overline{E}_0 used to calculate shift for the given (intermediate) determinant.

Syntax: frontier_energy <real energy>

Default: frontier energy is evaluated automatically as the maximum energy of main model space determinants:

$$\overline{E}_0 = \max_{\Phi_M \in \text{main}} E_M, \qquad E_M = \langle \Phi_M | H_0 | \Phi_M \rangle.$$

Properties

hermit perform symmetric orthogonalization of model vectors. Orthogonalized model vectors are used only to calculate properties and approximate density matrices; matrices of effective Hamiltonians flushed to HEFF files (see Sec. 6.1) remain non-Hermitian.

Syntax: hermit

Default: "hermitization" is disabled.

enable calculations of the model-space approximations to transition dipole moments (wavefunctions are represented by model vectors) [14, 15, 10]). To avoid huge outputs, use option nroots to restrict the set of target roots.

Note that left model vectors are used in bra:

$$d_{if,\eta} = \langle \tilde{\psi}_i^{\perp \perp} | d_{\eta} | \tilde{\psi}_f \rangle \qquad \eta = x, y, z$$

and if "hermitization" of the effective Hamiltonian matrix is disabled, the TDM matrix will be (normally slightly) non-Hermitian $(|d_{if}| \neq |d_{fi}|)$.

This approach typically overestimates transition moments by 30-50%, hence we recommend using it only for semiquantitative estimations, e.g. detection of the most intensive transitions and discerning between 0^+ and 0^- states (such transitions are strictly forbidden).

Syntax: mstdm

Default: disabled.

mdprop

enable calculations of the model-space approximations to arbitrary properties presented in the DIRAC's MDPROP unformatted file. To avoid huge outputs, use option nroots to restrict the set of target roots. To be used for semiquantitative estimations only.

Syntax: mdprop "roperty-name>" [transpose]

Default: disabled.

Example: calculation of x,y,z-components of transition moments.

mdprop "XDIPLEN"
mdprop "YDIPLEN"
mdprop "ZDIPLEN"

txtprop

enable calculations of the model-space approximations to arbitrary properties. Property matrix is loaded from two formatted files. To avoid huge outputs, use option **nroots** to restrict the set of target roots. To be used for semiquantitative estimations only.

Syntax: txtprop <string file_re> <string file_im> [transpose]

Default: disabled.

Example:

mdprop PropIntsRe.txt PropIntsIm.txt

The code for properties is currently experimental; please, we kindly ask you to report all failures and obviously incorrect results.

analyt_prop analytic calculation of coupled cluster expectation values of arbitrary property operators. Matrix elements of property operators must be stored in the DIRAC's MDPROP unformatted file. Λ -equations, analytic density matrix and properties are currently available only for the 0h0p sector (vacuum state).

Syntax: analyt_prop property-name>

Default: disabled.

Example: calculation of x,y,z-components of the dipole moment.

analyt_prop XDIPLEN
analyt_prop YDIPLEN
analyt_prop ZDIPLEN

density calculation of analytic density matrix in the 0h0p sector. Prints occupation numbers

of CC natural orbitals.

Syntax: density OhOp

Default: disabled.

natorb calculation of "quasi-natural" spinors (NS) and transition spinors (NTS).

Note: density matrices $d_{pq}^{IJ} = \langle \tilde{\psi}_I^{\perp \perp} | a_p^{\dagger} a_q | \tilde{\psi}_J \rangle$ are constructed with model vectors instead of true wavefunctions.

Syntax: natorb [<rep>:<state> | <rep1:state1>-<rep2:state2>] ...

Electronic states are given by the <rep>:<state> pair, where:

<re> name of representation in square braces. See Sec. C for the list of representation names used in EXP-T.

 \langle state \rangle sequential number of the state in the irrep (1, 2, ...).

Pair of electronic states separated by hyphen shows that NTS between these states are to be calculated. You can ask EXP-T to calculate NS and NTS for arbitrary number of states (or pairs of states).

Default: calculation of NS and NTS is disabled.

Examples:

• NS for the ground and the first excited states of the RaCl molecule:

```
natorb [1/2+]:1 [1/2+]:2
```

• NTS for the $(1)0^+ - (2)0^+$ transition in the RbCs molecule:

```
natorb [0]:1-[0]:3
```

(note that in the latter example the [0]:2 state is excluded, since it is the $(1)0^-$ state).

Iterative solution and convergence

arith allows one to force EXP-T to use the complex arithmetic everywhere.

Syntax: arith complex

nroots

specifies the number of roots (electronic states) of interest in each irrep (in the target Fock space sector). Model vectors analysis will be performed only for the nroots lowest states.

Syntax: nroots <pairs irrep:number-of-roots>

Default: all roots are of interest.

Example: consider the electronic states of the RbCs molecule, corresponding to the first three dissociation limits (see, for example, [2, 16] and references therein). In the Hund's case a these electronic states are $X^1\Sigma^+$, $a^3\Sigma^+$, $b^3\Pi$, $A^1\Sigma^+$, $c^3\Sigma^+$, $B^1\Pi$. In the Hund's case c these states are classified according the projection of the total electronic angular momentum and can be rearranged as follows: $(1-3)0^+$, $(1-3)0^-$, (1-4)1, (1)2 states. The names of irreps used in EXP-T are given in Appendix C (the point group of RbCs is $\overline{C}_{\infty v}$). All output and analysis of model vectors will be restricted to these 11 states using the following command:

nroots [0]:6 [1+]:4 [1-]:4 [2+]:1 [2-]:1 (note that as in DIRAC 0^+ and 0^- states belong to the same irrep).

roots_cutoff specifies the upper energy bound for the electronic states to be analyzed and printed. Model vectors analysis and properties calculations will be performed only for the states with energy lower than roots_cutoff. Allowed units of energy are eV, cm⁻¹ and Hartrees.

Syntax: roots_cutoff <real max_energy> (ev || cm || au)

Default: all roots are of interest.

Example: roots_cutoff 30000 cm

maxiter maximum number of iterations allowed to solve the CC equations.

Syntax: maxiter <integer max>

Default: maxiter 50.

conv specifies the requested convergence threshold (in amplitudes):

$$\left| |t_K^{(n+1)}| - |t_K^{(n)}| \right| < conv \quad \forall K \quad \to \text{converged}$$

Syntax: conv <real thresh>

Default: 1e-9.

div_thresh specifies the divergence threshold (in amplitudes). Solution of amplitude equation is diverged if for some t_K holds:

$$\left| |t_K^{(n+1)}| - |t_K^{(n)}| \right| > div_thresh \rightarrow diverged$$

May be useful if few initial steps of the iterative solution are divergent.

Syntax: div_thresh <real thresh>

Default: 1e+3.

diis enables the DIIS extrapolation technique [17, 18].

Syntax: diis [(off || triples || <integer dim>)]

Disable DIIS: diis off

Enable DIIS for triples: diis triples

Default: diis 10 (DIIS is enabled for singles and doubles, disabled for triples)

It is highly recommended to enable DIIS for triples if the CCSDT-n or CCSDT models are employed.

damping enable damping in the given FS sector. Amplitudes obtained at the n-th step will be mixed with amplitudes from the (n-1)-th step using the damping factor α :

$$T^{(n)'} = \alpha T^{(n-1)} + (1 - \alpha)T^{(n)}$$

Damping can aid the convergence of equations in some non-pathological cases (when the lack of convergence is caused by amplitude oscillations rather than intruder states). Note that in other cases damping can significantly slow down convergence.

Syntax: damping <H>h<P>p <integer last_step> <real factor>

<h><h>>p FS sector in which these damping parameters will be applied

last_step at which iteration damping will be disabled

factor damping factor α . Can be negative.

Default: disabled in all sectors

Example: damping 0h2p 100 0.5

shifttype indicates which formula for denominator shifts should be used [2, 3].

shifttype is the same for all FS sectors. If shifttype is not equal to none, you must specify the shift parameters for at least one FS sector (see keyword shift; except for 0h0p).

 $\operatorname{Syntax}:$ shifttype (none || real || realimag || imag || taylor)

Possible values:

none no shift

real real shift

$$D_K' = D_K + S \left(\frac{S}{D_K + S}\right)^n \tag{1}$$

realimag real simulation of imaginary shift [11]

$$D_K' = D_K + \frac{S^2}{D_K} \left(\frac{S^2}{D_K^2 + S^2} \right)^n \tag{2}$$

imag imaginary shift [11]

$$D_K' = D_K + iS \left(\frac{|S|}{|D_K + iS|}\right)^n \tag{3}$$

taylor extrapolated intermediate Hamiltonian-like shift [19]

$$D_K' = (D_K + S) \frac{\left(1 - \frac{S}{D_K + S}\right)}{1 - \left(\frac{S}{D_K + S}\right)^{n+1}} \tag{4}$$

Default: shifttype none

Shift is never applied in the 0h0p sector!

shift sets the denominator shift parameters in the $n_h h n_p p$ FS sector.

shift <H>h<P>p <int n> <real shift_S1> <real shift_S2> [<real shift_S3>]

n attenuation parameter (see eqs. (1) and (4));

shift_S1 shift value for the $S_1^{n_h h n_p p}$ cluster operator (singles). Must be omitted in the 2h0p and 0h2p FS sectors. It is recommended to set shift_S1=0.0 in the 1h1p sector, unless you really understand the consequences of using nonzero values. Note that shift_S1 value in this sector does not affect the resulting energies;

```
shift_S2 shift value for S_2^{n_h h \ n_p p} (doubles);
```

shift_S3 shift value for $S_3^{n_h h n_p p}$ (triples). Must be used only for CC models including non-perturbative triples.

Default: n = 3, all shifts = 0.0

Examples:

shift 0h1p 3 -0.3 -0.6 -0.9

shift 0h1p 3 -0.6

shift 1h1p 4 0.0 -0.5

reuse

indicates that sorted integrals and cluster amplitudes from the previous calculation should be reused. Cluster amplitudes will be used as an initial guess. If EXP-T fails to locate the files containing amplitudes, they will be recalculated.

Syntax: reuse <list-of-arguments>

Possible values of arguments:

```
integrals load one- and two-electron integrals (from a scratch directory);
```

 ${\tt 1-integrals} \quad {\rm one\text{-}electron\ integrals\ only;}$

2-integrals two-electron integrals only;

amplitudes load amplitudes for all FS sectors occuring in the calculation;

OhOp, Oh1p, 1h0p, 1h1p, Oh2p, 2h0p, Oh3p

load amplitudes for the given sector only

Default: no reuse

Note: files with converged amplitudes are flushed to the scratch directory after each calculation (see also Appendix B). To prevent the deletion of files with amplitudes or integrals it is necessary to use the --no-clean command-line option.

Example: reuse integrals 0h0p 0h1p

flush

indicates that cluster amplitudes must be flushed to disk every N iterations. Can be useful for very time-consuming calculations with poor convergence. Allows one to change the convergence acceleration strategy.

Syntax: flush <integer N> iter

Default: no flush.

Example: flush 5 iter

skip

indicates which sectors will be skipped in the FS-CC calculation. Can be useful in very costly calculations (like CCSDT) in order to avoid recalculation of the first step in already converged sectors. To be used with the reuse option.

Syntax: skip <list-of-sector-symbols>

Default: no skip.

Example: skip 0h0p 0h1p

Interfaces

integrals sets paths to files containing transformed molecular integrals.

Arguments:

- one-electron integrals file;
- two-electron integrals file;
- properties transformed integrals file.

Default: integrals MRCONEE MDCINT MDPROP

Example: integrals ../MRCONEE-Cinfv ../MDCINT-Cinfv /home/user/MDPROP

x2cmmf since DIRAC's unformatted files do not contain any information about the relativistic Hamiltonian used, the x2cmmf flag must be used if the molecular-mean-field X2C Hamiltonian (X2Cmmf) was employed at the DIRAC stage.

Syntax: x2cmmf

oneprop imports one-electron property integrals from the MDPROP unformatted file. The one-electron operator \hat{O}_1 is added to the Fock operator at the integral sorting stage:

$$\hat{f}' = \hat{f} + \lambda \hat{O}_1$$

where λ is some perturbation parameter. EXP-T does not assume Kramers symmetry, so both T-odd and T-even operators are allowed.

L_re, L_im real and imaginary parts of λ ;

property-name string identifier used to encode property matrix in the MDPROP file (can be also found in the DIRAC output file).

Default: disabled.

Example: add interaction with an external electric field:

oneprop 1e-5 0.0 "ZDIPLEN"

Memory management

memory specifies the amount of memory that EXP-T can use for the job (just for *dynamically* allocated memory).

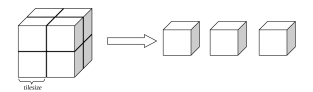
Syntax: memory <real size> (mb || gb)

Min value: 10 Mb

Default: memory 1024 mb (1 Gb)

tilesize the whole set of one-electron functions (spinors or spin-orbitals) is divided into blocks: (1) by irrep; (2) then into sub-blocks with size not exceeding tilesize (actually the same as "tiles" in NWChem-TCE):

The N-dimensional arrays of integrals and amplitudes (tensors) also turn out to be divided into blocks containing $\sim (tilesize)^N$ elements:



tilesize must be large enough to place at least one N-dimensional array of size $(tilesize)^N$ in RAM. We recommend using as large tilesize as possible, otherwise memory management overheads can become enormous. However, it may be useful to decrease the tilesize parameter in cases of insufficient RAM or threaded execution (sufficient granularity of arrays leads to better dynamic load balancing among OpenMP threads).

Syntax: tilesize <integer size>

Default: tilesize 100

disk_usage indicates which data should be stored on the disk. Note that very intensive disk usage will slow down calculations; however, disk usage is unavoidable in case of large tasks.

Syntax: disk_usage <integer mode>

Possible values of <mode>:

- 0 all data is stored in RAM;
- 1 tensors of rank ≥ 6 are stored on disk (T_3 amplitudes etc);
- 2 tensors of rank ≥ 6 and $\langle pp||pp\rangle$ (4 particles) two-electron integrals are stored on disk;
- 3 tensors of rank ≥ 6 , $\langle pp||pp\rangle$ and $\langle *p||pp\rangle$ (3 particles) two-electron integrals are stored on disk;
- 4 the same as 3, + compression of all data. Is suitable for extremely large tasks only.

Default: $2 - \langle pp || pp \rangle$ and tensors of rank ≥ 6 are stored on disk.

compress

enables compression of all data written to disk. The LZ4 algorithm [20] is employed (the fastest decompression to the date). Compression can slow down calculations slightly (depends on disk), but the disk space used can be reduced dramatically (up to 3 times).

Syntax: compress

Default: disabled (except for disk_usage=4)

Parallel execution

nthreads

number of OpenMP threads. Note that scaling with respect to number of threads is much better for large tasks (400+ spinors).

Syntax: nthreads <integer n_omp_threads>

Default: nthreads 1 (sequential execution).

At the moment parallelization is not used at the integral sorting stage, hence OpenMP can accelerate only the amplitude equations solution stage.

openmp

the same as nthreads.

cuda

enables parallel calculations on NVIDIA GPU (only a single GPU is supported at the moment). Requires nthreads 1 (the OpenMP+CUDA hybrid model is not implemented yet).

Syntax: cuda

Default: CUDA disabled.

Running typical calculations 5

The input and output files discussed in this section can be found in the examples directory of the EXP-T distro.

Ground state energy with CCSD: CO molecule 5.1

```
For input/output files see examples/CO_ccsd
```

```
Consider the calculation of the CCSD correlation energy of the CO molecule in the cc-pVDZ
[21] basis set.
   DIRAC input files:
   (CO.mol)
DIRAC
CO molecule, R = 2.132 bohrs
symmetry C1
    2
         6.
               1
          0.0
                    0.0
                              0.0
LARGE BASIS cc-pVDZ
         8.
               1
                              2.132
          0.0
                    0.0
LARGE BASIS cc-pVDZ
FINISH
   (TRA.inp)
**DIRAC
.TITLE
CO molecule
.WAVE FUNCTION
.4INDEX
**WAVE FUNCTION
.SCF
*SCF
. CLOSED
14
. ERGCNV
1E-12
**MOLTRA
.ACTIVE
all
*END OF INPUT
   We run DIRAC with the following command:
```

```
pam --inp=TRA --mol=CO --get="MRCONEE MDCINT"
```

Note that the --get option is used so that pam exports the transformed molecular integrals (stored in the MRCONEE and MDCINT files) from DIRAC's scratch directory to the working directory.

In some cases, it can be convenient to rename the files containing the integrals (this is not mandatory):

```
mv MRCONEE MRCONEE-C1
mv MDCINT MDCINT-C1
   Now the EXP-T input file input-C1 is set up:
# CO molecule, CCSD/cc-pVDZ
# nonrelativistic, symmetry C1
# task title (of type "string")
title "CO/CCSD/cc-pVDZ"
# print level, default medium
print medium
# max number of iterations
maxiter 30
# convergence threshold (by cluster amplitudes)
conv 1e-9
# target Fock space sector
sector OhOp
# CC model: ccsd, ccsdt-1, etc
model ccsd
# import integrals from
integrals MRCONEE-C1 MDCINT-C1
To run EXP-T, enter:
expt.x input-C1
```

The string with the correlation energy can be found at the end of the output. Note that by default EXP-T flushes all output simply to stdout.

```
SCF reference energy = -112.820480227130

CCSD correlation energy = -0.298117912040

Total CCSD energy = -113.118598139170
```

5.2 Simple atomic FS-CCSD calculation: the Pb atom

For input/output files see examples/Pb_fs_ccsd

We now turn to simple atomic relativistic calculations employing the FS-CCSD approach. Since wave functions of most low-lying electronic states of one-electron atoms and atomic ions are effectively single-reference, it is not very interesting to apply the multireference methodology. In contrast, electronic states of the "two-electron" atoms can hardly be simulated using any other approaches except those based on the multireference wave functions. Consider for example the $6s^26p^2$ low-lying states of the Pb atom. The reference state will be Pb²⁺ $6s^2$, and one should select 6p-spinors as the active ones (overall 6 spinors).

Pb atom is a purely relativistic system, and one should use four-component Hamiltonian or relativistic pseudopotential approach. The latter allows one to incorporate bulk of the Breit interaction and QED corrections. Here the semilocal valence version of the N.S. Mosyagin's RECP for Pb is used (see [22] and references therein).

Input file for the FS-CCSD calculation will be as follows:

```
sector 0h2p
nactp 6
maxiter 500
memory 110 gb
conv 1e-7
disk_usage 0
nthreads 8
model ccsd
```

After the FS-CCSD calculation is finished, one can compare the calculated FS-CCSD excitation energies of Pb with their experimentally measured counterparts (7819, 10650, 21458 and 29467 cm⁻¹, respectively [23]):

Le	evel	Re(eigenvalue)	•		•	Rel eigv, eV	Rel eigv, cm-1	deg	symmetry
				•					
@	1	-0.8258538771				0.000000000	0.000000	1	0g
@	2	-0.7916687782				0.9302239316	7502.761992	3	0g 1g+ 1g-
@	3	-0.7784232250				1.2906537953	10409.824894	5	0g 1g+ 1g- 2g+ 2g-
@	4	-0.7298818976				2.6115306047	21063.414836	5	0g 1g+ 1g- 2g+ 2g-
@	5	-0.6892150215				3.7181326781	29988.762480	1	0g

The remaining error can be significantly reduced by inclusion of triple excitations [8].

5.3 Electronic states of the HgH²⁺ molecular ion. Energy denominator shifts technique

For input/output files see examples/HqH2+_shifts

In almost every problem involving the exploration of potential energy surfaces using the FS-CC method in a fairly wide range of internuclear distances, the intruder state problem arises. The presence of intruder states usually manifests itself as (but does not not simply arise from!)

the appearance of small energy denominators in amplitude equations (see Sec. ?? for details); the presence of small denominators makes the iterative procedure unstable. A straightforward solution of the intruder-state problem can be obtained by modifying (shifting) all energy denominators in the problematic Fock space sector(s) in such a way that the shifts are significant for ill-defined (nearly zero or positive) energy denominators and remain negligibly small for large negative denominators [2, 3]. Note, however, that this does not imply any modification of the right-hand side of the amplitude equations and, thus, introduces an additional (although controllable) approximation.

The denominator shift technique is implemented in EXP-T; different formulas are available (see Sec. 4, keyword shifttype). The example discussed below employs the simplest *real* shifts.

Consider the HgH^{2+} molecular ion at R=2.0 bohr. The FS-CC equations cannot be solved at all without using denominator shifts. Let the active space be comprised of the 8 lowest virtual spinors. The EXP-T input file will look like this:

```
# Test:
# (1) HgH2+ ion, FSCC scheme: HgH3+ -> HgH2+
# (2) hamiltionian: 2-comp gatchina ECP
# (3) symmetry Cinfv
# (4) sector Oh1p
title "Test HgH3+ -> HgH2+/gatchina ECP/Cinfv"
maxiter 200
conv 1e-9
active -10 -0.5
sector Oh1p
# import integrals from:
integrals MRCONEE-Cinfv MDCINT-Cinfv
# shifts of denominators:
# real shift:
shifttype real
# shift parameters:
# 1. <n> -- attenuation parameter (here: 3)
# 3. <S1> -- shift for all S1 amplitudes (here: -0.5)
# 3. <S2> -- shift for all S2 amplitudes (here: -1.0)
shift 0h1p 3 -0.5 -1.0
```

Approaches to the selection of shift parameters are discussed in [2, 3]. You can also perform Pade extrapolation of the series of effective Hamiltonian matrices to the zero shift limit in order to minimize distortions introduced by the shift [3] (see also Sec. 6.1).

5.4 Intermediate Hamiltonian for incomplete main model spaces: C atom

For input/output files see examples/C_atom_ih-imms

Let us consider low-lying electronic states of the C atom. Using the NIST ASD database https://www.nist.gov/pml/atomic-spectra-database we find that for the states below $80000~\rm cm^{-1}$ the following configurations dominate in wave functions:

- $2s^22p^2$;
- $2s^12p^3$;
- $2s^22p^13s^1$;
- $2s^22p^13p^1$;
- $2s^22p^13d^1$;
- $2s^22p^14s^1$.

Within the Fock space coupled cluster formalism, electronic states dominated by the $2s^12p^3$ configuration (e. g. the $^5S_2^o$ state, 33735 cm⁻¹) can be accessed only in 1h3p or 0h4p sectors. Implementation of corresponding CC models is not available to date, thus we will not consider these states in the further discussion. In contrast, other states of C can be easily obtained in the 0h2p Fock space sector provided that the $2s^2$ state of the C^{2+} ion is considered as the Fermi vacuum. For the sake of simplicity we will not also consider electronic states involving shells with principal quantum number n=4 (the generalization of the IH-IMMS procedure to such states is straightforward).

At this point we have several useful observations:

- active space should comprise at least 2p, 3s, 3p, 3d spinors of the C^{2+} ion;
- only the 2p shell can be doubly occupied in the target electronic states;
- all target electronic states involve configurations with at least one 2p-electron;
- we can expect that electronic states dominated by the $3s^2$, $3s^13p^1$, etc configurations will be very high in energy (and even lie above IP). These states arise inevitably in the 0h2p sector since the "classical" FS-CC is formulated for quasi complete model spaces. Moreover, these *intruder states* will destroy convergence of amplitude equations;
- the linear span of these "malicious" determinants can be considered as the *intermediate* space;
- main model space should include only determinants belonging to the $2p^2$, $2p^13s^1$, $2p^13p^1$, $2p^13d^1$ configurations. These determinants will be dominating in the model vectors representing target electronic states of C.

Now we start actual calculation with the molecular integral transformation: (C.mol)

```
C atom
doubly augmented triple-zeta basis set (d-aug-cc-pVTZ)
symmetry Dinfh
    1
        6.
С
     0.000000000
                     0.000000000
                                      0.000000000
LARGE EXPLICIT
                  4
                        2
                             2
                                  1
                                       1
# s functions
F 10 2
       8236.00000000
                         0.00053100
                                      -0.00011300
       1235.00000000
                         0.00410800
                                      -0.00087800
        280.80000000
                         0.02108700
                                      -0.00454000
         79.27000000
                         0.08185300
                                      -0.01813300
         25.59000000
                         0.23481700
                                      -0.05576000
          8.99700000
                         0.43440100
                                      -0.12689500
          3.31900000
                         0.34612900
                                      -0.17035200
          0.90590000
                         0.03937800
                                       0.14038200
                        -0.00898300
          0.36430000
                                       0.59868400
          0.12850000
                         0.00238500
                                       0.39538900
F
  4 0
          0.90590000
          0.12850000
          0.04402000
          0.01510000
# p functions
F
   5
     1
         18.71000000
                         0.01403100
          4.13300000
                         0.08686600
          1.20000000
                         0.29021600
          0.38270000
                         0.50100800
          0.12090000
                         0.34340600
F
   4 0
          0.38270000
          0.12090000
          0.03569000
          0.01050000
# d functions
  4 0
          1.09700000
          0.31800000
          0.10000000
          0.03140000
# f functions
F 3 0
          0.76100000
          0.26800000
```

0.09440000

FINISH

(TRA.inp)

- **DIRAC
- .TITLE

Carbon atom

- .4INDEX
- .WAVE FUNCTION
- **HAMILTONIAN
- .X2Cmmf
- .GAUNT
- **WAVE FUNCTION
- .SCF
- *SCF
- . CLOSED
- 4 0
- . ERGCNV
- 1E-12
- **MOLTRA
- .SCHEME
- 4
- .ACTIVE
- all
- *END OF INPUT

Command line:

Let us now look at the DIRAC output. We find that one-electron energies of our active spinors are (in a.u.):

$$\varepsilon_{2p} \sim -0.86$$

$$\varepsilon_{3s} \sim -0.36$$

$$\varepsilon_{3n} \sim -0.28$$

$$\varepsilon_{3d} \sim -0.23$$

We split these spinors into two subsets. Spinors from the first subset can be doubly occupied in the main model space determinants, and spinors from the second subset can be only singly occupied. In some cases it can be useful to define the third subset of spinors which must be unoccupied in all main determinants. This procedure resembles that used to define active space in the RASSCF method. For the C atom these subsets are:

- 1. 2p;
- 2. 3s, 3p, 3d.

In terms of these two subspaces of active spinors the main model space determinants can be defined using the following directives inside the ih_imms block in the EXP-T input file:

- main_occ 2 0: both electrons occupy spinors from the first subspace, the $2p^2$ configuration;
- main_occ 1 1: one electron occupy spinor from the first subspace (2p), the other one occupy spinor from the second subspace (3s, 3p or 3d). These determinants correspond to the $2p^13s^1$, $2p^13p^1$ and $2p^13d^1$ configurations.

We will rely on the automatic determination of the frontier_energy parameter (by default, the uppermost energy of main model space determinants is used). Finally, the EXP-T input file will be as follows:

```
sector Oh2p
model ccsd
nactp 24
x2cmmf
# intermediate Hamiltonian parameters.
# spinor subspaces are defined via energy ranges (in a.u.)
ih_imms
  sectors Oh2p
  subspace energy -1.0 -0.5 # 2p spinors
 subspace energy -0.5 -0.2 # 3s, 3p, 3d spinors
 main_occ 2 0
                            # 2p^2 determinants
 main_occ 1 1
                             # 2p3s, 2p3p, 2p3d determinants
end
# memory
memory 4 gb
disk_usage 0
# convergence
maxiter 300
diis 300
conv 1e-9
# print compositions of model vectors in terms of
# effective occupation numbers of active spinors
print "eff config"
```

relativistic CC model

Let us take a look at the table with calculated energy levels in the 0h2p sector in the EXP-T output file:

```
Level Re(eigenvalue) ...Rel eigv, cm-1 % main deg symmetry
-----
@ 1 -1.2955562077 ... 0.000000 100.0 1 0g
```

```
0
     2
         -1.2954943219 ...
                                 13.582372
                                               100.0
                                                       3
                                                            0g 1g+ 1g-
0
     3
         -1.2953712703 ...
                                 40.589069
                                               100.0
                                                       5
                                                            0g 1g+ 1g- 2g+ 2g-
     4
@
         -1.2503086362 ...
                               9930.694080
                                                99.9
                                                       5
                                                            0g 1g+ 1g- 2g+ 2g-
     5
@
         -1.1923569864 ...
                              22649.611052
                                                99.8
                                                       1
                                                            0g
     6
         -1.0208724411 ...
                              60286.118415
                                                99.9
0
                                                       1
                                                            0u
     7
@
         -1.0207910577 ...
                              60303.980008
                                                99.9
                                                       3
                                                            0u 1u+ 1u-
0
     8
         -1.0206248742 ...
                              60340.453071
                                                99.9
                                                       5
                                                            0u 1u+ 1u- 2u+ 2u-
0
     9
         -1.0089659280 ...
                              62899.295993
                                                99.9
                                                       3
                                                            0u 1u+ 1u-
    10
         -0.9902128248 ...
                                               100.0
                                                       3
0
                              67015.126416
                                                            0g 1g+ 1g-
@
                                               100.0
    11
         -0.9824896577 ...
                              68710.165670
                                                       3
                                                            0g 1g+ 1g-
0
    12
         -0.9824017917 ...
                              68729.450018
                                               100.0
                                                       5
                                                            0g 1g+ 1g- 2g+ 2g-
                                                       7
0
    13
         -0.9822669729 ...
                              68759.039338
                                               100.0
                                                            0g 1g+ 1g- 2g+ 2g- 3g+ 3g-
0
    14
         -0.9751409822 ...
                              70323.013514
                                               100.0
                                                       3
                                                            0g 1g+ 1g-
0
    15
         -0.9723223910 ...
                              70941.622775
                                               100.0
                                                       1
                                                            0g
0
    16
         -0.9722734205 ...
                              70952.370551
                                               100.0
                                                       3
                                                            0g 1g+ 1g-
    17
0
         -0.9721863073 ...
                              70971.489684
                                               100.0
                                                       5
                                                            0g 1g+ 1g- 2g+ 2g-
0
    18
         -0.9616074654 ...
                              73293.277111
                                                99.8
                                                       5
                                                            0g 1g+ 1g- 2g+ 2g-
0
    19
         -0.9497229655 ...
                              75901.623363
                                                99.9
                                                       1
                                                            0g
@
    20
                                               100.0
         -0.9322863070 ...
                              79728.527553
                                                       5
                                                            0u 1u+ 1u- 2u+ 2u-
@
    21
         -0.9307044048 ...
                              80075.714943
                                               100.0
                                                       5
                                                            0u 1u+ 1u- 2u+ 2u-
0
    22
         -0.9306318565 ...
                              80091.637473
                                               100.0
                                                       7
                                                            0u 1u+ 1u- 2u+ 2u- 3u+ 3u-
@
    23
         -0.9305012392 ...
                              80120.304651
                                               100.0
                                                       9
                                                            0u 1u+ 1u- 2u+ 2u- 3u+ 3u- ...
                                                            0u 1u+ 1u-
0
    24
         -0.9303011549 ...
                              80164.218067
                                               100.0
                                                       3
    25
         -0.9302612163 ...
                                               100.0
                                                            0u 1u+ 1u- 2u+ 2u-
0
                              80172.983578
                                                       5
                                                       7
                                                            0u 1u+ 1u- 2u+ 2u- 3u+ 3u-
@
    26
         -0.9302131715 ...
                              80183.528205
                                               100.0
    27
                                                       7
                                                            0u 1u+ 1u- 2u+ 2u- 3u+ 3u-
@
         -0.9264080659 ...
                              81018.652356
                                               100.0
@
    28
         -0.9244722663 ...
                              81443.511260
                                               100.0
                                                            0u 1u+ 1u- 2u+ 2u-
                                                       5
0
    29
         -0.9243950805 ...
                              81460.451580
                                               100.0
                                                       3
                                                            0u 1u+ 1u-
@
    30
         -0.9243547369 ...
                              81469.305985
                                               100.0
                                                            0u
                                                       1
                                                99.9
         -0.9218925156 ...
                                                            0u 1u+ 1u-
0
    31
                              82009.701090
                                                       3
         -0.5545617414 ... 162629.487352
0
    32
                                                 9.6
                                                       1
                                                            0g
0
    33
         -0.5088229529 ... 172667.991098
                                                 0.0
                                                       1
                                                            0u
         -0.5088105565 ... 172670.711776
                                                       3
0
    34
                                                 0.0
                                                            0u 1u+ 1u-
    35
         -0.5087857581 ... 172676.154402
                                                       5
                                                            0u 1u+ 1u- 2u+ 2u-
                                                 0.0
 Ionization potential Oh2p -> Oh1p = ... = 10.9819 eV =
                                                                88575.16 cm<sup>-1</sup>
```

The column named "% main" contains total weight of main model space determinants in model vectors describing the given electronic states. It can be seen that these weights are very close to 100% for all target (or main) electronic states. Intermediate states (with weighs of main model space determinants not exceeding several percents) are also obtained as a by-product. However, they can hardly be considered as physical ones (compare their energies with the IP₁ $\sim 90000~\rm cm^{-1}$). The greater the weight of main model space determinants, the less the dependence on the parameters of the applied denominator shifting scheme.

195766.56 cm⁻¹

Ionization potential $Oh1p \rightarrow Oh0p = ... = 24.2720 \text{ eV} =$

It is worth noting that in some cases convergence of amplitude equations can be achieved only by the introduction of additional active spinors, which are unoccupied in all main model space determinants. These spinors serves as buffer ones. In calculations accounting for triples such an extension of active space can result in an increase of computational cost. So it is recommended to thoroughly choose the upper bound of the active space to make it as small as possible. However, this is absolutely irrelevant for calculations employing the CCSD model only. Introduction of additional buffer states also greatly improves stability of energy levels with respect to the shifting parameters (see Figures 3 and 4 in [4] for the case of Lu⁺ electronic states).

5.5 Finite-field transition dipole moments calculations: Rb atom

For input/output files see examples/Rb_atom_TDMs

See [15, 24, 5] for detailed discussions on the finite field technique (FF) for transition dipole moments. The algorithm for any FF calculation is as follows:

- solve the HF problem for the non-perturbed Hamiltonian (no external fields);
- transform the molecular integrals to the basis of "non-perturbed" one-electron functions;
- perform two FS-CC calculations with these integrals at external fields strengths -F and +F;
- estimate TDMs are estimated using the finite-difference-type formula:

$$\boldsymbol{d}_{\eta,if}^{FF} = (E_i - E_f) \lim_{\substack{F_{\eta} \to 0 \\ F_{\eta'} = 0, \eta' \neq \eta}} \frac{\langle \tilde{\psi}_i^{\perp \perp} (-F_{\eta}) | \tilde{\psi}_f (+F_{\eta}) \rangle}{2F_{\eta}} \quad (\eta = x, y, z), \tag{5}$$

where $\tilde{\psi}_i$ and $\tilde{\psi}_f$ are model vectors corresponding to **i**nitial and **f**inal electronic states, respectively.

Consider the FF calculation of TDMs for the ${}^2S_{1/2} \leftrightarrow {}^2P^o_{1/2}$ and ${}^2S_{1/2} \leftrightarrow {}^2P^o_{3/2}$ transitions in the Rb atom. The basis set and effective core potential employed were taken from [24]; basis set was reduced to [7s7p5d3f2g]. The simplest active space required to describe ${}^2S_{1/2}$, ${}^2P^o_{1/2}$ and ${}^2P^o_{3/2}$ states of Rb comprised 5s- and 5p-spinors (overall 8 spinors, or 4 Kramers pairs).

Firstly, we perform the Hartree-Fock calculation and subsequent transformation of integrals. Note that transformed integrals of one-electron properties (in this case, the dipole moment operator) are stored in the MDPROP unformatted file which should be copied from DIRAC scratch to the working directory:

Other integrals are saved to the MRCONEE (one-electron Fock) and MDCINT (two-electron) unformatted files. Note that the .PRPTRA option as well as the **PROPERTIES/.DIPOLE specification is required in the TRA.inp input file:

```
**MOLTRA
.PRPTRA
.SCHEME
4
.ACTIVE
energy -100.0 20.0 0.5
**PROPERTIES
.DIPOLE
```

These lines are used to ask DIRAC to calculate and transform dipole moment integrals (these integrals are used later to construct the perturbation due to the external field). Also note that the external uniform electric field lowers spherical symmetry of the atom to $C_{\infty v}$.

Now we are ready to run FSCC calculations in EXP-T. At this step we use the oneprop directive to add an external field F contribution to the Fock operator:

$$f_{pq}^{'} = f_{pq} + (d_x)_{pq}F_x + (d_y)_{pq}F_y + (d_z)_{pq}F_z$$

Here we use only the d_z component, stored in the MDPROP file in the ZDIPLEN record. The operator which is added to the Fock matrix must be fully symmetric. The first EXP-T input file (for field strength -F):

```
# title for the task (of type "string")
title "Rb atom -- relativistic EA-CCSD calculation"

# max number of iterations
maxiter 50

# convergence threshold (by cluster amplitudes)
conv 1e-9

# target Fock space sector
sector 0h1p

# active space specification: 8 lowest virtual spinors (5s,5p1/2,5p3/2)
nactp 8

# add the external field contribution to the Fock matrix:
# Fock = Fock + d_z * F_z
# (F_z = 1e-5)
oneprop -1e-5 0.0 "ZDIPLEN"
```

For the other field strength +F it is convenient to reuse two-electron integrals, which are already sorted (the **reuse** option). One-electron integrals depend on the finite perturbation parameter and must be recomputed. Cluster amplitudes converged at -F can also serve as an excellent initial guess for the calculation at +F. This will lead to significant computational savings. Use the --no-clean option to prevent the deletion of files containing sorted two-electron integrals and converged amplitudes.

The second EXP-T input file (for field strength +F):

```
title "Rb atom -- relativistic EA-CCSD calculation"
maxiter 50
conv 1e-9
sector 0h1p
nactp 8

reuse 2-integrals amplitudes
oneprop +1e-5 0.0 "ZDIPLEN"
```

Run EXP-T and save formatted files HEFF with effective Hamiltonian matrices to the working directory:

```
expt.x --no-clean input_F- | tee ccsd_F-.out
mv scratch/HEFF HEFF1
expt.x --no-clean input_F+ | tee ccsd_F+.out
mv scratch/HEFF HEFF2
```

The electronic spectrum is given at the end of the EXP-T output file. Corresponding lines begin with the @ symbol:

Le	vel	Re(eigenvalue)	•	•	.Re	el eigv,	cm-1	deg	symmetry
0	1	-0.1531234146				0.0	00000	2	1/2+ 1/2-
0	2	-0.0959179975				12555.13	37832	2	1/2+ 1/2-
0	3	-0.0948635671				12786.5	58570	2	1/2+ 1/2-
0	4	-0.0948635506				12786.50	62182	2	3/2+ 3/2-

(compare with experimental values $12578.950 \text{ cm}^{-1}$ and $12816.545 \text{ cm}^{-1}$ [23]; note that the $^2P^o_{3/2}$ state is split in the external electric field).

Finally, we turn to the calculation of transition moments. The input file for the heffman.x utility is as follows (see Sec. 6.1 for details and explanation):

file: HEFF1
file: HEFF2
sector: Oh1p

rep: 1

step: 0.00002

print: 1

Run heffman.x:

heffman.x < ff.inp | tee ff.out

Transition moments and Einstein coefficients:

tran	siti	on	energy, cm^-1		d ^2	d	osc str	A,10^6 s-1
1	->	2	12555.138		2.627905	1.62108158	0.100220	10.537579
1	->	3	12786.559		5.173064	2.27443710	0.200921	21.911672
2	->	3	231.421		0.000000	0.00001167	0.000000	0.000000

Note: this is only the $|d_z|$ component of the full transition moment d. It should be mentioned that the small, but non-zero value of |d| in the last line is an artifact of the FF approach.

Taking into account the degeneracy of the ${}^{2}P_{3/2}^{o}$ state and the spherical symmetry of the Rb atom, the full transition moments |d| and full Einstein coefficients can be calculated:

$$A(^{2}S_{1/2} \leftarrow {^{2}P_{1/2}^{o}}) = 31.6 \text{ s}^{-1}$$

 $A(^{2}S_{1/2} \leftarrow {^{2}P_{3/2}^{o}}) = 32.9 \text{ s}^{-1}$

(compare with experimental values 34.0 s^{-1} and 37.0 s^{-1} [23]).

Results can be improved significantly by extending the basis set and active space [24].

5.6 «High-spin» CCSD for open shell systems

For input/output files see examples/02_highspin

The example considered in this subsection is borrowed from the DIRAC program manual [13]. Let us calculate the energy of the ground (triplet) state of the O₂ molecule.

The DIRAC input file contains directives for the HF calculation and subsequent integrals transformation:

```
(O2.mol)
```

```
test of open-shell CCSD calculation
molecular oxygen at eq distance taken from NIST
automatic symmetry detection: DIRAC will identify the Dinfh group
C
       8.
             2
0
       0.0
                 0.0
                           0.60376
                 0.0
                          -0.60376
       0.0
LARGE BASIS cc-pVDZ
FINISH
   (TRA.inp)
!
! calculation of the triplet ground state of molecular oxygen
! (Dirac-Coulomb Hamiltonian)
**DIRAC
.TITLE
molecular oxygen -- ground state
.WAVE F
.4INDEX
**WAVE FUNCTIONS
.SCF
*SCF
.CLOSED SHELL
 6 8
.OPEN SHELL
```

```
1
2/4,0
*END OF
```

DIRAC writes spinor occupation numbers to the MRCONEE unformatted file, but only spinors belonging to closed shell (the .CLOSED_SHELL keyword) are labeled as occupied. This is why EXP-T will calculate the energy for the ${\rm O}_2^{2+}$ cation without manually specified occupation numbers.

EXP-T provides two ways to set the occupation numbers of spinors:

• by setting the number of occupied spinors in each irrep (the occ_irreps keyword). For the example under consideration:

```
occ_irreps 3 2 0 1 0 0 0 0 0 0 0 0 0 0 0 2 2 1 1
```

Note that the 1s-orbitals of oxygen atoms are frozen here, and the corresponding electrons should not be accounted for in the occ_irreps string. In order to set the number of electrons in each irrep correctly, we recommend starting with a calculation for default occupation numbers and finding strings with an irrep order in the output:

```
0 1/2g+
1 1/2g-
2 3/2g+
3 3/2g-
4 5/2g+
5 5/2g-
...
16 1/2u+
17 1/2u-
18 3/2u+
19 3/2u-
20 5/2u+
21 5/2u-
...
```

For the example discussed:

- 3 electrons occupy the lowest spinors belonging to the 1/2g+ irrep;
- 1/2g-, 1/2u+, 1/2u-: 2 electrons in each irrep;
- 3/2g-, 3/2u+, 3/2u-: 1 electron in each irrep.
- you can also specify the occupation number for *each* spinor explicitly:

To identify spinors which should be populated in the electronic state under consideration, we again recommend starting the calculation with the default occupation numbers, then interrupting the task, finding information about spinors and restarting the calculation with the correct occupation numbers.

Note that, in fact, non-canonical orbitals are used, therefore the Fock matrix is not diagonal and the vacuum determinant energy $\langle \Phi_0 | H | \Phi_0 \rangle$ is not equal to the SCF energy value obtained before:

```
Total SCF energy = -149.686661451842
...

sorting one-electron integrals ...

Fock matrix reconstruction ...

SCF energy (energy of reference determinant) = -149.718144633810 a.u.

SCF energy (energy of reference determinant) was updated:

old energy = -149.686661451842 a.u.

new energy = -149.718144633810 a.u.

...

SCF reference energy = -149.718144633810

CCSD correlation energy = -0.366958682652

Total CCSD energy = -150.085103316462
```

6 Utility programs

6.1 heffman.x – manipulations with effective Hamiltonian matrices

The heffman.x program is designed to perform manipulations with FS-RCC effective Hamiltonians matrices. Implemented features are:

- the finite-field technique for transition dipole moments calculations [15, 24, 5];
- extraction of spin-orbit interactions from full-relativistic models by projection [2];
- Pade extrapolation of series of effective Hamiltonians.

heffman.x reads matrices of effective Hamiltonians from formatted files supported by both EXP-T and the latest versions of DIRAC.

- to compute finite-field transition moments [24, 5] two files with effective Hamiltonians must be provided, calculated at field 1 and field 2;
- to separate spin-orbit interactions by projection, two files with effective Hamiltonians must be provided, the first for spin-orbit (nearly) switched off and the second for spin-orbit switched on.

In this case you cannot freeze spinors at the FSCC step!

• to perform Pade extrapolation of the series of effective Hamiltonians, you have to put a sequence of effective Hamiltonians to a single file using the cat command:

```
cat heff_n_eq_1 heff_n_eq_2 heff_n_eq_3 > some_file_name
```

Recall that each Hamiltonian is calculated at its own attenuation parameter n, see Sec. ??. You can also perform extrapolation before TDM or SO calculation, just use such "composite" files with effective Hamiltonians.

Additional input file with options must be redirected to stdin:

heffman.x < input_file

heffman.x input files are very similar to EXP-T input files, but keywords are separated from their arguments with the ':' symbol, and the symbol '#' at the beginning of a comment must be placed in the first position of the line.

List of keywords:

file: file with the first effective Hamiltonian or with a sequence of heff's to be extrapolated, e.g.

- heff for the first field value:
- heff for the spin-orbit interaction (nearly) switched off;
- simply heff you want to get the eigenstates;
- sequence of heff's you want to Pade-extrapolate.

Syntax: file: <string path-to-first-heff>

If you are doing finite-field transition moment calculations or SO separation, you have to supply the second file (see the next item).

heff for another field value, or heff for the spin-orbit switched on, respectively. If there is no second file, the effective Hamiltonian eigenvectors and eigenvalues are written to the formatted file HEFFEVF.

Syntax: file: <string path-to-second-heff>

sector: the Fock space sector you are interested in (heff files for higher sectors contain several heff matrices).

Syntax: sector: <H>h<P>p

Example: 0h1p, 1h0p, 2h0p, 0h2p, 1h1p.

rep: irreducible representation you are interested in. As a rule, rep: 1 for the fully symmetrical rep.

Syntax: rep: <integer n>

Enumeration of irreps begins from 1!

main: for the FF-TDM calculations or Pade extrapolation: simply the number of lowest states or the range of states of given sector and symmetry you want to see in the output file. For the SO extraction by the projected approximant construction [3] this keyword defines the subspace of states used for projection.

Syntax: main: <integer number> or main: <integer number1>-<integer number2>.

Default: all states.

Example: main: 10 (equivalently, main: 1-10)

[L/M] (optional) to perform Pade extrapolation, you have to indicate which Pade approximant [L/M] should be constructed ([0/1], [1/1], [0/2] etc). You should have L+M+2 effective Hamiltonian matrices to build the [L/M] approximant. You can cut off all states above subspace done by the main keyword (see above); to do this, add the cut keyword.

Syntax: [L/M] [cut]

Default: Padé extrapolation disabled

Example: [0/1] cut

step: only for FF calculations. Difference of field strengths (a.u.) in two finite-field calculations. The presence of this line activates the FF TDM calculation mode (do not use this keyword in other types of calculations).

Syntax: step: <real value>

scale: only for SO calculations. Fraction of effective spin-orbit retained in "scalar" calculations to get good relativistic symmetries (typically of order 10⁻⁴. Omit this line if you do not want to diabatize anything.

Syntax: scale: <real value>

ground: ground state energy (a.u.). Useful only when you have to compute energies with

respect to some state (normally ground) which belongs to another rep or sector.

Syntax: ground: <real energy>

Default: ground: 0.0

print: print level. print: 0 or print: 1 are normally ok; enlarging this number, you

can get any amount of superfluous information listed.

Syntax: print: <integer level>

Default: print: 1

Bloch activates the computation of finite-field transition dipoles with Bloch effective

Hamiltonian eigenvectors (note: TDM matrices will be non-Hermitian, this has no physical meaning). Otherwise, ff computations are performed with des Cloizeaux

effective Hamiltonian eigenvectors ("hermitization").

Syntax: Bloch

Default: "hermitization" is enabled in finite-field calculations

When using any of the techniques implemented in the heffman.x program, please, cite the corresponding paper:

• finite-field transition dipole moment calculations: [5]

• quasidiabatization by projection and SO extraction: [2]

• Pade extrapolation: [3]

6.2 expt_transform_natorb.x-transformation of quasi-natural spinors to the AO basis

Having model vectors $|\tilde{\psi}_i\rangle$ we can construct approximate density $d_{pq} = \langle \tilde{\psi}_i | a_p^\dagger a_q | \tilde{\psi}_i \rangle$ and transition density matrices $d_{pq} = \langle \tilde{\psi}_i | a_p^\dagger a_q | \tilde{\psi}_j \rangle$ and then calculate "quasi-natural" (transition) molecular spinors (NS and NTS, respectively). These operations can be performed by the EXP-T itself (see Sec. 4, keyword natorb). EXP-T stores NS and NTS expanded in the basis of molecular spinors in formatted files named NATORB*.dat. To use visualization tools one have to transform NS or NTS to the AO basis. This transformation can be performed by the expt_transform_natorb.x utility.

expt_transform_natorb.x requires two command-line arguments:

- path to the DIRAC output file containing molecular spinors expanded in the basis of atomic orbitals;
- path to the NATORB*.dat formatted file produced by EXP-T and stored in its scratch directory (contains natural spinors expanded in the basis of molecular spinors).

To ask DIRAC to print molecular spinors (vectors) to the output, enable the <code>.ANALYZE</code> module in the **DIRAC section and then add the following text to the **ANALYZE section of the DIRAC <code>.inp</code> input file:

- **ANALYZE
- .PRIVEC
- *PRIVEC
- .AOLAB
- .VECPRI

all

(.AOLAB = "print vectors in the AO basis"). Note that for the groups with the inversion one must add the second string specifying the range of molecular spinors (e.g. all).

Running expt_transform_natorb.x (for example):

expt_transform_natorb.x dirac.out NATORB_Oh1p_1:1.dat > nts.out

The transformed spinors are flushed to stdout again in the DIRAC four-column format and can be redirected to file.

6.3 expt_diatomic.x - rotational-vibrational levels and properties of diatomic molecules

If one have a potential energy curve U(R) for a diatomic molecule, one can solve the radial Schrödinger equation and obtain rovibrational levels and wavefunctions as well as any other properties (molecular constants, transition moments, Franck-Condon factors). The radial Schrödinger equation is written as:

$$\left[-\frac{\hbar^2}{2\mu} \frac{d^2}{dR^2} + \frac{J(J+1)}{2\mu R^2} + U(R) \right] \psi_{vJ}(R) = E_{vJ} \ \psi_{vJ}(R), \tag{6}$$

where

- $\mu = \frac{M_1 M_2}{M_1 + M_2}$ reduced mass,
- \bullet v vibrational quantum number,
- J rotational quantum number.

Normally this second-order differential equation is solved numerically using some mesh of radial points (see, for example, [25, 26, 27]). Within the current implementation, the U(R) function is interpolated using the cubic natural spline. The matrix version of either the Numerov method [28] or the 2nd order finite-difference method [25] is used to perform integration. Only bound states can be treated to the moment.

6.3.1 Input file syntax

The syntax of input files for the solver is quite similar to that of the coupled-cluster program. Single line comments start with the # symbol. The following directives are allowed:

masses of isotopes M_1 and M_2 (in amu).

Syntax: masses <float M1> <float M2>

charge net charge of the molecule Q. For charged molecules the "charge-modified reduced mass" formula [29, 27] is used:

$$\mu = \frac{M_1 M_2}{M_1 + M_2 - m_e Q},\tag{7}$$

where m_e stands for the electron mass. Can be useful for light molecular ions. Syntax: charge <integer Q>

viblevels range $[v_{min}, v_{max}]$ of vibrational levels of interest.

Syntax: viblevels <integer vmin> <integer vmax>

Default: viblevels 0 0

rotlevels range $[J_{min}, J_{max}]$ of rotational levels of interest.

Syntax: rotlevels <integer Jmin> <integer Jmax>

Default: rotlevels 0 0

grid_size number of radial mesh points.

Syntax: grid_size <integer npoints>

Default: grid_size 300

solver integration algorithm:

solver numerov Numerov method [28, 27] (default)

solver fd2 2nd order finite-difference method [25]

mapping meshkov08 <float ra in Angstroms> <float beta>

mapping formula for the radial variable (disabled by default). Reduced variable mapping first proposed by V. V. Meshkov and co-authors [25]:

$$y(r) = \frac{r^{\beta} - r_a^{\beta}}{r^{\beta} + r_a^{\beta}} \quad y \in [-1, 1]$$
 (8)

Mapping is available only for the finite-difference solver.

write_psi flush wavefunctions $\psi_{Jv}(R)$ to the formatted text files (disabled by default).

potential point-wise potential energy curves and/or property matrix element. The first (r) and the second $(U_1(r))$ columns are mandatory. The third (property) and the fourth $(U_2(r))$ columns are optional. If two potentials are specified, transition matrix elements will be calculated. To calculate Franck-Condon factors, use M(r) = 1 in the third column.

Syntax:

```
potential <units of distance> <units of energy>
  <float r> <float U1(r)> [<float M(r)> [<float U2(r)>] ]
   . . .
end

Allowed units of distance are: angstrom, atomic, bohr, au.
Allowed units of energy are: cm-1, cm, atomic, hartree.
```

6.3.2 Example: Franck-Condon factors for the AcOH⁺ molecular ion

To estimate the suitability of the molecule for the direct laser cooling one should calculate vibrational branching ratios and Franck-Condon factors for the working electronic transition. Recently the AcOH⁺ molecular ion was proposed as a candidate for direct laser cooling [30].

This molecule is linear in both the ground (1)1/2 and the first excited (1)3/2 electronic states. If we neglect the anharmonic couplings between stretching and bending modes, the total FCF between the (v'_1, v'_2, v'_3) and the (v''_1, v''_2, v''_3) vibrational states can be factorized:

$$FCF = |\langle v_1'v_3'|v_1''v_3''\rangle|^2 \times |\langle v_2'|v_2''\rangle|^2$$

$$(9)$$

The latter factor is the one-dimensional bending mode FCF and it is expected to be very close to unity since the corresponding sections of potential energy surfaces are perfectly parallel to each other. The overlap intergal over the v_3 stretch mode is expected to be nearly equal to unity and, which is more important, this factor will be the same for the $(v'_1, v'_2, 0)$ and $(v''_1, v''_2, 0)$ vibrational states analyzed here since this stretching mode is not excited in the states under consideration. Thus the total FCF can be approximated with the squared one-dimensional $\langle v'_1|v''_1\rangle$ overlap integral. To calculate this integral one should solve the radial Schrödinger equation, considering the OH group as a pseudoatom.

The input file for the expt_diatomic.x program:

```
#
# calculation of Frank-Condon factors for the (1)1/2-(1)3/2 transition in AcOH+
# potential curves were calculated by the relativistic FS-CCSD(0h1p) method,
# for more details, see:
# A. Oleynichenko, L. V. Skripnikov, A. V. Zaitsevskii, V. V. Flambaum
# Laser-coolable AcOH+ ion for CP-violation searches (2022)
  arXiv:2112.02307 [physics.atom-ph]
#
           Ac
                      OH
masses 227.028000 17.003000
rotlevels 0 0
viblevels 0 5
                       property(r)=1.0
# r
       potential1(r)
                                           potential2(r)
# units:
          dist
                 energy
potential atomic atomic
0.340150702E+01
                -0.451045595E+03
                                    1.0
                                         -0.451004746E+03
```

```
0.349599333E+01
                 -0.451064998E+03
                                     1.0
                                          -0.451025470E+03
0.359047964E+01
                 -0.451079018E+03
                                     1.0
                                          -0.451040834E+03
0.368496594E+01
                 -0.451088543E+03
                                     1.0
                                          -0.451051696E+03
0.377945225E+01
                 -0.451094322E+03
                                     1.0
                                          -0.451058791E+03
0.387393856E+01
                 -0.451096994E+03
                                     1.0
                                          -0.451062746E+03
0.396842486E+01
                 -0.451097103E+03
                                     1.0
                                          -0.451064095E+03
0.406291117E+01
                 -0.451095111E+03
                                     1.0
                                          -0.451063297E+03
0.415739747E+01
                 -0.451091413E+03
                                     1.0
                                          -0.451060740E+03
0.425188378E+01
                 -0.451086341E+03
                                     1.0
                                          -0.451056758E+03
0.434637009E+01
                 -0.451080177E+03
                                     1.0
                                          -0.451051631E+03
0.444085639E+01
                 -0.451073161E+03
                                     1.0
                                          -0.451045598E+03
0.453534270E+01
                 -0.451065491E+03
                                     1.0
                                          -0.451038858E+03
end
```

Run the job:

```
expt_diatomic.x AcOH+_FCF.inp | tee AcOH+_FCF.out
```

At the end of the output file one can find the table with transition moments (and their squares). We select only the FCFs with the wavenumber of the upper state v'' = 0:

J' 3	J,,	v, 1	$_{_{J}},,$		delta e,cm^-1	prop	prop ^2
\$ 0	0	0	0		7279.8734	-0.942947	0.889149
\$ 0	0	1	0		6627.6841	0.325405	0.105888
\$ 0	0	2	0		5979.2731	0.069678	0.004855
\$ 0	0	3	0		5334.3122	0.010337	0.000107
\$ 0	0	4	0		4692.7001	0.001070	0.000001
\$ 0	0	5	0		4054.6587	-0.000079	0.000000

The sum of first three FCFs (see the last column) is equal to 0.9999, thus allowing one to conclude that the closed optical cycle can exist for the AcOH⁺ molecular cation [30].

Appendix A Alphabetical index of keywords

```
active 15
analyt prop 18
arith 19
compress 25
conv 20
cuda 25
damping 20
degen thresh 12
density 18
diis 20
disk_usage 24
div thresh 20
flush 22
ih imms 15
ih_imms: frontier_energy 17
ih imms: main occ 16
ih imms: npower 17
ih_imms: scale 17
ih imms: sectors 16
ih imms: shift type 16
ih imms: subspace 16
integrals 23
maxiter 20
mdprop 18
memory 23
model 13
mstdm 17
nacth 14
nactp 14
natorb 18
hermit??
nroots 19
nthreads 25
occ 14
occ irreps 14
openmp 25
print 12
reuse 22
roots_cutoff 20
sector 13
shift 21
shifttype 21
skip 22
spinor_labels 13
```

tilesize 24 title 12

txtprop 18 x2cmmf 23

Appendix B Temporary files

During the calculation, EXP-T operates with temporary files stored in the *scratch* directory. We recommend to remove these temporary files after every calculation (except for those planned to be used in the future, like the HEFF files or files with cluster amplitudes). All types of temporary files generated by EXP-T are listed and described below.

- HINT one-electron integrals core Hamiltonian matrix + one-particle part of perturbation (if presented). Recommended for removal after calculation.
- VINT-*-*-* two-electron integrals. These files are generated during the integral sorting stage. Recommended for removal after calculation.

Approximate number of files: $\sim N_b^4$ (N_b – number of blocks into which the whole set of spinors is divided by symmetry and the tilesize parameter)

- *.sb single symmetry block of molecular integrals/cluster amplitudes. It is just a part of a diagram. During the calculation each file is mapped to some structure in RAM. Recommended for removal after calculation.
 - Approximate number of files: $\sim 10 \times N_b^4$ for CCSD, $\sim N_b^6$ for CCSDT-n and CCSDT models can be very large!
- *.dg file containing diagram. Diagram files containing cluster amplitudes (t1c.dg, t2c.dg etc) can be used in subsequent calculations when constructing the initial approximation to amplitudes (see Sec. 4, keyword reuse).
- HEFF formatted file containing effective Hamiltonian matrices. These files can be read by the heffman.x utility program used to perform Padé extrapolation of series of effective Hamiltonians [3], finite-field transition moments calculations [24, 5], spin-orbit coupling calculations [2].
- *DIPLEN formatted files with dipole moment integrals (just its electronic part) in the basis of molecular spinors.
- MVCOEF** binary files containing model vectors expanded in the basis of model determinants. These files are used to organize data flow inside EXP-T in the most convenient and logical manner; model vectors are used for calculation of model-space approximations of TDMs and "quasi-natural" spinors, etc.
- NATORB*.dat formatted text files containing "quasi-natural" spinors expanded in the basis of molecular spinors. These expansions can be transformed to the AO basis with the mknatorb.x utility program (see Sec. 6.2).

Appendix C Symmetry and irreducible representations

This section lists the names of irreducible representations (irreps) used in EXP-T.

- For nonrelativistic groups, irrep is determined by the spatial symmetry (Mulliken notation is used) and the M_s value (for $M_s >= 2$ only symbol 2 is used);
- For relativistic double groups we use Mulliken-type notation (completely inherited from DIRAC);
- For $U(1) = \overline{C_{\infty v}}$ and $U(1) \times C_i = \overline{D_{\infty h}}$ irreps are determined by Ω values.

 C_1

 A_a

A_b

 $A_{-3/2}$

A_+3/2

A_0

 A_4

 A_+1

 $A_{-}1$

 C_2

A_a B_a

A_b B_b

A_-3/2 B_-3/2

A_+3/2 B_+3/2

B_0

A_0

A_2 B_2

A_+1 B_+1

A_-1 B_-1

 C_s

A'_a A"_a

A'_b A"_b

A'_-3/2 A"_-3/2

A'_+3/2 A"_+3/2

A'_0 A"_0"

A'_2 A"_2"

A'_+1 A"_+1

A'_-1 A"_-1

 C_i

Ag_a Au_a
Ag_b Au_b
Ag_-3/2 Au_-3/2
Ag_+3/2 Au_+3/2
Ag_0 Au_0
Ag_2 Au_2
Ag_+1 Au_+1
Ag_-1 Au_-1

 C_{2v}

A1_a B2_a B1_a A2_a A1_b B2_b B1_b A2_b A1_-3/2 B2_-3/2 B1_-3/2 A2_-3/2 A1_+3/2 B2_+3/2 B1_+3/2 A2_+3/2 A1_0 B2_0 B1_0 A2_0 A1_2 A2_2 B2_2 B1_2 A1_+1 B2_+1 B1_+1 A2_+1 $A1_-1$ B2_-1 B1_-1 $A2_{-1}$

 C_{2h}

Ag_a Bg_a Bu_a Au_a Ag_b Bu_b Bg_b Au_b Ag_-3/2 Bg_-3/2 Bu_-3/2 Au_-3/2 Ag_+3/2 Bg_+3/2 Bu_+3/2 Au_+3/2 Ag_0 Bg_0 Bu_0 Au_0 Ag_2 Bg_2 Bu_2 Au_2 Ag_+1 Bg_+1 Bu_+1 Au_+1 Ag_-1 Bg_-1 Bu_-1 Au_-1

 D_2

 A_a B3_a B1_a B2_a A_b B3_b B1_b B2_b A_-3/2 B3_-3/2 B1_-3/2 B2_-3/2 A_+3/2 B3_+3/2 B1_+3/2 B2_+3/2 A_0 B3_0 B1_0 B2_0 A_2 B1_2 B2_2 B3_2 A_+1 B3_+1 B1_+1 B2_+1 $A_{-}1$ B3_-1 B1_-1 B2_-1

```
D_{2h}
        Ag_a
                  B1u_a
                             B2u_a
                                        B3g_a
                                                   B3u_a
                                                               B2g_a
                                                                          B1g_a
                                                                                     Au_a
        Ag_b
                  B1u_b
                             B2u_b
                                        B3g_b
                                                   B3u_b
                                                               B2g_b
                                                                          B1g_b
                                                                                     Au_b
        Ag_-3/2 B1u_-3/2 B2u_-3/2 B3g_-3/2 B3u_-3/2 B2g_-3/2 B1g_-3/2 Au_-3/2
        Ag_+3/2 B1u_+3/2 B2u_+3/2 B3g_+3/2 B3u_+3/2 B2g_+3/2 B1g_+3/2 Au_+3/2
        Ag_0
                  B1u_0
                             B2u_0
                                        B3g_0
                                                   B3u_0
                                                               B2g_0
                                                                          B1g_0
                                                                                     Au_0
        Ag_2
                  B1u_2
                             B2u_2
                                        B3g_2
                                                    B3u_2
                                                               B2g_2
                                                                          B1g_2
                                                                                     Au_2
                             B2u_+1
                                                               B2g_+1
        Ag_+1
                  B1u_+1
                                        B3g_+1
                                                    B3u_+1
                                                                          B1g_+1
                                                                                     Au_+1
        Ag_-1
                  B1u_-1
                             B2u_-1
                                        B3g_-1
                                                   B3u_-1
                                                               B2g_-1
                                                                          B1g_-1
                                                                                     Au_-1
\overline{C_1}
        Α
        a
\overline{C_2}
        1E 2E
        a b
\overline{C_s}
        1E 2E
        a b
\overline{C_i}
        AG AU
        ag au
\overline{C_{2v}}
        1E 2E
        a b
\overline{D_2}
        1E 2E
        a b
\overline{D_{2h}}
```

1Eg 2Eg 1Eu 2Eu ag bg au bu $\overline{C_{\infty v}}$

```
1/2+ 1/2- 3/2+ 3/2- 5/2+ 5/2- 7/2+ 7/2-
 9/2+ 9/2- 11/2+ 11/2- 13/2+ 13/2- 15/2+ 15/2-
17/2+ 17/2- 19/2+ 19/2- 21/2+ 21/2- 23/2+ 23/2-
25/2+ 25/2- 27/2+ 27/2- 29/2+ 29/2- 31/2+ 31/2-
               1-
                            2-
                                  3+
                                        3-
   0
         1+
                     2+
   4-
         5+
               5-
                     6+
                           6-
                                  7+
                                        7-
                                              8+
         9+
               9-
   8-
                    10+
                           10-
                                 11+
                                       11-
                                             12+
  12-
              13-
                           14-
        13+
                    14+
                                 15+
                                       15-
                                             16+
```

 $\overline{D_{\infty h}}$

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