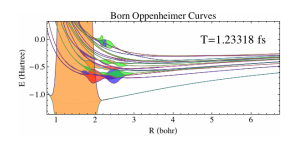
## LBNL-AMO-MCTDHF

Multiconfiguration Time-Dependent Hartree Fock
for ultrafast electronic and nonadiabatic nuclear dynamics
of atoms and molecules in intense laser fields

Version 1.22 Atomic / Diatomic Vibronic / Polyatomic Fixed-Nuclei



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LBNL-AMO-MCTDHF is a suite of codes for Multiconfiguration Time-Dependent Hartree-Fock applied to ultrafast laser dynamics of atoms and molecules. It calculates nonadiabatic electronic and nuclear wave functions for the nonrelativistic Schrödinger equation. Currently it uses the dipole approximation in length and velocity gauge and has options for a variety of laser pulses. It contains analysis and output routines and auxillary scripts for calculating absorption and stimulated emission, populations, total and partial photoionization, wave mixing, and other capabilities. It supports

- Electronic wave functions for atoms (chmctdhf\_atom)
- Vibronic wave functions for diatoms using prolate coordinates (chmctdhf\_diatom)
- Electronic wave functions for polyatomics with fixed nuclei (chmctdhf\_sinc)

#### Future versions will support

- Electronic wave functions for atoms (chmctdhf\_atom)
- Rovibronic wave functions for diatoms using prolate coordinates (chmctdhf\_diatom)
- Rovibronic wave functions for diatoms using modified prolate coordinates (chmctdhf\_diatom2)
- Vibronic wave functions for polyatomics and general Cartesian treatment (chmctdhf\_sinc)

#### The method and equations are described in

- MCTDHF treatment of electronic and nuclear dynamics in diatomic molecules. D. J. Haxton, K. V. Lawler, C. W. McCurdy, Phys. Rev. A 83, 063416 (2011)
- Two methods for restricted configuration spaces within the MCTDHF method. D. J. Haxton and C. W. McCurdy, Phys. Rev. A **91**, 012509 (2015)

#### Some of the largest nuts and bolts are due to

- Exporting A Software Package for Computing Matrix Exponentials. R. B. Sidje, ACM Transactions on Mathematical Software 24, 130 (1998)
- Adiabatic formulation of heteronuclear hydrogen molecular ion. B. D. Esry and H. R. Sadeghpour, Phys. Rev. A 60, 3604 (1999)
- Solving the three-body Coulomb breakup problem using exterior complex scaling. C. W. McCurdy, M. Baertschy, T. N. Rescigno, J. Phys. B. **37**, R137 (2004)
- Grid-based methods for diatomic quantum scattering problems: A finite-element discrete-variable representation in prolate spheroidal coordinates. L. Tao, C. W. McCurdy, T. N. Rescigno, Phys. Rev. A 79, 012719 (2009)
- An efficient basis set representation for calculating electrons in molecules. J. R. Jones, F.-H. Rouet, K. V. Lawler, E. Vecharynski, K. Ibrahim, S. Williams, B. Abeln, C. Yang, D. J. Haxton, C. W. McCurdy, X. Li, T. N. Rescigno, submitted.

The original open-source MCTDH code of H.-D. Meyer *et al.* at Heidelberg has driven our interest in MCTDHF and guided this numerical implementation directly, providing the foundation for the nomenclature and notation used here and in our articles in press, the input style, the variable names, and the layout of this manual. Reviews and guides available at the MCTDH website may help fill in the blanks.

- An efficient & robust integration scheme for the equations of motion of the multiconfiguration timedependent Hartree (MCTDH) method. M. H. Beck and H.-D. Meyer, Z. Phys. D 42, 113 (1997)
- The multiconfiguration time-dependent Hartree method: A highly efficient algorithm for propagating wavepackets. M. H. Beck, A. Jäckle, G. A. Worth, H.-D. Meyer, Physics Reports **324**, 1 (2000)

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This project is dedicated to the memory of Donovan Emil Smith.

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## **Preliminaries**

#### 0.1 Git install

The code is distributed through GitHub.com. Please visit

```
http://commons.lbl.gov/display/csd/LBNL-AMO-MCTDHF
```

to request access to the code. Send us a message via this website. We will receive the message and you will be added to Team Users on the LBNL-AMO-MCTDHF GitHub organization. You will get an email invitation when this happens. Then you can view the repository on your browser

```
https://github.com/LBNL-AMO-MCTDHF/V1
```

and perform a git clone to obtain the code.

```
prompt> git clone https://github.com/LBNL-AMO-MCTDHF/V1 master
```

Instead of git clone, you can download the zipfile

```
https://github.com/LBNL-AMO-MCTDHF/V1/archive/master.zip
```

The README that is visible on the website and included with the code is included below, and includes quick installation instructions for mac.

- \*\* LBNL-AMO-MCTDHF VERSION 1.22 \*\*
- v1.20 multiple cation files for partial photoionization action 17 using numcatfiles, catspffiles, catavectorfiles in namelist &parinp cation.configlist.BIN no longer needed
- v1.19 better improvedquadflag=2 and 3 (orbitals). Excited states may be calculated more easily.
- v1.18 predictor/corrector algorithm modified; results will be different, depending on stepsize par\_timestep
- v1.16 FACTOR OF 1/3 REMOVED FROM PHOTOIONIZATION (Actions 16,17)
  - \*\*\*\* photoionization RESULTS WILL BE 3x GREATER NOW BEWARE \*\*\*\*

the factor of 1/3 is historical. The cross section now reported in column 3 of xsec.spi.dat, xsec.proj.spi.datXXXX, is the quantum cross section in Megabarns for the problem at hand. Perform a 1:2 weighted average of parallel and perpendicular to obtain the cross section averaged over orientations

The ionization and absorption cross section for H2 is about 12~Mb at ion onset for both perpendicular and parallel. Not 4.~He xsec is about 8.~Mos

- ALSO v1.16 linear damping (fttriwindow=1) NEW DEFAULT actions 1,16,17,21 RESULTS WILL BE DIFFERENT (hopefully better)
  Probably better results for absorption/emission, action 21.
  CONSIDER fttriwindow=0, ftwindowpower=2 in namelist &parinp to invoke old windowing function for actions 1,16,17.
- ALSO v1.16 ABSOLUTE UNITS (megabarns) IMPLEMENTED FOR ACTION 21 emission/absorption. Column 9 in ZDipoleft.Dat etc is cross section in Mb. See H2ABSORPTION and HEABSORPTION examples.
- ALSO v1.16 par\_consplit.ne.0 attempted first time. Total MPI parallelization of LBNL-AMO-MCTDHF calculation with par\_consplit.ne.0 and parorbsplit.eq.3 (parorbsplit.eq.3 for polyatomic only)

CHANGES TO PULSE FOURIER TRANSFORM SUBROUTINES IN VERSION 1.10. VARIABLES EFLUXLO, EFLUXHI, NEFLUX, FLUXNBINS, FLUXSINEOPT DEPRECATED. NOTE NEW VARIABLE pulseft\_estep USED FOR Pulseft.Dat OUTPUT ONLY

-----

Please watch the commits to stay notified about bugs and bug fixes: https://github.com/LBNL-AMO-MCTDHF/V1/commits/master

Known bugs appear as issues:
 https://github.com/LBNL-AMO-MCTDHF/V1/issues

\_\_\_\_\_\_

The scripts in the example directories use bash and gnuplot.

Using the bash shell is recommended but may not be necessary.

To begin a bash shell, if it is not your default, simply execute the command prompt> bash

To set bash as your default shell on mac or other linux, execute the command prompt> chsh -s /bin/bash

Lots of scripts use gnuplot. Gnuplot should be installed on your system, if you want to have an easy time seeing the results of the example calculations.

If gnuplot is installed on your system then the command prompt> which gnuplot

should return the location of the executable file. If it returns nothing then it is not installed. The best way to install gnuplot is probably with macports.

Possible workflow described below for mac. The general idea is that you can work entirely within the directory that you clone and to which you pull, not copy the git distribution elsewhere for compilation.

-----

On Mac: minimal demonstration. BIN\*mac\* directories use GFORTRAN. You must have gfortran installed to use the BIN\*mac\* directories. To install gfortran -- which you will need to do if the Makeme step fails -- visit

https://gcc.gnu.org/wiki/GFortranBinariesMacOS

00) git clone https://github.com/LBNL-AMO-MCTDHF/V1 master

or download using a web browser

https://github.com/LBNL-AMO-MCTDHF/V1/archive/master.zip and unzip this file.

You then have a V1 or V1-master directory with the code in it. change to that directory:

cd ./V1

or

cd ./V1-master

PERFORM ALL OTHER STEPS IN THIS V1 DIRECTORY.

0) You must get the examples now separately.

```
cd EXAMPLES-DEPOT
     git clone https://github.com/LBNL-AMO-MCTDHF/EXAMPLES-DEPOT master
          or download using a web browser
     https://github.com/LBNL-AMO-MCTDHF/EXAMPLES-DEPOT/archive/master.zip
          and unzip this file to get the EXAMPLES-DEPOT examples files.
1)
     cd COMPDIRS/BIN.ecs.hermnorm.mac
     ./Makeme
     if this fails, you need gfortran. Also you may then
     cd ../debug.BIN.ecs.hermnorm.mac
     ./Makeme
     cd ../BIN.ecs.hermnorm.mac.mpi
     ./Makeme
     cd ../debug.BIN.ecs.hermnorm.mac.mpi
     ./Makeme
    You can delete the directories in COMPDIRS that you don't want.
2)
    You need to have the code in your path. Safe, temporary way:
2A) cd COMPDIRS/BIN.ecs.hermorm.mac
     export PATH=$PATH:$PWD
2B) Permanent way, system-wide, links in /opt/local/bin, requires root password:
     cd COMPDIRS/BIN.ecs.hermorm.mac
     ./LinkMe
     cd ../debug.BIN.ecs.hermorm.mac
     ./LinkMe debug
     cd ../BIN.ecs.hermorm.mac.mpi
     ./LinkMe mpi
     cd ../debug.BIN.ecs.hermorm.mac.mpi
     ./LinkMe mpi.debug
2C) Permanent way, user-specific, script that edits your .bashrc file
     cd COMPDIRS/BIN.ecs.hermnorm.mac
     ./LinkMeLocal
    You must restart the terminal after step 2C.
    mkdir EXAMPLES
                             (you can also make other directories e.g.
    mkdir MYRUNS
     cp -R -p EXAMPLES-depot/H2-EXAMPLE-PLAIN-SCRIPTS EXAMPLES
```

```
cd EXAMPLES/H2-EXAMPLE-PLAIN-SCRIPTS
    follow the README;
    ./Relax.Bat
    ./Fourier.Bat
     ./Flux.Bat 500
    ./gnu.xsec
    for total photoionization; see the README for more.
If you want to update the code then perform step 5, like steps 1-4, in the
V1 working directory,
    git pull https://github.com/LBNL-AMO-MCTDHF/V1 master
     ... then ./Makeme should be sufficient, in the BIN directories,
        but ./Makeme clean; ./Makeme would certainly be fail safe
        after updating the code with git pull.
                             MODULES TO LOAD
_____
----- ON NERSC & LAWRENCIUM ------
 COMPDIRS/BIN.ecs.hermnorm.law,
  COMPDIRS/BIN.ecs.hermnorm.edison, etc.
LAWRENCIUM:
module load mkl
CARVER:
module unload pgi openmpi
module load intel openmpi-intel mkl
module unload torque
module load torque
HOPPER:
module swap PrgEnv-pgi PrgEnv-intel
#yes last
module unload cray-libsci
EDISON:
module unload cray-libsci
module load mkl
```

#### 0.2 Running the code

The code looks for an Input.Inp input file. If another input file is desired, use e.g. > chmctdh\_atom Inp=Input.Inp.myinput

The code will perform a calculation with default parameters, if no input is given.

#### 0.3 File system etiquette

See also section 5.3.1 and note script findwork at top level in the distribution.

If they do not exist in the working directory, the code makes the subdirectories

WALKS Flux Dat Bin timing

For more information on WALKS see section 3, Output. For more information on Flux, see 5.3.1. Bin is generally used to save the wave function, and generally Dat contains numerical text output from actions, section 2.5. Timing output, what is discussed in section ??, is sent to directory timing if &parinp namelist variable notiming is set less than the value 2, its default, and is described in section ??.

Some large files may end up in these directories. Beware of this. Again, see also section 5.3.1 and note script findwork at top level in the distribution. On a supercomputer e.g. NERSC machines, you MUST be reading and writing these files to the \$SCRATCH directory. One could run on the axl condo cluster home file system /clusterfs/axl but make symbolic links to the scratch directory. E.g.

```
> mkdir $SCRATCH/blah5
> ln -s $SCRATCH/blah5 ./WALKS
```

#### 0.4 Run Control

A running calculation can be stopped by creating a file named stop in its working directory.

## 0.5 Optimizing your runs

To successfully use the LBNL-AMO-MCTDHF package you must pay attention to timings. The mean field treatment has several components and it is necessary to intelligently select the basic parameters of the propagation, which are among the &parinp namelist variables, to achieve best performance.

A basic part of the mean field implementation is the separation of the orbital and A-vector propagation. The orbitals and the A-vector are propagated separately over a small time step, the mean field time step, the &parinp namelist variable par\_timestep. Thus, a basic question is, which is taking longer, orbital propagation or A-vector?

We recommend performing rough draft calculations with grids as low-resolution as possible, such that the orbital calculation runs fast. We recommend not attempting to "converge" with respect to number of orbitals, namelist variable nspf. For first row diatomics, the ten-orbital calculation is the obvious

starting point. However, if more orbitals are desired, the A-vector propagation may begin to take the majority of the computation time.

For the polyatomic version <code>chmctdhf\_sinc</code>, you will find best performance using one process per node (as opposed to one process per core, for <code>chmctdhf\_atom</code> and <code>chmctdhf\_diatom</code>, which are generally run as sequential programs, not parallel). There are both vectorized (xSSE4.2, with intel FFT) and OpenMP (using FFTPACK) compilation directories provided for the lawrencium machine in <code>COMPDIRS</code>. The sequential (no shared memory, only MPI parallelization) version is directory <code>BIN.ecs.hermnorm.law.seq</code>. On this machine <code>\$OMP\_NUM\_THREADS</code> controls the number of threads per process and for the vectorized and OpenMP versions, all threads are taken if it is not set.

To ask the question, "what is the breakdown in timings for my run?" please set &parinp namelist variable notiming=0. The code will answer, proceeding to output large amounts of information into separate text files in the timing directory. See sections 0.3 and 5.3.1 on file system etiquette.

• There is Main.time.dat but it should show that the time is all spent in propagation. Below you see there were about 6000 time units spent at the beginning, counted under Non MPI. MPI +Non MPI is the total time and if the parallel propagation is working ok then MPI (communication) should be smaller than Non MPI (computation). If you don't include the startup time which is only counted in the MPI and Non MPI columns, roughly speaking, MPI+ Non MPI should add to the sum of the others. TO-DO: ADD BIORTHOGONALIZATION COLUMN.

	Spfs	Prop	Act	Final	MPI	Non MPI
0.000	0	508	0	2	16	6381
0.050	0	857	0	5	23	6726
2.150	0	10362	0	121	232	16139
2.200	0	10539	0	124	242	16310

• To see the breakdown of the Prop column, examine cmf\_prop.time.dat:

	Time	matel	denmat	reduced	spfprop	aprop	advance	constrain	#DERIVS
T=	0.100	40	2	0	146	640	3	0	81
T=	0.150	61	4	0	229	865	5	0	80
T=	0.200	81	5	0	282	1090	7	0	48
T=	0.250	103	6	0	360	1315	9	0	58
T=	0.300	124	8	0	424	1539	10	0	58
T=	0.350	145	9	0	521	1763	12	0	104
T=	0.400	166	10	0	592	1988	14	0	63
T=	0.450	186	12	0	664	2213	16	0	63

If A-vector propagation is taking quite a bit longer than orbital (spf) propagation, as above, run in parallel and try both parorbsplit=0 and 1 if you want to be systematic and agnostic about it. If matrix elements, density matrices, or any of the in-between steps are taking a long time, CERTAINLY consider using a large par\_timestep and littlesteps> 1. Most of the time is usually taken in orbital propagation, in which case par\_timestep shouldn't affect overall speed that much, unless very small.

#### #DERIVS

is the number of evaluations of the mean field operator required to take the time step (both predictor and corrector \*\*CHECK\*\*) and should never get too much bigger than  $400 \times 2 = 800$  or so. To get time per mean field evaluation, divide the other columns by the number of DERIVS.

• The timing file actreduced.time.dat is very important and tells about column "spfprop" in cmf.prop.time.dat above. However, note that this column spfprop is not only actreduced, which

is the name of a subroutine; there is also the operation of the projector, which is counted separately, and so to see the breakdown of column spfprop in cmf.prop.time.dat, go to jacoperate.time.dat, note that the first column is quite a bit smaller than the second of the two columns, and then look at actreduced.time.dat which goes like this:

		rmult	ke	pot	pulse	nuc	twoe	invdenma	project	constrai	MPI
T=	0.4500	169	881	88	106	0	318	18	4	0	55
T=	0.1500	333	1688	177	212	0	632	35	8	0	110
T=	0.3500	496	2496	266	318	0	947	52	12	0	165
T=	0.6500	659	3303	355	424	0	1261	69	16	0	219

Note that there are two steps to the mean field propagation of the orbitals, the predictor step and the corrector step, so the times, the first numerical column, may not be in order. Orbital parallelization with parorbsplit set nonzero should affect the columns ke through twoe because those refer to steps that are distributed over processors by this option. Generally speaking, with nprocs=nspf, if the orbital parallelization is working well you should observe that the steps rmult, perhaps invdenmat, and MPI take a significant proportion of the time.

• Examine expo.dat which tells you about the orbital propagation. In the example below, notice the numbers 1 and 48 at the bottom. That's one Krylov iteration, no restarts, with 48 matrix-vector multiplications required. This is the output near the start of the run. It has increased the krylov dimension up to 46 and at that point it does not need to restart, whereas at the first step it required 2 steps, one restart, with 66 matrix vector multiplications total. There have been steps rejected. This will happen at the beginning of the run when it is adjusting the internal step size, but should NOT happen afterwards that much. If it is, something is wrong. It will increase the Krylov order (thisexpodim as reported below) from where it starts (namelist input expodim) up to the maximum krylov order (namelist input maxexpodim), and then if it is restarting then the number of steps will be greater than one, but it should not be rejecting steps and changing its stepsize. In a nutshell: increase maxexpodim up to 300, or something like that, and then if it is restarting, (steps > 1) so be it, but it should not be rejecting steps very often, after the start of the run. Consider increasing littlesteps if you are trying to take big steps (large par\_timestep).

```
31
step 5.0000000000000E-002
     ...REJECTED 0.5000E-01 0.3849E-06 0.1000E-05
     ...REJECTED 0.4000E-01 0.1023E-06 0.1000E-05
  End expo. Orthog error, steps, stepsize, iterations 6.050711511541938E-015
         2 3.2000000000001E-002
Go Orbital Expoprop. Tinit= 0.100000000000000
                                                thisexpodim=
                                                                    38
      5.00000000000000E-002
      ...REJECTED 0.5000E-01 0.2653E-06 0.1000E-05
  End expo. Orthog error, steps, stepsize, iterations 6.213364751595364E-015
         2 4.0000000000001E-002
Go Orbital Expoprop. Tinit= 0.150000000000000
                                                thisexpodim=
                                                                    46
 step 5.0000000000000E-002
  End expo. Orthog error, steps, stepsize, iterations 1.136962511459106E-014
         1 4.999999999999E-002
                                        48
```

- Check file avecexpo.dat, for the A-vector propagation, similar to expo.dat above.
- If running in parallel, check to be sure you get the same results (certainly to six digits for everything) for different numbers of processors.
- improvednatflag to enforce natural orbitals for relaxation has been extensively debugged but it involves a difficult transformation of the wave function. For very large A-vectors, especially, be sure it is working ok; do the run without the flag as well. It is conceivable that the biorthogonalization routine could require tweaking for difficult cases, using the variables biodim and biotol. The only two reasons, in general, for which one needs to use this option improvednatflag in general is if,

first, one is going to perform excitations or annihilations subsequently – or alternatively, second, if using density matrix constraint (constraintflag=1) in the subsequent propagation step.

- Try larger time steps as described above. If you have a high frequency or strong laser and you are trying to take large time steps (e.g. for big A-vector), try the new variable littlesteps > 1 which may improve accuracy and stability in addition to speed, as described above.
- Use the smallest primitive basis possible until you are doing production runs. Go as small as you can without much static appearing in your result. First make sure things are qualitatively right. Absolute energies are not relevant. Gauge dependence is. Don't sweat getting transitions energies right, unless something is qualitatively wrong. Often transition/ionization energies can be off by an eV or more.

#### 0.6 Nomenclature

Here are some quick statements about the terms used in this manual.

We sometime use the terms configuration and Slater determinant interchangeably. Sometimes when we write configuration, we mean either a Slater determinant or a spin adapted sum of Slater determinants. But usually, we just mean Slater determinant.

The wave function is made of a time dependent linear combination of Slater determinants. The coefficients in the linear combination are time dependent. The Slater determinants are also time dependent, because they are made of orbitals that are time dependent. Sometimes (with allspinproject=1, default) the spin symmetry of the wave function is enforced. Then, the wave function is first represented as a linear combination of spin adapted linear combinations (SALCs is an acronym we have seen) of Slater determinants. Those linear combinations, the SALCs, are computed when the code starts.

So, not worrying about whether or not we are discussing Slater determinants (allspinproject=0) or spin adapted linear combinations thereof (allspinproject=1), we will generally use the term "configuration."

Again, we borrow from the MCTDH implementation of the Heidelberg group for not only the notation used in the equations, but also the terms by which they are referred to and often even the variable names in the code.

Orbitals are the same thing as single particle functions, SPFs. The configuration coefficients are the "A-vector,"  $\vec{A}$ . We have one- and two-electron reduced density matrix operators  $\rho$  and  $\Gamma$ , the reduced potential  $\mathbf{w}$ , the reduced mean field operator  $\mathbf{W}_{\alpha\beta} = \sum_{\gamma\delta} \Gamma^{\gamma\delta}_{\alpha\beta} \mathbf{w}_{\gamma\delta}$ , and the one-electron Hamiltonian  $h_0$ . The orbitals are  $\vec{\phi}$ ; the coefficients of  $\phi_{\alpha}$  are  $\vec{c}_{\alpha}$ .

For restricted configuration spaces, in which the orbitals are constrained to rotate into one another, we compute the  $g_{\alpha\beta} = \langle \phi_{\alpha} | i \frac{\partial}{\partial t} | \phi_{\beta} \rangle$  matrix. The matrix elements of g among configurations are called  $\tau$ .

$$i\frac{\partial}{\partial t}\vec{A} = (H - \tau)\vec{A}$$
$$i\frac{\partial}{\partial t}\vec{\phi} = \left[ (1 - P) \left( \rho^{-1}\mathbf{W} + h_0 \right) + g \right] \vec{\phi}$$

## Chapter 1

## Installation

System administrators may use this section for very quick startup especially on supported machines. Users may proceed to the next section.

The directory structure of the code makes extensive use of symbolic links and if you copy the directories you should ensure that you maintain the symbolic link structure. On mac, use cp -R to copy links as is. Otherwise, they are copied as new files, and you have duplicates. This is not a big deal, but it can be a waste of space and if you update the file then you have inconsistencies. You want to keep one copy of the files only. Look at the directories, see the symbolic links, and then proceed to use these instructions:

The code is provided with the following directory structure. The result of the 1s command is schematic and embellished to give you the idea.

```
prompt> tar -xvf MCTDHF.010115.VERSION1.0.tar.gz
prompt> cd MCTDHF.010115.VERSION1.0
prompt> 1s
  LBNL-AMO-MCTDHF.pdf
  EXAMPLES
     H2PH0T0
   JOB_SETUP_SCRIPTS
     corotating_counterrotating_streaking.py
        mpilaunchjob.slurm.template
        atomfile.txt
        Relax.Bat.template
  MCTDH.SRC
     DFFTPACK
        zfftf1.f
     SINCDVR
        sincDVR.f90
     H2PROJECT
        H2_params.f90
  COMPDIRS
     MCTDH.SRC -> ../MCTDH.SRC
     BIN.ecs.hermnorm.law.openmp.nofft
        Definitions.INC
        Name.Txt
        Makefile.header -> ../MCTDH.SRC/Makefile.header.lawrencium.openmp.nointelfft
        Definitions.ALL -> ../MCTDH.SRC/Definitions.ALL
        mctdhf.F90 -> ../MCTDH.SRC/mctdhf.f90
            Makefile.header -> ../Makefile.header
            Makefile -> ../../MCTDH.SRC/DFFTPACK/Makefile
```

```
zfftf1.f -> ../../MCTDH.SRC/DFFTPACK/zfftf1.f
SINCDVR
    Makefile.header -> ../Makefile.header
    Makefile -> ../../MCTDH.SRC/SINCDVR/Makefile
    sincDVR.f90 -> ../../MCTDH.SRC/SINCDVR/sincDVR.f90
debug.BIN.ecs.hermnorm.mac
BIN.mac
BIN.ecs.hermnorm.edison
```

etc.

There are three directories at top level, MCTDH.SRC, EXAMPLES, and COMPDIRS. The MCTDH.SRC directory at top level contains the actual hard copy of the fortran code. There are many directories ready-to-go for installation of the code on various machines, provided in COMPDIRS. There are optimized versions, and versions with debugging flags. Please compile the debugging version too and use it if the code seems to be producing an invalid or unstable result, or actually crashes. The supported machines, the machines with ready-to-go directories in COMPDIRS, are

- Lawrencium (extension .law, e.g. BIN.ecs.hermnorm.law.openmp.nofft). Intel sandybridge, intel fortran compiler with mkl.
- Edison on NERSC (extension .edison, e.g. debug.BIN.ecs.hermnorm.edison)
- Mac (extension .mac, e.g. debug.BIN.mac)

You may delete the directories in COMPDIRS that you don't want, but they don't take any space. All you need to do is go to the BIN directory you want and run ./Makeme. The code should then be compiled as, e.g.

Then you need to put the program in your \$PATH somehow so that you can use it in any working directory. We recommend

```
prompt> echo $PATH
   /usr/bin:/usr/sbin:/opt/local/bin:/home/me/bin
prompt> cd ~/bin
prompt> ln -s ~/myprogs/LBNL-AMO-MCTDHF/V1.0.INTELFFT/COMPDIRS/BIN.ecs.hermnorm/chmctdhf_diatom ./chmctdhf_diatom
prompt> ln -s ~/myprogs/LBNL-AMO-MCTDHF/V1.0.INTELFFT/COMPDIRS/debug.BIN/mctdhf_sinc ./mctdhf_sinc.debug
```

There are four versions of the code with different datatypes, but chmctdhf is all that is generally needed. Compilation directories are provided for chmctdhf and mctdhf only.

- mctdhf: REAL VALUED VERSION FOR RELAXATION. (e.g. BIN.mac)
- pmctdhf: COMPLEX VALUED, PROPAGATION WITHOUT ECS (e.g. BIN.complex.mac)
- chmctdhf: COMPLEX VALUED, WITH ECS: USUAL VERSION (e.g. BIN.ecs.hermnorm.mac)
- cmctdhf: COMPLEX VALUED, ECS, C-NORM: FOR RESONANCE RELAXATION (e.g. BIN.ecs.cnorm.mac)

If you are using an architecture different from all of these machines, then you should make a new BIN directory and a new Makefile.header.mymachine in MCTDH.SRC, linking the new header file as Makefile.header in the new BIN directory. There are parallel versions of Makefile.header for mac already done. Your fortran compiler must support the C preprocessor. You may have to figure out flags to invoke it. If you want to run the calculation in parallel (there is now some orbital parallelization!) then in Makefile.header you must include the line "MPIFLAG = -D MPIFLAG" and include paths or flags in variables MYINCLUDE and LAPACK if needed. Intel FFT libraries may be used by setting FFTFLAG = -D INTELFFT. Portland group compilers require PGFFLAG = -D PGFFLAG.

## Chapter 2

# Input

The code is run from the command line and takes command line arguments and also parameters from an input file, by default Input.Inp. Command line arguments supersede input from file. All variables that can be set are listed in Sec. 2.6. To use an input filename different from Input.Inp, specify it as follows:

C:> chmctdhf\_atom Inp=Input.Inp.myinput

If Input.Inp or the specified input filename does not exist it will do a default calculation. One can specify multiple command line arguments, e.g.

C: > chmctdhf\_atom Inp=Input.Inp.awesome T=1000.0 Nspf=4 Act=1

#### 2.1 Namelist input

Fortran includes a feature that makes it easy to input variable values from file, called namelist input. It is not case sensitive. Variable names in the namelists in the input file are the same as their names in the code. There are four namelists that can be used as input in the input file (default Input.Inp). The complete list of variables that can be set are listed in Sec. 2.6, which is a printout of parameters.f90 with comments. You can also look at the code: for instance, the &parinp namelist variables can be seen in getparams.f90, subroutine getparams(); that's where the &parinp namelist is defined.

### \*\* In your input file: \*\*

- You must have namelist &parinp.
- If variable tdflag in namelist &parinp is set to 1 (including a time-dependent pulse), or if doing a flux analysis calculation (action 16 or 17) (except if noftflag is set nonzero in parinp), you must also have namelist &pulse.
- You must have &h2parinp for diatoms (chmctdhf\_diatom), &heparinp for atoms (chmctdhf\_atom), or &sincparinp for polyatomics (chmctdhf\_sinc).

### 2.2 Specifying the wave function

The orbitals and A-vector are specified independently. One may load either of them, or obtain them by diagonalization. For orbitals, the core Hamiltonian is used for diagonalization. One needs to specify the symmetry constraints for each. Restricting the symmetry of the A-vector requires restricting the symmetry of the orbitals, but not vice-versa.

#### 2.2.1 Symmetry constraints

By default the  $S_z$  spin angular momentum projection quantum number is restricted. It is specified by restrictms=X. This is  $2 \times$  the  $S_z$  quantum number.

By default the wave function is restricted to be an eigenfunction of spin,  $\hat{S}^2$ , via spinwalkflag=1 and allspinproject=1. The high spin case is calculated  $(S = M_s)$ . Thus one specifies the multiplicity by setting restrictms = multiplicity-1, i.e. for a doublet, restrictms=1.

One generally wants to restrict spatial symmetry as well. To restrict the  $l_z$  quantum number (angular momentum projection) of the orbitals set spfrestrictflag=1. Then specify the  $m_z$  values of the orbitals by spfmvals=0,1,-1,... To restrict the parity of the orbitals set spfugrestrict=1. Then specify the parity by spfugvals=1,-1,0,0,... where 0 means no parity restriction.

Without spfrestrictflag=1, spfmvals is only used if you are starting from core orbitals, for your initial relaxation run for instance, in which case it is just used to select the initial orbitals. Similarly for spfugrestrict.

To restrict the overall angular momentum projection  $L_z$  of the wave function, spfugrestrict must be set nonzero; then set mrestrictflag=1 and mrestrictval to  $L_z$ . Similarly with ugrestrictflag and ugrestrictval.

To restrict  $L_z$  to a range of values, do not set mrestrictflag; instead set mrestrictmin and mrestrictmax. This is useful for relaxation or propagation of multiple wave functions. See warnings in the section on relaxation.

If one is reading in orbitals from file and spfrestrictflag or spfugrestrict is set, then the angular momentum projections or parities of the orbitals must match those in the present calculation.

#### 2.2.2 Restricted configuration spaces

Restricted configuration spaces, non-full-configuration interaction, are a new capability and the numerical implementation is more difficult than the full CI method. There are more things to do, and the integrator ends up taking more steps to calculate the more rapidly changing orbitals. There are profound issues relating to symmetry broken solutions of a relaxation calculation, to prepare the initial state, and therefore we advise using the real-valued versions BIN/mctdhf\_XXX to prepare the initial state, or at least checking that the same energy is obtained with both complex and real versions.

For these reasons using restricted configuration spaces be avoided until you have pushed the limit of a one node calculation. Doing the code in parallel over many nodes is not something we want to do if we want to make the best use of computer time. For instance, with neon, with greater than 14 orbitals, one is going to need a restricted configuration space, despite the use of sparseconfigflag=0, sparseopt=0, and small values for Krylov space parameters (maxaorder, aorder, even maxexpodim if the orbitals take up some space that can be freed).

You can also try frozen orbitals, numfrozen, which reduces the number of electrons in your full CI calculation, to avoid having to use a restricted configuration space. Frozen orbitals work, but they also require more things to do, slowing the calculation down, and make the problem more numerically difficult. But, if you want, do try frozen orbitals if your physics does not perturb the core orbitals! This will require consultation as the manual is not complete. See section 2.2.4.

There are two methods for restricted configuration spaces (see the paper). Density matrix (constraintflag=1) and dirac-frenkel (constraintflag=2). A restricted configuration space can be used with constraintflag=0, such that the  $g/\tau$  matrix remains set to zero, but one is making uncontrolled error in the wave function.

With constraintflag not equal to zero, dfrestrictflag must be set nonzero. Unless you are running action 22, it should be set dfrestrictflag=1.

Then, **ALL THE FOLLOWING VARIABLES** should be incremented by the chosen value of dfrestrictflag. That is because, in the code, they will be decremented by that amount, to obtain the included list of configurations. "Incremented by 1" means, change the value by 1 in the direction that enlarges the restricted configuration space.

minocc maxocc numexcite vexcite

Variables (namelist &parinp) specifying a restricted configuration list are as follows. The variables numshells

shelltop

come first. They divide the nspf orbitals into shells. By default there is one shell, numshells= 1, and shelltop(1)=nspf. shelltop denotes the last spatial orbital in each shell. You must enter shelltop for shells 1 through numshells-1. The value of shelltop for the last shell is automatically set to nspf.

One way of doing a restricted configuration list, is to use one or the other or both of the two variables numexcite

vexcite

and leave it at that. Variable numexcite may be set for shells 1 through numshells-1; numexcite(i) corresponds to the maximum number of holes in shells 1 through i (so the list, if numshells> 2, should be never decreasing). vexcite is the opposite of numexcite and is only input for the last shell; it corresponds to the maximum number of electrons in the last shell and so can be used to easily restrict the configuration list to all singles and doubles (CISD), etc.

The other available options are minocc and maxocc which can be input for shells 1 through numshells. These are minimum and maximum occupation numbers.

To reiterate, vexcite, numexcite, minocc, and maxocc must all be incremented by the value of dfrestrictflag (1, unless using action 22, or unless debugging) from what you want them to be for the restricted configuration space you want. The configuration lists are built using the incremented values, but the wave function is constrained to occupy only the restricted configuration space you want. If you are just playing around and constraintflag and dfrestrictflag are both zero then just set these variables how you want them; you can do that, but you are making uncontrolled error.

TO PREPARE THE INITIAL STATE: To prepare the initial state for a propagation with the density matrix constraint (constraintflag=1), perform a relaxation calculation with improvednatflag=1, constraintflag=0, dfrestrictflag=0. You could also set dfrestrictflag to the same value (probably 1) that you use for the propagation, in which case your values for vexcite, minocc, etc. will be the same. But constraintflag=0, improvednatflag=1 for density matrix initial state calculation. For Dirac-Frenkel (constraintflag=2), perform a relaxation calculation with the restricted configuration space machinery just as done for propagation. You may even find you have to do regular imaginary time relaxation calculation (improvedrelaxflag=0, threshflag=1). For relaxation or improved relaxation you will need small time steps. Set constraintflag=2, dfrestrictflag=1 (or whatever nonzero), just like the propagation calculation.

Always, if you are indeed taking the bold leap into restricted configuration space calculations, you should definitely do it BOTH WAYS, density matrix and dirac-frenkel, constraintflag=1 and 2. Hope-

fully they will both work and if they do, it is a powerful convergence check. Remember that you can check the overlap between two time-dependent wave functions using Actions 15 and 23.

#### 2.2.3 Specifying the initial orbitals

By default the orbitals are obtained by diagonalizing the core hamiltonian. These would be used for a relaxation calculation. Thus, for a relaxation calculation, no extra input is required.

One loads orbitals from file with loadsppflag=1.

```
&parinp
  loadspfflag=1
  spffile="Bin/spfs.bin.mine"
```

By default, the orbitals are loaded from ./Bin/spfs.bin, the same file that is written at the end of a run. In practice you would save this file elsewhere and specify its location as above.

If you are loading orbitals with <code>loadsppflag=1</code> and there are fewer orbitals in <code>spfs.bin</code> than in the calculation, then the additional orbitals are taken from core orbital eigenfunctions. This is useful if you want to converge an excited state with a small number of orbitals first, then add orbitals to get a better wave function.

For a relaxation run, obtaining the initial guesses for the orbitals from core diagonalization, even if specifying spfmvals and spfugvals as appropriate, one may find that the initial guess orbitals are incorrect for the desired calculation. For instance, the ordering of 3s and  $3d_{z^2}$  will depend on small numerical errors. One may skip over the orbitals as follows: say that you want the 1st, 2nd, and 5th orbitals for m=0 and the 1st and 3rd for  $m=\pm 1$ . One would specify

```
&h2parinp
  num_skip_orbs=3
  orb_skip=3,4,2
  orb_skip_mvalue=0,0,1
```

Thus you list the orbitals to skip and the corresponding m-values.

To read in two or more small sets of orbitals to combine into one set (useful for interacting fragment type description) use numspffiles and spffile(1), etc. For instance you might have a run with three orbitals and one with four and you could combine these for a seven orbital calculation. You can skip over orbitals that are read in this step, by specifying numskiporbs, in namelist &parinp.

#### 2.2.4 Frozen orbitals

If your calculation is getting out of hand (can't fit on one node, in memory), and it only involves valence electron dynamics, then you have two options – restricted configuration spaces, or frozen orbitals. If it involves core electron dynamics, you only have the restricted configuration space option. Otherwise, if you are really at this point, perhaps, try both.

Frozen orbitals work like this. Variable numfrozen in namelist &parinp is number of frozen orbitals (spatial orbitals, containing two electrons). So you might start with a small full CI calculation. Say Neon, 6 orbitals. Don't do hartree-fock! We do not have an actual fock Hamiltonian. We are missing the "scalar terms" in the reduced operator  $\hat{\mathbf{W}}$ . They do not matter when (1-P) is put in front of them. Therefore we do not have a good 1s energy eigenfunction. All we have is natural orbitals – or, now, the Dirac-Frenkel constraint – to define orbitals.

By "define orbitals" we mean, in better terms, "resolve orbitals". Given the space of orbitals – six, say, for neon, with an extra 3s orbital beyond the Hartree-Fock – resolve orbitals means specify a specific set of six orthonormal orbitals.

So, the easiest thing is to start with non-Hartree-Fock, improvednatflag=1 calculation, with the restricted configuration space (no need for dfrestrictflag nonzero).

Take the orbitals on disk, and read them in for a calculation with frozen orbitals. Start with neon, six orbitals, then read those orbitals in, with nspf=5, numfrozen=1. nspf is the number of unfrozen orbitals. That will do a calculation with one frozen orbital, and five unfrozen orbitals, starting with the six orbitals from your first calculation; the frozen orbital is now the 1s natural orbital from the first calculation. You can go on and add more orbitals.

Specify spfmvals and spfugvals only for the unfrozen orbitals.

So, you could have done nspf=8, numfrozen=1, and it would have loaded the six orbitals, then added three more (the p orbitals) gotten from diagonalization of the core Hamiltonian.

Going from six orbitals to nspf=13, numfrozen=1, one would set the thirteen values of spfmvals and spfugvals appropriately, then, we think this is how it goes, one would need to specify num\_skip\_orbs=1 and orb\_skip\_mvalue=0 in &h2parinp or &heparinp, to get the usual set of fourteen orbitals.

#### 2.2.5 Specifying the initial A-vector

Similar to the orbitals, one may specify

```
&parinp
  loadavectorflag=1
  avectorfile="Bin/avector.bin.mine"
```

With mcscfnum > 1 the code propagates multiple A-vectors, but still uses only one set of orbitals. To read in multiple A-vectors for such a run, use numavectorfiles and avectorfile(1)=, etc.

One may also excite or annihilate orbitals and the following subsections describe that.

When the excitation or annihilation is performed, it is done so in terms of the spin orbitals and the spin value is not conserved. What is obtained from the excitation or annihilation is then projected upon the spin value specified in restrictval, if spinwalkflag=1.

One should take care that the orbitals one is annihilating or exciting are meaningful. If one does a regular improved relaxation run, the orbitals will be arbitrarily mixed. If you want to have meaningful molecular orbitals, use <code>improvednatflag=1</code> in namelist <code>parinp</code> in the relaxation run; the orbitals will then be natural orbitals. Fock eigenfunctions are not implemented.

#### 2.2.6 Annihilation

One may read in an (N+1)-electron wave function and then annihilate an electron. Use avectorhole= in namelist &parinp.

```
&avectorhole=1
```

This will annihilate the first spin orbital. As it happens the even numbered spin orbitals are spin up and the odd numbered are spin down. So if one is reading in a triplet wave function (restrictms=2) into a doublet calculation (restrictms=2) and annihilating an electron, because the high spin case is calculated one annihilates a spin up electron, odd numbered, as above.

One may use a linear combination of hole wave functions. This would be useful for annihilating the left or right core hole. To do this specify

This makes the (+) combination of orbitals (which each come out with an arbitrary phase). To make the (-) combination use &avectorhole=1,-3.

#### 2.2.7 Excitation

One may read in an A-vector and then perform an excitation or annihilation.

To perform an excitation, use avectorexcitefrom and avectorexciteto in namelist &parinp. These variables should be assigned to the spin orbital indices. With the default value of orderflag, this means that to perform an excitation from the first spatial orbital, spin down, to the third spatial orbital, spin up, you'd do avectorexcitefrom=2, avectorexciteto=5. As for annihilation, the excitation does not preserve the S(S+1) quantum number and the wave function obtained is subesequently projected on the given spin space if spinwalkflag=1.

#### 2.3 Primitive basis and Hamiltonian

The primitive basis and hamiltonian are specified in the Fortran namelists h2parinp, heparinp, or &sincparinp in the input file. The namelist read depends upon the version of the code that is run (chmctdhf\_diatom, chmctdhf\_atom, or chmctdhf\_sinc).

```
nspf=5
spfmvals=0,0,1,1,1
&h2parinp
NucCharge1=1.d0
NucCharge2=5.d0
LBIG=13, MBIG=1
pro_hmass=1836.152701d0
pro_dmass=20213.07d0
bornopflag=0
xinumpoints=14
xinumelements= 8
xicelement=7
xiecstheta= 0.157d0
xielementsizes = 2.0d0, 10.0d0, 10.0d0, 10.0d0, 10.0d0, 10.0d0, 10.0d0, 10.d0, 10.d0,
rnumelements=1
rnumpoints=32
relementsize = 2.5d0
rstart=1.836d0
```

The most important options are:

- bornopflag: Performs a Born-Oppenheimer calculation, one is default. Zero: full nonadiabatic.
- xicelement: First element that is complex-scaled.
- xinumelements: Total number of elements in xi.
- rnumelements: Number of elements in R
- LBIG: Number of points in  $\eta$  minus one.
- xinumpoints, rnumpoints: number of points per element, including bridge functions.

The masses affect a fixed-nuclei (bornopflag=1, default) calculation because different masses will shift the origin in the prolate coordinate system. Different masses may give superior or inferior results for a given heteronuclear. If in doubt, don't include masses which will set masses equal.

#### 2.4 Propagation parameters

See the advice at the top of the document about making runs go faster. The most important factor in speeding up the calculation is to use the maximum grid spacing possible. We advise using length gauge, if stable, and even grid spacing (no small first element) which we believe will give you the fastest possible converged calculation.

The following options may be useful for fine tuning the propagation.

```
&parinp
aorder
                   !! Krylov dimension for A-vector propagation
                   !! Krylov dimension for A-vector propagation
maxaorder
aerror
                   !! Error tolerance for A-vector propagation OR improvedquadflag=1
expodim
                   !! Corresponding Krylov dimension
                   !! Corresponding Krylov dimension
maxexpodim
expotol
                   !! Error tolerance
littlesteps
                   !! Divide par_timestep into many steps for calculation with pulse
                   !! density matrix regularization parameter (rarely matters)
denreg
lioreg
                   !! if constraint, ESPECIALLY if DF constraint (constraintflag=2)
                   !! regularization parameter for lots of matrix inversions/linear solves
invtol
```

#### 2.4.1 Analyzing the propagation

See the section at the beginning of the manual about examining and improving the performance of your runs.

#### 2.4.2 Using sparse configuration routines

Set sparseconfigflag=0 to use nonsparse configuration routines; otherwise sparse is default. If sparseconfigflag=0 is specified, there is a maximum number of configurations that is allowed and if it is exceeded the program quits. To override this you can specify nosparseforce=1. In practice there are few reasons to do this. If you have something like 100-300 configurations, you should check whether sparse or nonsparse is faster.

With sparseconfigflag=0, the default behavior is that the sparse CI Hamiltonian is constructed in sparse format. Sometimes – NO with 12 orbitals on Lawrencium, for instance – the calculation can't fit on one node with this default behavior. Setting sparseopt=0, as opposed to the default, 1, performs a direct CI instead. No sparse hamiltonian is constructed; memory use is less, but it is slower.

### 2.5 Action input

Things that one can optionally do to or with the wave function in-between time steps are called actions. Everything you want to do (fourier transform dipole moment, output orbitals, whatever) is an action, and specified by numactions=XX, action=YY,ZZ,....

### 2.6 Complete list of namelist variables

Below we provide verbatim copies of the files parameters.f90, H2PROJECT/H2\_params.f90, HEPROJECT/He\_params.f90, and SINCDVR/sinc\_params.f90. The first is for all calculations, namelist &parinp; the others are for namelists &h2parinp, &heparinp, and &sincparinp, which are required for chmctdhf\_diatom, chmctdhf\_atom, and chmctdhf\_sinc, respectively. You will find description of lots of minor options in the code here.

```
!! Parameters for MCTDHF calculation; parinp NAMELIST input from Input.Inp (default)
11
!! Type, variable, default value !! Command-line !! Description
                               !! option
!!
!!
     SPARSE - if sparseconfigflag .ne. 0
module lan_parameters
integer :: lanprintflag=0
integer :: lanczosorder=200
                               !!
                                               !! lanczos order used in A-vector eigen.
integer :: lancheckstep=20
                               1.1
                                               !! lanczos eigen routine checks for convergence every this # steps
real*8 :: lanthresh=1.d-9
                                               !! convergence criterion.
end module lan_parameters
module sparse_parameters
integer :: sparseprime=1
                                               !! For reordering config list (experimental)
integer :: sparsesummaflag=0
                                               !! 0=gather 1=summa(bad) 2=sendrecv(ok)
integer :: sparsedfflag=1
                                               !! If zero disable separate restricted config walk list
integer :: sparseconfigflag=0
                               !! Sparse
                                               !! Sparse configuration routines on or off (for large # configs)
integer :: sparseopt =1
                                               !! O= direct CI 1= sparse matrix algebra (faster, more memory)
integer :: nonsparsepropmode=1   !! 0 = ZGCHBV expokit; 1 = mine expmat
logical :: use_dfwalktype=.false.
                                               !! internal
end module sparse_parameters
module ham_parameters
     HAMILTONIAN PARAMETERS
integer :: nonuc_checkflag=1
                                               !! Turn off deriv operators in nuclear dofs.
                               !! Pulse
integer :: tdflag=0
                                               !! Use pulse?
integer :: velflag=0
                               !!
                                               !! Length (V(t)) or velocity (A(t))
!! Constraintflag=1: Density matrix constraint: assume nothing, keep constant off block diag
!! 2: Dirac-Frenkel (McLachlan/Lagrangian) variational principle.
integer :: constraintflag=0
                               !! Constraint= !! As described below (see CONSTRAINT)
integer :: denmatfciflag=0
                                               !! If .ne. O then does denmat constrant as programmed
                                               !! before Miyagi's help
DATATYPE :: energyshift=0d0
                                               !! complex shift for making energy real for imperfect CAP/ECS
integer :: drivingflag=0
                                               !! Solve for the change in the wave function not wave function
real*8 :: drivingproportion=0.9999999999900
                                                   -- "psi-prime" treatment.
DATATYPE :: timefac=&
                               !! Prop/
                                               !! d/dt psi = timefac * H * psi
       DATANEGONE
                               !! Relax
                                               !!
real*8 :: mshift=0d0
                                               !! shift configurations based on m-value.. to break
                                               !! degeneracy for state averaged sym restricted
                                               !! (mrestrictmin, mrestrictmax) mcscf; good idea.
integer :: offaxispulseflag=0
                                               !! internal (not namelist)
end module ham_parameters
module basis_parameters
    For restricted configuration lists (not full CI): SEE MANUAL about dfrestrictflag
integer :: numshells=1
                                               !! number of shells. greater than one: possibly not full CI.
!!$ integer :: shelltop(100)=-1 !! Numfrozen=
                                               \verb|!!| shelltop is namelist input in parinp; the internal variable
                                               !!
                                                    is allshelltop. shelltop(1) only may be assigned via
                                               !!
                                                    command line, with Numfrozen.
integer :: numexcite(100)=999
                                !! Numexcite=
                                               !! excitations from core shells (i.e. defined for shells 1
                                                    through numshells-1). Only numexcite(1) may be
                                               !!
                                                    assigned via command line input.
integer :: minocc(100)=-999
                                               !! minimum occupation, each shell
integer :: maxocc(100)=999
                               1.1
                                               !!
                                                    maximum
integer :: vexcite=999
                               !!
                                               !! excitations INTO last shell. Use to restrict to doubles, etc.
integer :: shells(100)=1
integer :: allshelltop(0:100)=0
```

**!!** CONFIGURATIONS

```
integer :: mrestrictflag=0
                                               !! If spfrestrictflag=1, restrict wfn to given total M.
                                1.1
integer :: mrestrictval=0
                                1.1
                                               !! This is the value.
                                               !! If doing state averaged MCSCF, can include a range of m vals;
integer :: mrestrictmax= 99999
                                !!
integer :: mrestrictmin=-99999
                                !!
                                               !! set these variables, with mrestrictflag=0, spfrestrictflag=1
integer :: ugrestrictflag=0
                                !!
                                                !! like mrestrictflag but for parity
                                                !! like mrestrictval but for parity (1=even,-1=odd)
integer :: ugrestrictval=1
                                !!
integer :: restrictflag=1
                                                !! Restrict spin projection of determinants?
" ORBITALS (SINGLE PARTICLE FUNCTIONS, SPFS)
integer :: spfrestrictflag=0
                                                !! Restrict m values of orbitals?
integer :: spfmvals(1000)=0
                                                !! M-values of orbitals
                                !!
                                                !! Restrict parity of orbitals?
integer :: spfugrestrict=0
                                !!
integer :: spfugvals(1000)=0
                                                !! Parity (+/-1; 0=either) of orbitals (ungerade/gerade)
                                !!
end module basis_parameters
module output_parameters
integer :: iprintconfiglist=0
end module output_parameters
module timing_parameters
   Timing
integer :: notiming=2
                                !!NoTiming=0,1,2!! O=write all 1=write some 2= write none
                                                      controls writing of all timing and some info files
                                !! Timing=2,1,0!!
integer :: timingout=499
                                1.1
                                               !! various routines output to file (timing info) every this
                                !!
                                                !! # of calls
character(len=200):: timingdir="timing"
end module timing_parameters
module tol_parameters
real*8 :: lntol=1d-8
real*8 :: invtol=1d-8
end module tol_parameters
module bio_parameters
   Biorthogonalization
integer ::
               maxbiodim=100, &
    biodim=10
                                !! Krylov dim for biorthogonalization
real*8 ::
             biotol=1.d-6
                                !! 1=old way complex zg/hpiv 0=always real
integer :: biocomplex=0
integer :: auto_biortho=1
                                !! do we want to use biorthonormalization or permutation overlaps? 0 perm overlaps, 1 biortho
end module bio_parameters
module spfsize_parameters
integer :: spfsize,spfsmallsize
                                                !!internal
integer :: reducedpotsize = -1
                                                !!internal
integer :: parorbsplit=1
                                                !! Parallelize orbital calculation. Might speed up, might
                                                   slow down; check timing.
end module spfsize_parameters
module constraint_parameters
   CONSTRAINT: With constraintflag. NEED FOR RESTRICTED CONFIG LIST.
real*8 :: lioreg= 1d-9
                                !!
                                                !! Regularization for linear solve constraint
                                                !! for constraintflag=2, dirac frenkel constraint
integer :: conway=0
                                                    O=McLachlan 1=50/50 mix 2=Lagrangian
                                                !!
                                                    3=Lagrangian with epsilon times McLachlan
real*8 :: conprop=1d-1
                                                !! epsilon for conway=3
real*8 :: condamp=1
integer :: liosize
end module constraint_parameters
module denreg_parameters
real*8 :: denreg=1d-10
                                !! Denreg=
                                                !! density matrix regularization parameter.
end module denreg_parameters
module df_parameters
```

#### " CONFIGURATIONS

```
integer :: df_restrictflag=0
                                  !!
                                                 !! apply constraint to configuration list? Must use this
                                                !! option if constraintflag /= 0. 1 is sufficient;
                                                !! dfrestrictflag=2 necessary for action 22.
                                                !! SEE MANUAL FOR PROPER USE OF dfrestrictflag/shell options.
end module df_parameters
module parameters
 use littleparmod; use fileptrmod; use r_parameters; use sparse_parameters; use tol_parameters
 use ham_parameters; use basis_parameters; use timing_parameters; use spfsize_parameters; use df_parameters
  MAIN PARAMETERS
integer :: mcscfnum=1
                                !! MCSCF=
                                                !! Number of A-vectors (state avgd mcscf or prop)
integer :: numelec=2
                                                !! NUMBER OF ELECTRONS
                                11
integer :: orbcompact=1
                                1.1
                                                !! Compact orbitals for expo prop with spfrestrictflag? Probably ok.
integer :: saveflag=1
                                                !! if zero does not save wave function at the end
integer :: save_every=0
                                                !! if nonzero saves wave function every save_every mean field steps
integer :: walkwriteflag=0
                                                !! Turning OFF writing of walks by default
integer :: spf_flag=1
                                                !! IF ZERO, FREEZE SPFS. (for debugging, or TDCI)
                                !!
integer :: avector_flag=1
                                !!
                                                !! IF ZERO, FREEZE AVECTOR. (for debugging)
!! FOR TOTAL ORBITAL PARALLELIZATION with SINC DVR, SET PARORBSPLIT=3
!! and orbparflag=.true. in &sinc_params. parorbsplit=3 not supported for atom or diatom.
integer :: par_consplit=0
character (len=200) :: &
                                                !! MAY BE SET BY COMMAND LINE OPTION ONLY: not namelist
                                !! Inp=filename !! input. (=name of input file where namelist input is)
  inpfile="Input.Inp
  PROPAGATION/RELAXATION
                                                !! MEAN FIELD TIMESTEP
real*8 :: par_timestep=0.1d0
                                !! Step=
integer :: improvedrelaxflag=0
                                !! Relax
                                                !! For improved versus regular relaxtion.
integer :: threshflag=0
                                1.1
                                                !! Set to 1 for regular relaxation
real*8 :: expotol=1d-8
                                !!
                                                !! Orbital krylov convergence parameter
integer :: maxexpodim=100
                                                !! Orbital maximum kry dimension OR DGMRES DIM improvedquad=2,3
                                 !!
real*8 :: expostepfac=1.2d0
                                11
                                                !! Miscellaneous algorithm parameter
     SPARSE - if sparseconfigflag .ne. 0
integer :: maxaorder=100
                                 1.1
                                                     lanczos order for sparse a-vector prop and improvedquad=1,3
  PROPAGATION
integer :: littlesteps=1
                                                !! Sub intervals of mean field time step for avector prop
                                1.1
real*8 :: finaltime=4d4
                                !! T=
                                                !! length of prop. Overridden for pulse and relax.
                                                !! Starting krylov size for orbital propagation (expokit)
integer :: expodim=10
                                1.1
     SPARSE - if sparseconfigflag .ne. 0
integer :: aorder=30
                                 1.1
                                                1.1
real*8 :: aerror=1d-9
                                 !!
                                                !! lanczos error criterion for sparse a-vector CMF propagation
                                                !! within aerror to stop.
   RELAXATION
integer :: improvednatflag=0
                                                !! If improved relax, replace with natorbs every iteration
real*8 :: stopthresh=1d-5
                                !!
                                                !! Spf error tolerance for relaxation convergence (PRIMARY)
real*8 :: astoptol=1d-7
                                                !! Avector error tolerance for relax (BACKUP - WAS STOPTHRESH)
real*8 :: timestepfac=1d0
                                                !! accelerate relax. multiply par_timestep by this each time
real*8 :: max_timestep=1d10
                                                    maximum time step (limit on exponential growth)
integer :: improvedquadflag=0
                                 !!
                                                !! Use newton iteration not diagonalization for improvedrelax.
                                                1.1
                                                       (1 = A-vector, 2 = orbitals, 3 = both)
                                                !! Waits to turn on orbital quad (2 or 3) until this time
real*8 :: quadstarttime=-1d0
real*8 :: aquadstarttime=-1d0
                                                !! Waits to turn on avector quad (1 or 3) until this time
real*8 :: maxquadnorm=1d10
                                                !! brakes to use if improvedquadflag=2 or 3 is diverging
```

```
real*8 :: quadtol=1d-1
                                                !! Threshold for solution of Newton solve iterations orbitals.
                                11
integer :: quadprecon=1
                                1.1
                                                !! Precondition newton iterations for A-vector?
integer :: quadorthflag=0
                                !!
                                                !! If eigenfunctions are becoming linearly dependent try this
integer :: normboflag=0
                                1.1
                                                !! Enforce norm at each r value
   ORBITALS (SINGLE PARTICLE FUNCTIONS, SPFS)
integer :: nspf=1
                                 !! Nspf=
                                                !! number of orbitals
integer :: numfrozen=0
                                 !!
                                                !! number of doubly occ orbs (removed from calculation)
   CONFIGURATIONS
                                                !! For restrictflag=1: 2*m_s: 2x total m_s (multiplicity of
integer :: restrict_ms=0
                                 !!
                                                      lowest included spin states minus one)
integer :: spin_restrictval=0
                                                !! For all spin project=1: determines spin. Default high spin S=M_s.
                                 !!
                                                !! To override use this variable. Equals 2S if S^2 eigval is S(S+1)
integer :: all_spinproject=1
                                 !!
                                                !! Constrain S(S+1) for propagation?
" INITIALIZATION
integer :: loadavectorflag=0
                                 !! A=file
                                                !! load avector to start calculation?
                                                !! number of avector files containing a-vectors to load into the
integer :: numavectorfiles=1
                                !!
                                                !! mcscfnum available slots, or if load_avector_product.ne.0,
                                                !! number of one-wfn files for product, w/ total numelec=numelec
integer :: load_avector_product=0
                                                !! make product wave function for multiple-molecule load
integer :: loadspfflag=0
                                !! Spf=file
                                                !! load spfs to start calculation?
                                                !! (Otherwise, core eigenfunctions.)
integer :: reinterp_orbflag=0
                                                !! sinc dvr only, half spacing interpolation for orb load
integer :: spf_gridshift(3,100)=0
                                                !! sinc dvr only, shift orbitals on read, slow index spffile
integer :: numspffiles=1
                                !!
                                                !! for multiple-molecule (e.g. chemistry) calcs, load many
integer :: numskiporbs=0
                                !!
                                                !! Reading orbs on file(s), skips members of combined set.
integer :: orbskip(1000)=0
                                1.1
                                                !! Which to skip
character (len=200) :: &
                                !! A=file
                                                !! A-vector binary file to read. Can have different configs
    avectorfile(MXF)="Bin/avector.bin"
                                                !! but should have same number of electrons.
character (len=200) :: &
                                !! Spf=file
                                                !! Spf file to read. Can have fewer m vals, smaller radial
    spffile(MXF)="Bin/spfs.bin"
                                                     grid, or fewer than nspf total orbitals.
integer :: avecloadskip(100)=0
integer :: numholes=0
                                                !! Load a-vector with this many more electrons and annihilate
integer :: numholecombo=1
                                                !! Number of (products of) annihilation operators to combine
                                                !!
                                                     (for spin adapt)
integer :: numloadfrozen=0
                                                !! For loading a vector with orbitals to be frozen (dangerous)
integer, allocatable :: myavectorhole(:,:,:)
                                                !! Namelist input is avectorhole. Fast index numholes (#
                                                     annihilation operators to multiply together); then
                                                     numholecombo, number of such products to combine; then
                                                !!
                                                     mcscfnum, wfn of current propagation.
                                                !!
integer :: excitations=0
                                                !! Similar to holes: number of products of excitation ops
integer :: excitecombos=1
                                                    number of products to linearly combine
integer, allocatable:: myavectorexcitefrom(:,:,:) !! Similar to avectorhole. Namelist input avectorexcitefrom, etc.
integer,allocatable:: myavectorexciteto(:,:,:)
                                                !!
                                                !! For both excite and hole: value is spin orbital index
                                                !! 1=1alpha, 2=1beta, 3=1alpha, etc.
                                                !! negative input -> negative coefficent
  INPUT / OUTPUT
character(len=200):: finalstatsfile="Dat/finalstats.dat"
                                                             !! output for relaxation calculation
real*8 :: pulseft_estep=0.01d0
                                                !! energy step in hartree for Pulseft.Dat output files
                           avectoroutfile="Bin/avector.bin"
character (len=200) ::
                                                            !! A-vector output file.
                           spfoutfile="Bin/spfs.bin"
character (len=200) ::
                                                             !! Spf output file.
character(len=200):: psistatsfile="Dat/psistats.dat"
                                                             !! for action 25
integer :: psistatfreq=1
                                                             11 "
character(len=200):: dendatfile="Dat/denmat.eigs.dat"
                                                             !! if notiming=0
character(len=200):: denrotfile="Dat/denmat.rotate.dat"
                                                             11 "
character(len=200):: rdendatfile="Dat/rdenmat.eigs.dat"
                                                             !! deprecated
```

```
character (len=200) :: ovlspffiles(50)="Bin/ovl.spfs.bin"
                                                              !! for actions 20 and 26
character (len=200) :: ovlavectorfiles(50)="Bin/ovl.avector.bin"!!
                                                                        (see numovlfiles in ACTIONS)
character(len=200):: outovl="Dat/Overlaps.dat"
                                                              !! for action 20
character(len=200):: outmatel="Dat/Matel.dat"
                                                              !! for action 20
character(len=200):: zdipfile="Dat/ZDipoleexpect.Dat"
                                                              !! for action 21
character(len=200):: zdftfile="Dat/ZDipoleft.Dat"
                                                              !! "
character(len=200):: ydipfile="Dat/YDipoleexpect.Dat"
                                                              !!
character(len=200):: ydftfile="Dat/YDipoleft.Dat"
                                                              1.1
                                                              !! "
character(len=200):: xdipfile="Dat/XDipoleexpect.Dat"
                                                              !! "
character(len=200):: xdftfile="Dat/XDipoleft.Dat"
character(len=200):: corrdatfile="Dat/Correlation.Dat"
                                                              !! for action 1
character(len=200):: corrftfile="Dat/Corrft.Dat"
                                                              !! "
character(len=200):: fluxmofile="Flux/flux.mo.bin"
                                                              !! for actions 15,16,17,23
character(len=200):: fluxafile="Flux/flux.avec.bin"
                                                              11
character(len=200):: spifile="Dat/xsec.spi.dat"
                                                              !! for action 16 (cross section)
                                                              !! " (total flux(t) without e_ke resolution)
character(len=200):: gtaufile="Dat/gtau.dat"
character(len=200):: projspifile="Dat/xsec.proj.spi"
                                                              !! for action 17 (partial cross section)
character(len=200):: projgtaufile="Dat/gtau.dat"
                                                              !! " (projected flux(t))
character(len=200):: projfluxfile="Flux/proj.flux.wfn.bin"
                                                                  !! " (see numcatfiles in ACTIONS)
character (len=200):: catspffiles(50)="Bin/cation.spfs.bin"
character (len=200):: catavectorfiles(50)="Bin/cation.avector.bin"!! "
character(len=200):: fluxafile2="Flux/flux.avec.bin"
                                                              !! for action 23
character(len=200):: fluxmofile2="Flux/flux.mo.bin"
                                                              !! "
character(len=200):: natplotbin="Bin/Natlorb.bin"
                                                              !! for actions 2,8
character(len=200):: spfplotbin="Bin/Spfplot.bin"
                                                              !! for actions 3,9
character(len=200):: denplotbin="Bin/Density.bin"
                                                              !! for actions 4,10
character(len=200):: rnatplotbin="Bin/RNatorb.bin"
                                                              !! for actions 5,11
character(len=200):: denprojplotbin="Bin/Denproj.bin"
                                                              !! for actions 6,12
character(len=200):: natprojplotbin="Bin/Natproj.bin"
                                                              !! "
" PULSE. (If tdflag=1)
integer :: numpulses=1
integer :: pulsetype(100)=1
                                                  !! Pulsetype=1: A(t) = pulsestrength * sin(w t)^2,
                                  !!
real*8 :: omega(100)=1.d0
                                  !!
                                                  !! 2: A(t) = strength * sin(w t)^2
real*8 :: omega2(100)=1.d0
                                  1.1
                                                 1.1
                                                                 * sin(w2 t + phaseshift),
real*8 :: pulsestart(100)=0.1d0
                                  1.1
                                                  1.1
real*8 :: phaseshift(100)=0.d0
                                  !!
                                                  !!
                                                        pulsestart < t < pulsestart + pi/w; 0 otherwise</pre>
real*8 :: chirp(100)=0d0
                                  1.1
                                                  1.1
real*8 :: ramp(100)=0d0
real*8 :: longstep(100)=1d0
                                  1.1
                                                 !! Pulsetype 3 available: monochromatic, sinesq start+end
!! NOW COMPLEX
DATATYPE :: pulsestrength(100)=.5d0 !!
                                                  !! A_0 = E_0/omega (strength of field)
real*8 :: intensity(100) = -1.d0 !!
                                                  !! overrides pulse strength. Intensity, 10^16 W cm^-2
real*8 :: pulsetheta(100)=0.d0
                                                  !! angle between polarization and bond axis (radians)
real*8 :: pulsephi(100)=0.d0
                                 1.1
                                                 !! polarization in xy plane
real*8 :: maxpulsetime=1.d20
                                 11
real*8 :: minpulsetime=0.d0
                                 !!
                                                 !! By default calc stops after pulse (overrides finaltime,
                                                 !!
                                                    numpropsteps); this will enforce minimum duration
	ext{!!} ACTIONS may also be specified by Act=X where X is an integer on the command line
integer :: numactions=0
integer :: actions(100)=0
                                                !! ACTIONS
                                1.1
    Act=1
              Autocorrelation; set corrflag=1 for fourier transform
    Act=2
              Save natorbs
11
   Act=3
             Save spfs
   Act=4
              Save density
   Act=5
11
             Save R-natorbs
11
    Act=6
              Save projections of natural configurations (with Mathematica data in NatCurves/)
!!
    Act=7
              Save curve data files in LanCurves/ for Mathematica plotting.
   Act=8
             Enter plotting mode (do not run calculation) and plot natorbs
!!
!! Act=9
              Enter plotting mode and plot spfs
```

```
Act=10
             Enter plotting mode and plot density
11
    Act=11
             Enter plotting mode and plot natorbs in R
!!
    Act=12
             Enter plotting mode and plot projections from act=6
11
    Act=13
             Nuclear FLUX
!!
    Act=14
             Enter plotting mode and analyze nuclear flux
   Act=15
!!
             Save ELECTRONFLUX
             Enter plotting mode and analyze ELECTRONFLUX
   Act=16
   Act=17
             Enter plotting mode and analyze ELECTRONFLUX (projected)
11
11
    Act=18
             Plot denproj from act=6
!!
    Act=19
             Enforce natorbs between steps (experimental)
   Act=20
             Overlaps with supplied eigenfunctions
!!
   Act=21
             Fourier transform dipole moment with pulse for emission/absorption
!!
!!
   Act=22
             With Dirac Frenkel restrition, constraintflag=2, check norm of error - NEEDS
!!
                  dfrestrictflag=2 not 1
             Enter plotting/analysis mode and read flux.bin files from Act=15 for overlaps
11
    Act=23
11
                 between two time dependent wave functions
!!
   Act=24
             keprojector
   Act=25
1.1
             make psistats.dat
             mcscf_matel with supplied eigenfunctions
   ACTION VARIABLES (also see filenames in INPUT/OUTPUT above)
integer :: numovlfiles=1
                                !! see ovlspffiles and ovlavectorfiles in INPUT/OUTPUT
                               !! For keprojector ACTION 24
integer :: nkeproj=200
real*8 :: keprojminenergy=0.04d0 !!
real*8 :: keprojenergystep=0.04d0!!
real*8 :: keprojminrad=30
                               !!
real*8 :: keprojmaxrad=40
                                !!
real*8 :: eground=0.d0
                                               !! energy to shift fourier transform for ACTIONS 1 16 17
                               !! Eground=
                                               !! (AUTOCORRELATION AND PHOTOIONIZATION)
complex*16 :: ceground=(0.d0,0d0)!!
                                               !! input as complex-valued instead if you like
                               !! ACTIONS 1 and 21 (autocorrelation and emission/absorption):
real*8 :: autotimestep=1.d0
                                !! time step for fourier transform
" AUTOCORRELATION, PHOTOIONIZATION and EMISSION/ABSORPTION (ac-
tions 1,16,17,21)
integer :: ftwindowlength=-99 !! FOR ACTIONS 1,16,17,21 options for damping function of time to be transformed
                                  if .ge.0, only damp at end (high frequency cutoff): last ftwindowlength
                             1.1
                                  points in fourier transform are damped by cosine function
                             !! if ftwindowlength not set, use previous (v1.16) windowing function: multiply
integer :: ftwindowpower=1
                                 all points by cos(pi t / 2 / tmax)**ftwindowpower
integer :: fttriwindow=1
                             !! If nonzero override other two options, do straight linear damping function
                             1.1
                                  NOW DEFAULT linear damping looks best v1.16
integer :: ftdiff=0
                             !! fourier transform derivative of dipole moment not dipole moment
   EMISSION/ABSORPTION (action 21)
integer :: hanningflag=0
                                !! for hanning window set nonzero action 1 autocorr
integer :: diptime=100
                                !! For act=20, outputs copies every diptime atomic units
integer :: dipmodtime=100
                                !! do ft every autotimestep*dipmodtime
real*8 :: dipolesumstart=1d10,& !! range for integration of oscillator strength (e.g. for sum rule)
    dipolesumend=1d9
                                     photon energy, atomic units (Hartree), start and end
  PHOTOIONIZATION (actions 15,16,17)
integer :: computeFlux=500, &
                               ! O=All in memory other: MBs to allocate
    FluxInterval=50,&
                                !! Multiple of par_timestep at which to save flux
    FluxSkipMult=1
                                !! Read every this number of time points. Step=FluxInterval*FluxSkipMult
integer :: nucfluxopt=0
                                !! Include imaginary part of hamiltonian from nuc ke
                               !! O=Full ham 1=halfnium
integer :: FluxOpType=1
integer :: numcatfiles=1
                               !! see catspffiles and catavectorfiles in INPUT/OUTPUT for action 17
!!$ IMPLEMENT ME (DEPRECATE fluxinterval as namelist input)
!!$ real*8 :: fluxtimestep=0.1d0
```

#### **!! PLOTTING OPTIONS**

integer :: debugflag=0
real\*8 :: debugfac=1d0

```
integer :: plotmodulus=10
                                !! PlotModulus= !! For saving nat/spf (Act=2, 6), par_timestep interval
                                                     for saving natorbs. plotskip is for stepping over
real*8 :: plotpause=0.25d0
                                !! PlotPause= !!
real*8 :: plotrange=0.2d0
                                !! PlotZ=
                                                !!
                                                      the saved natorbs on read. others are dimensions
real*8 :: plotcbrange=0.001d0
                                                !!
real*8 :: plotxyrange=2.d0
                                !! PlotXY=
                                                !!
real*8 :: plotview1=70.d0
                                !!
                                                !! viewing angle, degrees
real*8 :: plotview2=70.d0
                                1.1
                                                !! viewing angle, degrees
integer :: plotnum=10
                                !! PlotNum=
                                                !! Max number of plots
integer :: plotterm=0
                                !!
                                                !! 0=x11, 1=aqua
integer :: pm3d=1
                                !! PM3D
                                                !! Turn pm3d on when plotting
integer :: plotres=50
                                !!
                                                !! Resolution of plot
integer :: plotskip=1
                                !! PlotSkip=
                                                !! For plotting (Act=3,5,7), number to skip over
                                !! Mult df3 data by factor. For small part of orbs.
real*8 :: povmult=1d0
integer :: povres=10
                                !!
                                                !! Povray resolution
integer :: numpovranges=1
                                11
                                                !! number of magnifications to plot
real*8 :: povrange(10)=(/ 5,15,& !!
                                                !! Povray plotting ranges (unitless - each magnification)
 80,80,80,80,80,80,80,80 /)
real*8 :: povsparse=1.d-3
                                                !! Sparsity threshold for transformation matrix in povray
!! MISC AND EXPERIMENTAL
integer :: conjgpropflag=0
                                 !! for complex Domcke
integer :: pulsewindowtoo=1
                                 !! use window function for E(omega) as well
integer :: nucfluxflag=0
                                !! 0 = both 1 =electronic 2= nuclear NOT nuclear flux action 13,14
logical :: readfullvector=.true.
                                                !! if you have problems with MPI i/o, maybe try this
logical :: walksinturn=.false.
integer :: turnbatchsize=5
integer :: nosparseforce=0
                                !!
                                                !! to override exit with large number of configs, no sparse
integer :: noftflag=0
                                !!
                                                !! turns off f.t. for flux. use for e.g. core hole propag'n.
integer :: timefacforce=0
                                1.1
                                                !! override defaults
integer :: timedepexpect=0 !! expectation value of H_O(t) or H(t) reported
                                !! CMF/VMF
integer :: cmf_flag=1
                                               !! CMF/LMF/QMF or VMF?
integer :: intopt=3
                                !! RK, GBS
                                                !! SPF/VMF Integrator: 0, RK; 1, GBS, 2, DLSODPK
                                                !! for CMF: 3=expo 4=verlet
                                                !! Number of verlet steps per CMF step
integer :: verletnum=80
integer :: jacprojorth=0
                                !! 1: projector = sum_i |phi_i> <phi_i|phi_i>^-1 <phi_i|</pre>
                                !! 0:
                                                 sum_i |phi_i> <phi_i|
                                                                          default
integer :: jacsymflag=0
                                !! 1: use WP - PW not (1-P)W 0: default (1-P)W
integer :: jacgmatthird=0
                                !! 0: default g (constraintflag.ne.0) is linear operator
                                !! 1: g |phi_c> -> sum_ab |phi_a> g_ab <phi_b|phi_c>
real*8 :: autopermthresh=0.001d0 !! Autoperm=
real*8 :: autonormthresh=0.d0
```

```
!! PARAMETERS FILE FOR PROLATE SPHEROIDAL DVR BASIS and HAMILTONIAN :
             h2parinp NAMELIST input.
!!
module myparams
implicit none
integer :: debugflag=0
!! HAMILTONIAN
real*8 :: nuccharge1=1, &
                              !!
                                               !! Nuclear charges
   nuccharge2=1
                               !!
integer :: twoeattractflag=0
                               !!
                                              !! make 1/r12 attractive
integer :: reducedflag=0
                               !!
                                              !! Use reduced e^- masses? Default 1 (yes) with bornopflag=0
integer :: bornopflag=1
                                               !! Born-Op calculation, or with nuclear KE?
                               !!
                                               !! (Sets &parinp nonuc_checkflag=1.)
real*8 :: pro_Hmass=1836.152701d0!!
                                              !! Masses of nuclei: pro_Hmass is the mass of nucleus one and
!! real*8 :: pro_Dmass=3670.483014d0
real*8 :: pro_Dmass=1836.152701d0!!
                                              !! pro_Dmass is the mass of nuc 2.
integer :: JVALUE=0
                                               !! J value for improved adiabatic
!! Additional atoms - hardwired Z=1 (hydrogen) for now
!! old way: hlocs puts hatoms on gridpoints, hlocrealflag=0
!! new way: turn on hlocrealflag; hlocreal is position r,theta (h2 or he) phi not yet
integer :: numhatoms=0
                                1.1
                                               !! number of h atoms
integer :: hlocs(3,100)=1
                                               !! dimension (3, numhatoms): first 2 indices xi,eta gridpoint
                                !!
                                !!
                                               !! then -1 or 1 for left or right -- all h's in plane for now
integer :: hlocrealflag=0
real*8 :: hlocreal(2,100)=0d0
!! BASIS
integer :: lbig=3,&
                                !!
                                               !! Number of points in eta minus 1
    mbig=0
                                !!
                                               !! Number of m-values exp(imphi)
!! DVR / FEM-DVR for R
II R
integer :: rnumelements=1
                               !!
                                               !! Number of elements in R
                                               !! First element at which ecs begins
integer :: rcelement=100
                               !!
real*8 :: rthetaecs=0.d0
                                              !! ECS scaling angle
                               !!
integer :: rnumpoints=3
                              !!
                                             !! Number of points per element including endpoints in R
real*8 :: relementsize=2d-16
                              !!
                                              !! Size of R elements
real*8 :: rstart=1.4d0
                               !!
                                              !! First (excluded) gridpoint in R
integer :: capflag=0
                               !!
                                              !!
real*8 :: capstrength=0.d0
                               !!
                                               !!
integer :: cappower=2
                               !!
                                               !!
!! XI / r radial electronic DOF
integer :: xinumpoints=14
                                !! Numpts=
                                               !! Number of points per element including endpoints in xi
                                !! Numel=
                                               !! Number of elements in xi
integer :: xinumelements=2
real*8 :: xielementsizes(100)=5 !!
                                               !! Sizes of elements
                                !! Celement=
integer :: xicelement=100
                                               !! First element which is scaled
real*8 :: xiecstheta=0.157d0
                                               !! Scaling angle, radians
                               !!
!! ORBITAL INITIALIZATION
integer :: ivoflag=0
real*8 :: loadedocc(200)=2d0
integer :: num_skip_orbs=0
                                               !! For skipping core orbitals for initial diagonalization.
integer :: orb_skip_mvalue(100)= 99
                                               !! Input how many you want to skip, their index (orb_skip)
integer :: orb_skip(20) = -1
                                               !! (order in energy, each m value) and m-value
```

```
!! PARAMETERS FILE FOR ATOM DVR INPUT: heparinp NAMELIST input.
module myparams
implicit none
integer :: debugflag=0
!! HAMILTONIAN AND BASIS
integer :: hecelement=100,&
                                !! Celement=
                                               !! Frist element that is scaled
    henumelements=2,&
                               !! Numel=
                                               !! Number of elements in r
    henumpoints=14
                               !! Numpts=
                                               !! Number of points per elements
real*8 :: heelementsizes(100)=5,&!!
                                               !! Sizes of elements in r
    heecstheta=0.157d0
                                               !! ECS scaling angle
                               !!
integer :: lbig=3,mbig=0
real*8 :: nuccharge1=1d0
!! ORBITAL INITIALIZATION
integer :: ivoflag=0
real*8 :: loadedocc(200)=2d0
integer :: num_skip_orbs=0
                                               !! For skipping core orbitals for initial diagonalization.
                                               !! Input how many you want to skip, their index (orb_skip)
integer :: orb_skip_mvalue(100)= 99
integer :: orb_skip(20)= -1
                                               !! (order in energy, each m value) and m-value
```

```
!! PARAMETERS FILE FOR SINC DVR POLYATOMIC BASIS AND HAMILTONIAN:
             sincparinp namelist input.
#include "Definitions.INC"
module myparams
implicit none
!! FOR TOTAL ORBITAL PARALLELIZATION, SET orbparflag=.true., AND parorbsplit=3 in &parinp
logical :: orbparflag=.false.
!! integer :: orbparlevel=3 !! in namelist, but not in myparams
!! THE FOLLOWING FLAG IS THEN RELEVANT. Option for parallel KE matvec, rate limiting step.
integer :: zke_paropt=1   !! O=sendrecv 1=SUMMA (bcast before) 2=reduce after
!! fft_batchdim: determines batch size for matrix elements and
!! fft_circbatchdim: determines sub batch size for FFT
11
     defaults set small (less memory, more MPI messages) to avoid MPI problems when doing large
     calculations. Otherwise bigger values will be faster. There is a message size sweet spot
11
    on many machines.
integer :: fft_batchdim=1
                             !! 1 = do nspf matrix elements in nspf batches (less memory)
                             !! 2 = do nspf^2 in one batch (faster unless MPI problems)
integer :: fft_circbatchdim=1 !! 0,1,2, circbatchdim < batchdim; larger faster unless MPI problems</pre>
                                    !! fft_mpi_inplaceflag:
integer :: fft_mpi_inplaceflag=1
                                    !! 0 = out-of-place fft, out-of-place fft inverse
                                          3d FFT + (summa/circ C.T. depending on fft_ctflag)
                                    !! 1 = 3 x (1d FFT , all-to-all index transposition)
integer :: fft_ct_paropt=1
                                    !! fft_ct_paropt, relevant if fft_mpi_inplaceflag=0
                                       like zke_paropt: 0 = sendrecv 1 = summa
integer :: num_skip_orbs=0
integer :: orb_skip(200)=-1
integer :: toothnsmall=40
integer :: toothnbig=240
integer :: numcenters=1
integer :: centershift(3,100)=0
                                 !! grid point index for each center
real*8 :: nuccharges(100)=2d0
integer :: numpoints(100)=15
real*8 :: spacing=0.25d0
integer :: orblanorder=500
                                 !! krylov order for block lanczos orbital calculation
integer :: orblancheckmod=10
                                 !! check every
real*8 :: orblanthresh=1d-4
integer :: capflag=0
                                 !! Number of complex absorbing potentials
integer :: capmode=0
                                 !! Capmode=1 is
integer :: cappower(100)=2
                                 !! v_i(r)= capstrength_i*(r/capstart_i)^cappower_i
real*8 :: capstrength(100)=0.01d0 !! v_i(r)= capstrength_i*max(0,r-capstart_i)^cappower_i
real*8 :: mincap=0d0 , maxcap=1d30 !! V_CAP = -i* max(mincap,min(maxcap,sum_i v_i))
integer :: maskflag=0
integer :: masknumpoints=0
                                 !! 1 = SMOOTH EXTERIOR COMPLEX SCALING
integer :: scalingflag=0
real*8 :: scalingdistance=10000d0 !! atomic units (bohr)
real*8 :: smoothness=5
                                 !! atomic units (bohr)
real*8 :: scalingtheta=0d0
                                !! scaling angle
real*8 :: scalingstretch=1d0
                                !! stretching factor
```

real\*8 :: tinv\_tol=1d-3

!! construct virtual orbtials using density based on loaded orbitals
integer :: ivoflag=0
real\*8 :: loadedocc(200)=2d0

## Chapter 3

# Output

The code outputs a substantial amount of information to screen, which can be redirected e.g. chmctdhf\_diatom Inp=Input.Inp.Relax | tee Outs/Out.relax

During the main propagation loop it shows the expectation value of the energy and the norm. By default the energy is the expectation value of the field-free Hamiltonian  $H_0$ . With the variable timedepexpect=1 set in &parinp, it will give the expectation value of  $H(t) = H_0 + V(t)$ .

```
T= 1.16000 Energy: -0.1170575477E+01 0.3036152171E-10 Norm: 0.1000000000E+01
T= 1.18000 Energy: -0.1170575477E+01 0.3036153835E-10 Norm: 0.1000000000E+01
Saving natorb! 18 3
Saving spf! 18
T= 1.20000 Energy: -0.1170575477E+01 0.3036156777E-10 Norm: 0.1000000000E+01
```

The code saves the wave function in Bin/spfs.bin and Bin/avector.bin at the end of the calculation; it will also save at intervals during the propagation if you set saveflag=1. The code makes the directories

```
WALKS
Flux
Dat
Bin
timing
```

Some large files may end up in these directories. Beware of this. On a supercomputer e.g. NERSC machines, you **MUST** be reading and writing these files to the \$SCRATCH directory. If one prefers to run in the home file system one may make symbolic links to the scratch directory. E.g.

```
> mkdir $SCRATCH/blah5
> ln -s $SCRATCH/blah5 ./WALKS
```

In WALKS it saves walks.BIN, which contains all the information about configurations and spin eigenfunctions, after that information is calculated at the start. Often this initial setup takes a long time; if you rename walks.BIN to savewalks.BIN, it will read the latter when you run a calculation, instead of recalculating the information. That is the reason for saving these large files. Unfortunately there is no option to disable this functionality (writing WALKS) a this time (TO-DO ITEM). It is possible that ln -s /dev/null ./WALKS might work on some systems.

Otherwise, the variable notiming sets the amount of output. notiming=2, least amount of output, is default. notiming=0 gives the maximum including all timing information.

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notiming=2 No optional output Dat/Pulse.Dat If tdflag=1, the pulse waveform A(t) or V(t)Dat/PulseFT.Dat Its F.T. In addition: notiming=1 Orbital occupation numbers. denmat.eigs.dat R-natorb occupation numbers. rdenmat.eigs.dat rdenmat.expect.dat Expectation value of bond length for each R-natorb. PsiStats.Dat Expectation values of various operators for atom DiatomStats.dat1 For diatom In addition: timing and other information notiming=0 expo.dat Information about exponential propagation of orbitals abstiming.dat Time stamp (wall clock) each step Overall timings of all parts of calculation. Main.time.dat Timings of steps of orbital propagation cmf\_prop.time.dat Timings of action of reduced operator during expo prop actreduced.time.dat actreduced.eops.time.dat Further breakdown of operator timings Timings for a-vector propagation cmf\_aprop.time.dat Timings for reduced hamiltonian construction reducedham.time.dat Timings for calculation of matrix elements matel.time.dat Breakdown for two-electron part twoe.time.dat Actions.time.dat Timings for actions

### 3.1 Action output (incomplete list)

Various actions that can be performed, described below, like overlaps, produce output. The main output files are listed below. They can be set in namelist &parinp using the variable name in the left column.

fluxmofile	Bin/flux.mo.bin	Molecular orbitals from Act=15 for photoionization calculation Act=16 or 17
fluxafile	Bin/flux.avec.bin	A-vector for photoionization calculation
spifile	Dat/xsec.spi.dat	Cross section from analysis of total photoionization (Act=16) in megabarns is colu
projfluxfile	<pre>Dat/xsec.proj.X.spi.dat</pre>	Partial cross section for state number X from projected photoionization (Act=17)
corrdatfile	Dat/Correlation.dat	Autocorrelation function for Act=1
corrftfile	Dat/Corrft.dat	Its fourier transform
zdftfile	Dat/ZDipoleft.Dat	Fourier transforms of field E and induced dipole moment D for absorption/emissic
		D(omega), E(omega), D(omega)* x E(omega) * is complex conjugate. Thus
		absorption is column 7. Absorption cross section in Mb is column 9.
zdipfile	Dat/ZDipoleexpect.Dat	Induced dipole moment D(t)
ovlfile	Overlaps.Dat	For Act=20, overlaps, these are the overlaps.

Less frequently used:

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Natlorb.bin Natural orbitals from Action 2; read by Action 8.

Spfplot.bin Similarly, with Act=3 and 9.

Density.bin Similarly, with Act=4 and 10.

RNatorb.bin Similarly, with Act=5 and 11.

Similarly, with Act=6 and 12.

NatCurves/ If Act=6, NatCurves/ contains data files and MC-Nat.txt1,

MC-Nat.txtx MC-Nat.txt2, etc. contain mathematica input.

LanCurves/ If Act=7, LanCurves/ contains data files and MC-Lan.txt1,

MC-Lan.txtx MC-Lan.txt2, etc. contain mathematica input.

Natorb/ If Act=8, plotting natural orbitals, and Povray output is

chosen, the data files go in this directory.

Spfs/ Similarly, with Act=9.
Density/ Similarly, with Act=10.

# Types of calculation

There are five general types of calculation that are generally performed:

- Relaxation for initial state
- Relaxation for many states for projection or overlap (called "mcscf" here)
- Propagation with a pulse
- Propagation without a pulse usually done only for photoionization calculation, after first running pulse
- Analysis, for flux, plotting, and other tasks that use an already calculated wave function: will be described in actions section

#### 4.1 Relaxation calculation

In a relaxation calculation the orbitals are propagated in imaginary time and the configuration coefficients are obtained as eigenvectors. The eigenvectors may be obtained through sparse diagonalization or, if an excited state is desired, by inverse iterations using improvedquadflag.

The most important options for relaxation are

loadavectorflag=0 mcscfnum=3 improvedrelaxflag=2 improvednatflag=1 improvedquadflag=1 stopthresh=1d-8

Given those input parameters, the code will diagonalize the Hamiltonian to start, take eigenvectors 2-4, then use inverse iterations on those vectors as the orbitals are optimized to convergence, always constraining the orbitals to be the averaged natural orbitals of the three states.

The three main types of relaxation calculation possible are:

• Regular time relaxation: solution of time dependent Schrodinger equation in imaginary time for ground state. Set improvedrelaxflag=0, threshflag=1 in &parinp.

- Improved relaxation: improvedrelaxflag>1. Orbitals are propagated in imaginary time and A-vector hamiltonian is diagonalized to obtain the wave function for a ground or excited state. Set by command line option >mctdh Relax=N or variable improvedrelaxflag=N to calculate the Nth eigenvector. Root flipping problems may emerge, in which case use:
- Iterative solve: improvedquadflag can be set to 1 to perform an iterative solve to lock onto an eigenvector, from a good initial guess. For instance, one could converge the first 5 roots using MCSCF, then optimize the 5th with a calculation on that state only using improvedquadflag=1.

One would want to adjust the variables par\_timestep (to a lower value than the default 0.5) and stopthresh (convergence parameter) for difficult cases.

#### • improvednatflag=1:

One can rotate the orbitals to be natural orbitals during a relaxation calculation using improvednatflag=1. Thus the final orbitals will be natural orbitals. This is useful if one is to subsequently excite or annihilate the orbitals using avectorexcite or avectorhole. Be sure to check the output in denmat.rotate.dat (second of three columns) to verify that the natorbs are correct. This column should be near zero. If not you should check your orbital integration tolerance (expotol usually). Sparse Avector eigenfunctions and solve may need increased tolerances too. For difficult cases use nonsparse if possible. improvednatflag is compatible with improvedquadflag; the iterations, energies, most things should be same. Improvednatflag should not hurt the performance of improvedquadflag.

Using mcscfnum > 1, one can perform a state-averaged MCSCF calculation, and calculate more than one A-vector for the one set of orbitals, for any of the three ways of relaxing mentioned above. This is most commonly done to obtain wave functions for analysis, projected flux (Action #17) or overlaps (Action #20). Just set variable mcscfnum to the number of states desired.

The default is to use core orbitals and eigenfunctions. However, one may wish to read in an arbitrary set of initial wave functions. For instance, one may wish to converge the first five states including an excited  $\Pi$  state using relaxation, then select one component of that  $\Pi$  state and use <code>improvedquadflag=1</code> or 3 to iteratively refine that state.

There are several options available for reading in initial orbitals and wave functions. Consider this contrived example: we start with a 12 orbital calculation, done on three sigma states. Then use those orbitals, loading them, and freezing them with  $spf_flag=0$ , in calculations on pi and delta states. Then, perform an 11 orbital, three state averaged calculation on the third sigma state, one pi state, and one delta state, discarding the 10th orbital, keeping 1-9, 11, 12. The input for that run would include:

```
&parinp
 mcscfnum=3
 loadspfflag=1
  spffile="Bin/spfs.bin.sigma"
 num_skip_orbs=1
 orb_skip=10
                 CHECK THIS
 loadavectorflag=1
 numavectorfiles=3
 avectorfile="Bin/avector.bin.delta", "Bin/avector.bin.pi", "Bin/avector.bin.sigma"
  avecloadskip=0,0,2
 improvedquadflag=1
 spfrestrictflag=1
 mrestrictflag=0
 mrestrictmin=0
 mrestrictmax=2
```

Because the lowest states aren't being calculated, improvedquadflag=1 or 3 must be invoked, otherwise the 3rd sigma state will collapse to the 1st.

When doing a MCSCF calculation, one often wants wave functions with different M-values. Thus, mrestrictflag=0 must be set to include all these configurations. However, spfrestrictflag should in most cases be set to one so that the orbitals have good m quantum numbers. And, one can use mrestrictmin and mrestrictmax as above.

As of version 1.10 a file is produced, Dat/finalstats.dat, after relaxation calculations, which contains information about the symmetry and matrix elements of the final state(s) and orbitals. The filename can be changed by setting variable finalstatsfile in namelist &parinp to another value.

#### 4.1.1 Utility

There is a program MCSCF-matel that may be compiled and used to compute matrix elements between mcscf wavefunctions (in mcscf.bin files). These can have different orbitals. The program takes the same input as the main program, and in addition the following input:

One should check that the usual namelist input includes the information defining configurations and primitive basis; other parameters are not used. The program produces various files:

```
MCMatel.Dat Energies
MCMatel.Ovl.Dat Overlap matrix elements
MCMatel.ZDipole.Dat Dipole matrix elements, parallel
MCMatel.XYDipole.Dat Dipole matrix elements, perpendicular
```

The namelist variable mcmskip can be set to one to just produce the MCMatel.Dat file.

#### 4.2 Pulse calculation

One specifies a pulse calculation using tdflag=1 in namelist &parinp, or Pulse on the command line. Namelist &pulse is then required, in which one specifies the pulse parameters. The calculation is stopped after the pulse is finished (finaltime in &parinp is overridden), except if minpulsetime or maxpulsetime is set in namelist &pulse. One may include multiple pulses.

```
&pulse
  numpulses=2
  pulsetype=3,1
  omega=0.152,0.1
  omega2=0.6,0.3
  intensity=1d-3,1d-2
  pulsetheta=0d0,0d0
  pulsephi=0d0,0d0
  phaseshift=0d0,0.3d0
  chirp=1d0,-1d0
  ramp=0d0,1d0
  longstep=3d0
```

The intensity is in units of 10<sup>16</sup> W cm<sup>-2</sup>. It is converted to pulsestrength, which is the coefficient

of the pulse waveform in the velocity gauge, and which may alternately be specified in the namelist. The pulse waveform is output in Pulse.Dat and its Fourier transform in PulseFT.Dat.

chirp and ramp are only available for velocity. chirp has units of energy and ramp is unitless.

The pulse types are as follows: NOT ALL CORRECT FIXME FIXME FIXME

```
\omega(t) = \quad \operatorname{omega2} + \operatorname{chirp} \times (t - \pi/\operatorname{omega}/2)/(\pi/\operatorname{omega}) A_0(t) = \quad \operatorname{pulsestrength} \times (1 + \operatorname{ramp} \times (t - \pi/\operatorname{omega}/2)/(\pi/\operatorname{omega}/2) t < \pi/\operatorname{omega} t < \pi/\operatorname{omega}/(2 \operatorname{longstep} + 1) \operatorname{or} (t > \pi/\operatorname{omega}/(2 \operatorname{longstep} + 1)) \operatorname{or} (t > \pi/\operatorname{omega}/(2 \operatorname{longstep} + 1))) t < \pi/\operatorname{omega}/(2 \operatorname{longstep} + 1) \operatorname{omega}/(2 \operatorname{longstep} + 1)) t < \pi/\operatorname{omega}/(2 \operatorname{longstep} + 1) \operatorname{omega}/(2 \operatorname{longstep} + 1)) t < \pi/\operatorname{omega}/(2 \operatorname{longstep} + 1) \operatorname{omega}/(2 \operatorname{longstep} + 1) t < \pi/\operatorname{omega}/(2 \operatorname{longstep} + 1) t < \pi/\operatorname{omega}/(2 \operatorname{longstep} + 1) \operatorname{omega}/(2 \operatorname{longstep} + 1) t < \pi/\operatorname{omega}/(2 \operatorname{longstep}/(2 \operatorname{lo
```

#### 4.3 Propagation calculation, no pulse (fourier run)

For flux or other analysis one needs to propagate the wave function with no pulse, called here a fourier run. Specify tdflag=0, improvedrelaxflag=0, and finaltime in namelist &parinp.

#### 4.4 Analysis / plotting

Certain actions (see below) correspond to analysis or plotting routines (actions 8-12, 14, 16-18, 23). If these actions are specified in the input file (numactions and actions in &parinp) or on the command line (e.g. chmctdhf\_diatom Inp=Inp.Inp Act=16 Eground=-49.545) then the code does not perform propagation and instead terminates after the analysis or plotting routine is complete.

## Actions

We call various things that can be done to the wavefunction "actions." Some actions are performed during a MCTDHF calculation; some actions skip the MCTDHF calculation and are used only for analysis or plotting of a run that has already been performed or that is in progress. One can specify multiple actions. However, if an analysis/plotting action is specified, then the first such specified action will be the only action that is performed.

> chmctdh Act=1 Act=2 Act=4...,

or in namelist &parinp, e.g. numactions=3, actions=1,2,4.

```
Act=1
              Autocorrelation;
11
    Act=2
             Save natorbs
!!
    Act=3
              Save spfs
1.1
    Act=4
             Save density
    Act=5
!!
             Save R-natorbs
             Save projections of natural configurations (with Mathematica data in NatCurves/)
!!
    Act=6
11
    Act=7
             Save curve data files in LanCurves/ for Mathematica plotting.
              Enter plotting mode (do not run calculation) and plot natorbs
    Act=8
1.1
    Act=9
             Enter plotting mode and plot spfs
    Act=10
             Enter plotting mode and plot density
!!
    Act=11
             Enter plotting mode and plot natorbs in R
!!
    Act=12
             Enter plotting mode and plot projections from act=6
             Nuclear FLUX
                                                 DEPRECATED
    Act=14
             Enter plotting mode and analyze nuclear flux
1.1
             Save ELECTRONFLUX
    Act=15
!!
    Act=16
             Enter plotting mode and analyze ELECTRONFLUX
!!
    Act=17
             Enter plotting mode and analyze ELECTRONFLUX (projected)
!!
    Act=18
             Plot denproj from act=6
!!
    Act=19
             Enforce natorbs between steps (experimental)
             Overlaps with supplied eigenfunctions
             Fourier transform dipole moment with pulse for emission/absorption
!!
    Act=21
!!
    Act=22
             With Dirac Frenkel restrition, constraintflag=2, check norm of error - NEEDS
!!
                  dfrestrictflag=2 not 1
!!
    Act=23
             Enter plotting/analysis mode and read flux.bin files from Act=15 for overlaps
!!
                   between two time dependent wave functions
!!
    Act=24
             keprojector
!!
    Act=25
             make psistats.dat
             mcscf_matel with supplied eigenfunctions
    Act=26
```

#### 5.1 Autocorrelation, action 1

Fourier transforms of autocorrelation functions are calculated by using &parinp namelist variables corrflag to 1 and setting an action to 1. On the command line these can be done with Act=1.

The autocorrelation is performed at the same time as the propagation, so these settings should go in that input file. The autocorrelation function is output in Correlation.Dat and its Fourier transform is output in Correlation.Dat. The zero of energy is set by eground.

# 5.2 Saving orbitals (actions 2-4) and plotting orbitals (actions 8-10)

See subsection 6.2.

# 5.3 Flux and projected flux for photoionization (actions 15, 16, 17)

To calculate photoionization use the flux and projected flux option. These are specified by actions 15, which saves the data during a propagation run, and actions 16, and 17, which compute total and projected flux, respectively. The analysis routines can be run before the calculation is finished; just specify T= on the command line to ensure you don't read past what's written. In other words you can run a flux calculation in the same working directory in which the correlation calculation is running.

The photon energy is defined as the difference between the absolute energy and eground so be sure to set it!

There are parameters that can be used to control the biorthogonalization for the linear solve via  $X = \exp(-\ln A)B$  using the expokit routine. These may not have a big effect but may need to be varied if the flux calculation is taking an extremely long time.

```
&parinp
biodim
biotol
biocomplex
```

#### 5.3.1 Options for saving wave function during propagation (Action 15)

In order to save the wavefunction you must specify action 15 during the propagation run; this propagation in which you save the wave function is often what we call a Fourier run. The wave function is saved in files Flux/flux.mo.bin and Flux.avec.bin, which filenames can be changed via &parinp namelist input variables fluxmofile and fluxafile, respectively. So that's Act=15 on the command line or namelist input as follows. When reading, action 16 or 17, you also want to specify or check in the input that FluxSkipMult > 1, if you have saved wave functions, given par\_timestep and FluxInterval in the Act=15 Fourier run that saves the wave function, more frequently than is required, to compute the flux-flux correlation function at a high enough photon frequency for your purposes.

```
&parinp
numactions=1
actions=15
FluxInterval Multiple of par_timestep at which the wave function is saved
```

The files Flux/flux.mo.bin and Flux.avec.bin MAY BE VERY VERY LARGE, especially if you are not careful. They contain the entire wave function propagated for the entire run. Do not use an excessively small save interval or you will waste large quantities of disk space. The save interval is FluxInterval times par\_timestep. The Flux directory is separate for a reason! Please note the section at the top of the document about "filesystem etiquette." On a supercomputer with multiple users, ensure that you do not write, e.g., WALKS, timing, and Flux, to the home filesystem; these should always be put on scratch. Many users prefer to operate on \$SCRATCH entirely; that is the most responsible option. Others run in a working directory in the home filesystem, such that the main results are preserved, and symbolically link the directories to scratch. The script findwork is included; this finds a new directory and symbolically links it as WORK in the working directory. One may run

```
prompt> rm -r Flux timing WALKS
prompt> findwork
prompt> mv WORK Flux
prompt> findwork
prompt> mv WORK timing
prompt> findwork
prompt> mv WORK WALKS
   et cetera (perhaps also Dat). The script findwork follows:
# lawrencium
scratchdir=$SCRATCH
# nersc
# scratchdir=$GSCRATCH
i=0; flag=0
while [[ $flag == 0 ]]
        (( i++ ))
        file="$scratchdir/WORK$i"
        if [[ ! -e $file ]]
       then
                flag=1
       fi
done
mkdir $file
# or not
rm -r WORK
ln --symbolic $file ./WORK
```

#### 5.3.2 Total photoionization (Action 16)

```
&parinp
FluxSkipMult Interval for flux read (F.T. timestep=par_timestep x FluxSkipMult x FluxInterval)
fluxoptype Approximation to Im(H): 0=no approx 1=halfnium potential approx (default)
```

(The option fluxoptype=0 is the exact expression, but it has not been used recently and may be buggy. TO-DO DEBUG.)

To calculate flux you run a calculation with action 16. For instance, say you have Input.Inp.Fourier that you are using for the propagation run (which you set with action 15). You could be running

```
> chmctdhf_sinc Inp=Input.Inp.Fourier |tee Outs/Out.Fourier
```

Let's say it has gotten to 300 time units. In the same directory you could run

> chmctdhf\_sinc Inp=Input.Inp.Fourier Act=16 T=300 | tee Outs/Out.Flux.T300

The flux calculation produces xsec.spi.dat (spi for single photoionization). This filename may be changed by &parinp namelist variable spifile.

The analysis of the wave function for photoionization is done separately from the propagation. During propagation, with Act=15, the only thing that is done is the wave function is saved. It is saved every FluxInterval times par\_timestep atomic units.

#### NO - DEPRECATE FLUXINTERVAL AS INPUT - DO FLUXTIMESTEP

Photoionization cross sections are calculated with e.g.

chmctdhf\_diatom Eground=-99.80004 Inp=Input.Inp.Fourier Act=16

for total photoionization.

The output in Dat/xsec.spi.dat is the quantum mechanical cross section in Megabarns ( $10^{-18}$  cm<sup>2</sup>):

$$(5.1)$$
 equation

The cross section in Megabarns is in column 3 and the photon energy in Hartree is in column 1 in the file Dat/xsec.spi.dat.

Note that the equation does NOT include a factor of  $\frac{1}{3}$ . (Now TRUE, NO FACTOR OF 1/3 as of version v1.16.) The perpendicular and parallel cross sections for fixed nuclei H<sub>2</sub>, just like the physical cross section for the molecule, is approximately 12 Megabarns at onset. Many definitions include a factor of  $\frac{1}{3}$  and sometimes  $\frac{2}{3}$  for the perpendicular fixed-nuclei cross section, such that the total cross section is variously the sum or the 1:2 weighted sum of the parallel and perpendicular cross sections, but we simply report the quantum mechanical cross section, that which becomes the classical cross section for the problem as defined, regardless of the dimensionality of nuclear motion that is included.

Remember, the photon energy is defined relative to eground. See the examples section for the script Flux.Bat, which reads eground by grepping from a computed output file from a relaxation run, automating the execution of the chmctdhf Eground= command above.

#### 5.3.3 Partial photoionization (Action 17)

```
&parinp

FluxSkipMult Interval for flux read (F.T. timestep=par_timestep x FluxSkipMult x FluxInterval)
numcatfiles=2 Number of cation state files
catspffiles(1)="Bin/cation.spfs.bin.pi_g" etc.
```

In order to calculate projected flux one must calculate the final (N-1)-electron states for the projection. These must be saved in the files Bin/cation.spfs.bin and Bin/cation.avector.bin. The setting of these filenames is not supported at this time.

One does this with a relaxation run. After you perform the (N-1)-electron relaxation, one must rename the binary files so that they may be read by the flux calculation. You add the prefix "cation." at the start of the filename. Thus you would have cation.spfs.bin and cation.avector.bin.

Projected flux produces files xsec.proj.spi.datN with N being the final state. The partial cross section in Megabarns is given in column 3 of this file.

#### 5.4 Calculating overlaps with given wave functions (action 20)

Using action 20 one may read in wave functions at the start of the calculation and calculate their overlaps with  $\Psi(t)$ . Perform one or more relaxation calculations to obtain the states upon which you want to project. Then, to read them in your propagation run, include the following.

```
&parinp
  numactions=1
  actions=20
  numovlfiles=2
  ovlspffiles(1:2)="spfs.bin.sigma","spfs.bin.pi"
  ovlavectorfiles(1:2)="avector.bin.sigma","avector.bin.pi"
```

# 5.5 Fourier transform of dipole moment for absorption/emission (action 21)

The absorption and emission spectrum is calculated via the "response function"

(5.2) 
$$S(\omega) = 2\omega Im(D(\omega)E^*(\omega))$$

In which  $D(\omega)$  is the fourier transform of the induced dipole moment, and  $E^*(\omega)$  is the complex conjugate of the Fourier transform of the field.

Relevant variables are

```
&parinp
autotimestep=0.5d0
dipmodtime=100
diptime=200
```

The variables don't have great names. The value of the dipole moment will be calculated every autotimestep atomic units. The Fourier transform will be performed every dipmodtime × autotimestep atomic units and the dipole moment and its Fourier transformed will be saved to file in files zdipfile, etc, in namelist &parinp. Every diptime atomic units (dipmodtime is integer and diptime is real\*8) the dipoles and transforms will be archived to files e.g. Dat/ZDipoleft.Dat000200 as per the values of zdipfile, etc.

The columns of the Fourier transform files ZDipoleft.Dat, etc. are as follows. D(omega) is the F.T. of the dipole moment; E(omega) is that of the field. As of v1.16 there is NOW a FACTOR of (2E) where E is photon energy, in columns 6 and 7.

```
## Photon energy (column 1); D(omega) (2,3); E(omega) (4,5); response (6,7); cross sect (9)
## UNITLESS RESPONSE FUNCTION FOR ABSORPTION/EMISSION ( 2 omega Im(D(omega)E(omega)*) ) IN COLUMN 7
## QUANTUM MECHANICAL PHOTOABSORPTION/EMISSION CROSS SECTION IN MEGABARNS (no factor of 1/3) IN COLUMN 9
```

#### 5.6 Overlaps between wave functions on file (actions 15, 23)

The wave function is saved to file with action 15 – which normally is used for actions 16 and 17, total and partial photoionization. However if multiple wave functions are saved to file, then they can be compared with action 23. Copy the Flux/ files for the second calculation to e.g. flux.mo.bin.wfn2 and use

&parinp
fluxafile="Flux/flux.avec.bin.wfn1"
fluxmofile="Flux/flux.mo.bin.wfn1"
fluxafile2="Flux/flux.avec.bin.wfn2"
fluxmofile2="Flux/flux.mo.bin.wfn2"

#### 5.7 Keprojector (action 24)

XXX

#### 5.8 Make psistats.dat (action 25)

In version 0 until version 1.09, file Dat/psistats.dat (this is variable psistatsfile in name list &parinp) was produced or not produced according to the value of variable notiming. In version 1 it was mostly gutted. As of version 1.10 it is now created by action 25 and again contains useful information about the symmetry and expectation value(s) of the wave function(s) being propagated.

#### 5.9 Unsupported/deprecated actions

Other saving (5-7) and plotting actions (11,12,18): Action 5 (R natural orbitals) may work; otherwise these options are not actively supported. 12 and 18 are redundant. Nuclear flux and plotting it (actions 13,14) are deprecated, not supported.

# Viewing the output

#### 6.1 Two auxillary files

There are two files which need to be included in the working directory for some of the plotting routines to work, for actions Act=6, 7, 8, 9, and 10. This functionality and the need for these files can be eliminated by answering no to the questions in the Install script that ask about Povray and Mathematica. If the files are not present in the working directory, the code will do what it can; you will still have the data files in NatCurves/ and LanCurves for Act= 6 and 7, and you will have .df3 files in Spfs/, Natorb/, and/or Density/, but Povray will not be called to produce .tga files, for Act= 8, 9, and 10.

Action		Required file
Act=6	Save natural projections	J.Mat. 00011
Act=7	Save B.O. curves	MC-LinesQQQ.txt
Act=8	Read spfs	)
Act=9	Read natorbs	Density.Bat if ploting with Povray, not gnuplot
Act=10	Read density	

# 6.2 Viewing natural orbitals, orbitals, density, R-natural orbitals, and projections of natural configurations

The code is built with a plotting mode (controlled by internal variable skipstuff) in which it reads the input as normal, but skips various parts of the setup and then goes directly to an interactive plotting subroutine. So if you want to plot any of these results, you re-run the code with the same input file and command line arguments, but additionally with one of the plotting mode actions specified. Viewing natural orbitals, orbitals, density, R-natural orbitals, and projections of natural configurations is accomplished by actions 8 through 12.

Action	Reads file	Comments
Act=8	Natlorb.bin	View natural orbitals using gnuplot or povray.
Act=9	Spfplot.bin	View orbitals using gnuplot or povray.
Act=10	Density.bin	View single particle density using gnuplot or povray.
Act=11	RNatorb.bin	View R-natural orbitals using gnuplot.
Act=12	Natproj.bin	View projections of natural configurations using gnuplot.

So you can re-run your calculation with Act=8, 9, 10, 11, or 12, respectively, as a(n additional) command line argument. You can do this as the MCTDHF calculation is running, if you'd like. Only one of





Figure 6.1: Example of gnuplot orbital plot.

these options may be used at a time. This works with the example directory scripts, so you can run C:> Third.Bat Act=9 for instance. Otherwise for example

```
C:> pmctdh Inp=Input.Inp.myinput Act=10
```

Some of the usual output will appear on screen, and then, for instance,

USE POVRAY? y for yes.

С

3

At this point the plot subroutine branches depending on whether you want to use gnuplot or Povray to produce the output. Gnuplot may output to screen as either an X11 or AquaTerm plot, or as a .gif file, via the "change terminal" option. Povray produces .tga files which must then be assembled into an animation. If you answer no to "USE POVRAY?" then the following menu will appear for gnuplot:

```
What should I do?
      s = change plotskip (now
                                           1)
      n = change plotnum (now
                                         10 )
      z = change zrange (now
                                 0.8000000000000000
      x = change xyrange (now
                                 2.00000000000000000
       t = change terminal ( now x11
      d = change pm3d (now
                                      0)
      v1= change view rotation 1 (now
                                         70.000000000000000
                                                                 ) degrees
      v2= change view rotation 2 (now
                                         70.000000000000000
                                                                  ) degrees
default = continue (plot with these options)
 Enter natorb number. Negative to stop.
```







Figure 6.2: Example of Povray natural orbital plot: the tenth  $(2\sigma_u)$  orbital of a valence-space CAS calculation, directory O2.Small. Left: at 0.47 fs, from a range of 40 bond lengths (which are 2.2819... bohr); middle and right, 0.96fs, from a range of 7 (middle) and 40 (right) bond lengths. The middle and left plots are the same orbital at the same time.

If the options are ok you can enter any input except for the labeled options (for instance, type "c" then enter) to continue to plot. Choose the natural orbital or spf that you want to plot. The program will output some information to screen as it builds the text file for gnuplot. This may take a while. Then, a plot should appear, as in Figure 6.1. You then have the option to plot again.

If you choose yes to "USE POVRAY?" then you will instead only have the following options:

```
What should I do?
    s = change plotskip (now 1)
    n = change plotnum (now 10)
default = continue (plot with these options)
```

If you need to change the other parameters for Povray plots, you will need to do this in the input file, in namelist &parinp:

```
&parinp
    povres=14
    numpovranges=2
    povrange=8.0d0, 50.0d0
    povsparse=1.d-3
    ... /
```

powres determines the resolution of the .df3 file; the data file is  $(2 \times powres+1)^3$  big. numpowranges is the number of distances, in units of R  $a_0$ , at which to view the orbitals or density, and the array powrange contains those distances. The parameter powsparse determines the zero cutoff for the prolate-to-cartesian transformation matrix – it is necessary to store this memory in sparse format for large powres. Select an orbital and the plotting subroutine proceeds:

```
Enter Natorb
                  number. Zero to change options. Negative to stop.
1
Good read at record
                                1
  Maxsparse ...
  Got maxsparse:
                            662316
                                                        11523120.
                            183544
  Got maxsparse:
                                                        11523120.
   Get spherical sparse.
      ...done.
  POV-plotting Natorb !!
                                                     8.0000000000000000
```

Persistence of Vision(tm) Ray Tracer Version 3.6.1 (/usr/bin/g++-4.2 4.2.1 @ i386-apple-darwin10)

...and the povray output will continue. To plot all the natural orbitals or spfs, choose a number of orbitals greater than the number in the calculation. The program outputs .tga files, which may be assembled by you into an animation. The output should appear as in Fig. 6.2. The plot is three dimensional and colors the wavefunction according to its phase on the color wheel. Concentrated parts of the wavefunction may be saturated in what you see. In Fig. 6.2, the middle and left panels are the same orbital at the same time, but at different magnifications; the intensity range is adjusted to different levels according to the magnification of the plot (variable povranges).

## **Examples**

There are several example directories now included in the EXAMPLES-DEPOT directory. As of version 1.20 the following examples are included, and others:

prompt> ls EXAMPLES-DEPOT/
02-photo-PLAIN-SCRIPTS
H2-EXAMPLE-PLAIN-SCRIPTS
HELIUM-EXAMPLE-PLAIN-SCRIPTS
HELIUM-POLY-ABSORPTION-1.8spacing-45pts-MAC-SCRIPTS
Helium-Neon-ICD-LAWRENCIUM-SCRIPTS
Helium.transient.absorption-bothrotating-weak-LAWRENCIUM-SCRIPTS
Methane.abs.0.39495414.36pts-ion2-MAC-SCRIPTS
Methane.abs.0.39495414.60pts-ion-LAWRENCIUM-SCRIPTS

etc. This chapter goes into these examples, but see the README file in each example directory for updated instructions.

In the example directories, you will find bash scripts for running the calculation and plotting the results with gnuplet and doing other things.

You probably want to start with the directories  $\tt H2-EXAMPLE-PLAIN-SCRIPTS$ ,  $\tt HELIUM-EXAMPLE-PLAIN-SCRIPTS$ , and  $\tt O2-photo-PLAIN-SCRIPTS$ .

# 7.1 Total and partial photoionization cross sections: directories H2-EXAMPLE-PLAIN-SCRIPTS, HELIUM-EXAMPLE-PLAIN-SCRIPTS, O2-photo-PLAIN-SCRIPTS

A total or partial photoionization cross section calculation is performed in three or four steps. The same primitive basis should be used for all steps.

- 1) Relaxation for initial state, Input.Inp.Relax in the examples, (e.g. script Relax.Bat) prompt> chmctdhf\_diatom Inp=Input.Inp.Relax | tee Outs/Out.Relax
- 2) For partial photoionization cross sections, one or more Input.Inp.Cations must be run to produce the (N-1)-electron wave functions that are used for projection.

```
prompt> chmctdhf_diatom Inp=Input.Inp.Cation.pi_g | tee Outs/Out.Cation.pi_g
```

• 3) A propagation that is used to construct the flux-flux correlation function as per the formalism of Meyer [CITE] and paper XX. The sharper the features in the photoionization cross section that

done

are to be resolved, and the longer the real part of the grid is, the longer the propagation must be. Useful durations range from 100 to 8000 atomic units (2.5 to 20fs). (script Fourier.Bat)

```
prompt> chmctdhf_diatom Inp=Input.Inp.Fourier |tee Outs/Out.Fourier
```

• 4) The final step is the analysis step for total (Action 16) or partial (Action 17) photoionization. It is run separate from the propagation and can be run while the propagation is still running as long as the final time is specified less than the maximum currently calculated time. (script Flux.Bat)

```
prompt> chmctdhf_diatom Inp=Input.Inp.Fourier Act=16 eground=-1.15645 T=50 | tee Outs/Out.Flux.t50
```

While the Fourier step is running, one may calculate the cross section so far computed (the Flux step). A script Flux.Bat is included in the example directories, e.g.

```
#!/bin/bash
if [[ "$1 " == " " ]]
then
      echo "need time"
      exit
fi
energy='grep -A 2 CONV Outs/Out.relax |tail -n 1|tr -s " "|cut -f 2 -d " "'
echo "ENERGY $energy"
cp -p Dat/xsec.spi$2.dat Dat/xsec.spi$2.dat.t$1
  For O2-photo, many cation states are calculated. Here is RunCat.Bat:
for ext in doub quart; do for sym in sig pi; do for ger in u g
mpirun -n 1 chmctdhf_diatom.seq Inp=Input.Inp.Cation.$ext.${sym}_${ger} | tee Outs/Out.cation.$ext.${sym}_${ger} &
done; done; done
  Script ProjFlux.Bat will calculate the partial photoionization cross sections.
#!/bin/bash
if [[ "$1 " == " " ]]
then
      echo "need time"
      exit
fi
energy='grep -A 2 CONV Outs/Out.relax |tail -n 1|tr -s " "|cut -f 2 -d " "'
echo "ENERGY $energy"
for file in Dat/xsec.proj.spi${2}*_*dat
      mv $file $file.t$1
```

For  $O_2$  partial photoionization there are many final cation states with different symmetries. As of version 1.20 these are now all loaded with namelist variable numcatfiles=8.

# 7.2 H2-EXAMPLE-PLAIN-SCRIPTS, HELIUM-EXAMPLE-PLAIN-SCRIPTS for action 21, absorption/emission, Fourier transform dipole moment

The Fourier step in these examples also produces photoabsorption results via action 21. See the README and scripts gnu.abs.here etc.

- 7.3 Benzene
- 7.4 Diels-Alder
- 7.5 Helium-Neon Interatomic Coulombic Decay
- 7.6 Transient absorption/emission and wave mixing

The code calculates the expectation value of the dipole moment and fourier transforms it using action 21. The output can be post processed. The example directory Helium.transient absorption is included in the EXAMPLES subdirectory.

- 7.6.1 O2.transient.absorption.ONE
- 7.6.2 2D spectra for  $O_2$  and He

# Programmers' guide

Again, the directory structure, right out of the package, is as follows.

```
MCTDH.SRC
   DFFTPACK
   DGMRES
   SINCDVR
   H2PROJECT
   HEPROJECT
COMPDIRS
   MCTDH.SRC -> ../MCTDH.SRC
   BIN.ecs.hermnorm.law.openmp.nofft
      Definitions.INC
      Name.Txt
      Makefile.header -> ../MCTDH.SRC/Makefile.header.lawrencium.openmp.nointelfft
      Definitions.ALL -> ../MCTDH.SRC/Definitions.ALL
      mctdhf.F90 -> ../MCTDH.SRC/mctdhf.f90
         Makefile.header -> ../Makefile.header
         Makefile -> ../../MCTDH.SRC/DFFTPACK/Makefile
         zfftf1.f -> ../../MCTDH.SRC/DFFTPACK/zfftf1.f
      SINCDVR
         Makefile.header -> ../Makefile.header
         Makefile -> ../../MCTDH.SRC/SINCDVR/Makefile
         sincDVR.f90 -> ../../MCTDH.SRC/SINCDVR/sincDVR.f90
   debug.BIN.ecs.hermnorm.mac
   {\tt BIN.ecs.hermnorm.edison}
```

Notice the following things.

- The \*BIN\* compilation directories are identical except for Name.txt, Makefile.header, and Definitions.INC.
- All of the subdirectories of these BIN directories are real subdirectories; they contain nothing but links.
- Notice that most of the symbolic links to fortran files end in .F90 whereas all the fortran files themselves in MCTDH.SRC end in .f90. This is done to invoke the c preprocessor with some compilers (with the .F90) but the fortran interpreter with emacs (with the .f90).
- The compilation directories have Symbolic links and directory-specific Name.txt, Makefile.header, and Definitions.INC.

These BIN directories contain the compiled version of the code and all the object files and miscellaneous files created during compilation. They have subdirectories, symbolic links, and two real files, Definitions.INC and Name.Txt:

Name.Txt contains the name of the program version, e.g. chmcdthf for chmctdhf\_diatom, etc.; alternatively mctdhf, pmctdhf, or cmctdhf; and is used by the Makefile. Definitions.INC is a header for most of the .f90 files that contains preprocesor directives that define data types and the corresponding LAPACK subroutines differently for the different BIN directories, implementing real or complex data types, ECS or no ECS, and hermitian norm or, for ECS, c-norm, mctdhf, chmctdhf, pmctdhf, cmctdhf. To effect the differences, the macros REALGO, ECSFLAG, and CNORMFLAG are either defined or undefined, and then there are conditional statements that do the rest of the work in Definitions.ALL.

```
Definitions.INC :
    #define REALGO
    #define ExxCSFLAG
    #define CxxNORMFLAG

#include "Definitions.ALL"

Definitions.ALL :
    #ifdef REALGO

    #define DATATYPE real*8
    #define MYGEMM DGEMM
    ...

The fortran files then use these c preprocessor macros,
    #include "Definitions.INC"

module automod
    implicit none

DATATYPE, allocatable :: overlaps(:,:)
    integer, allocatable :: calledflags(:)
    ...
```

#### 8.1 Main program files

The MCTDH.SRC directory contains the following files. They are grouped by nature and importance. parameters.f90 and main\_modules.f90 are the main dependencies; most of the other files depend on these files, and none other. Again we reiterate that we sometime use the terms configuration and slater determinant interchangeably. Sometimes when we write configuration, we mean either a slater determinant or a spin adapted sum of slater determinants. But usually, we just mean slater determinant.

```
-rw----- Definitions.ALL
                                          x C preprocessor macros
-rw----- Makefile.header.edison
                                          x Makefile.headers
-rw----- Makefile.header.mac.mpi.debug
                                          x . . . etc
-rw----- Makefile
                                          x Makefile
                                          x Compilation script
-rwxr-xr-x Makeme
drwx---- DFFTPACK
                                          x Fourier transform
drwx---- DGMRES
                                          x GMR.F.S
drwxr-xr-x COREPROJECT
                                          x coreproject.f90
```

drwx	HEPROJECT	x Atom project
drwx	H2PROJECT	x Diatom project
		WIRTHOUGH W. T. J. A. J.
	parameters.f90	x VARIABLES, Namelist &parinp
	main_modules.f90	x Modules
ı w	getparams.f90	x Routine to load input file, otherwise set variables
-rw	mctdhf.f90	x Main program
-rw	prop.f90	x Core of main program
-rw	derivs.f90	x Orbital propagation subroutines - working equation, call of expo_driver
-rw-rr	configstuff.f90	x Subroutine for configuration propagation and diagonalization
-rw	expo_driver.f90	x Orbital propagation subroutines - call of expokit
-rw	matel.f90	x Orbital and configuration matrix element subroutines
-rw	mean.f90	x Constructs 2-e reduced denmat
-rw	denmat.f90	x 1-e reduced denmat and miscellaneous (denmat constraint for restricted configuration list)
-rw	walks.f90	x Matrix elements among slater determinants
	walkmult.f90	x Use them to multiply vector of configuration coefficients
	newconfig.f90	x Configuration subroutines (get configuration list, get configuration index)
-rw		x Spin (S(S+1)) adaptation
	spinwalks.f90	x Matrix elements of spin operator among configurations
-rw-rr	biortho.f90	x Biorthogonalization workhorse
-rw-rr	driving.f90	x Psi-prime treatment
-rw	quad.f90	x Improvedquadflag - inverse iterations for diagonalization
-rw	second_derivs.f90	x Verlet intopt=4
-rw-rr	blocklanczos.f90	x Diagonalization
-rw	dfconstrain.f90	x Restricted configuration list - computation of g and tau, orbital derivatives
-rw	actions.f90	x Action driver routines.
-rw	autocall.f90	
-rw	autosub.f90	
-rw	readactions.f90	
-rw-rr	saveactions.f90	
-rw-rr	natprojaction.f90	
-rw	povactions.f90	
-rw-rr	orbvectoractions.f90	
-rw-rr	dipolecall.f90	
-rw	dipolesub.f90	
-rw	electronflux.f90	
-rw	projeflux.f90	
-rw	ovlsub.f90	
-rw-rr	keprojector.f90	
-rw	MPI.f90	x Parallel subroutines
	proputils.f90	x Utilities, including pulse subroutines
-rw-rr		x Utilities relating to orbitals
-rw		x LAPACK wrappers
-rw-rr	•	x Miscellaneous
	psistats.f90	
	•	x Expectation values of wave function $ imes I/O$
	configload.f90 loadstuff.f90	x 1/0 x 1/0
.T M - T T	100050011.130	A 1/U
-rw	expokit.f	x Exponential propagator for orbitals, also psi-prime & miscellaneous

```
Roger B. Sidje U Queensland modified in key places by me

-rw----- gaussq.f x Quadrature

-rw----- jacobi.f x Quadrature

-rw----- odex.f x Implicit integrator intopt=1

-rw----- opkda1.f x General purpose solver intopt=2

-rw----- rkf45.f x Runge-Kutta intopt=0

-rw----- arg.c x Some c routines
```

#### 8.2 Project directories

The project directories contain files required to build the different compiled versions of the code, running different coordinate systems. In the past, the code had a flag, and all coordinate systems were contained in one program; now it is different.

There are three projects currently, atom, diatom, and polyatomic, in H2PROJECT, HEPROJECT, and SINCPROJECT, respectively. The first two share one file; duplication is avoided by writing one file, MCTDH.SRC/COREPROJECT/coreproject.f90, that is shared. This is not the general operation; project directories, like SINCPROJECT, in general should be autonomous.

To add a new project directory, put the source code in a subdirectory of MCTDH.SRC, perhaps MCTDH.SRC/NEWPROJECT; then create a directory (not a link, mkdir) in one of the BIN directories, also called NEWPROJECT. Then, go into that directory and link each the files in MCTDH.SRC/NEWPROJECT to BIN/NEWPROJECT, with the same name except for capital-F .F90 extension.

That should set up BIN/NEWPROJECT; then you can recursively copy BIN.NEWPROJECT to BIN.ecs.hernorm.debug or any other compilation directories that you have set up (keeping symbolic links as symbolic links!!) and they should be ready to go, if you did your Makefile in NEWPROJECT like it is in the existing project directories.

#### 8.3 About the Makefile and Makeme

The code is meant to be compiled with the script Makeme which should be linked in the BIN directory in question. All it does is run make in the project directories, the other subdirectories, and then in the main directory. So if one adds a project, one adds a line in Makeme, to compile the code in the project directory that you have designed (with your makefile, which should be invoked by the "make" command that Makeme runs in your project directory). In the Makefile, one would add a new project called "newproject" as follows.

The makefile starts out by including the file Makefile.header which has your compiler options, and also the file Name.txt which just has the name of the code for the particular BIN directory. For instance, for BIN.ecs.hermnorm, Name.txt is as follows:

NAME=chmctdhf

#### 8.4 Important variables

It's all about the xarr data type and the variable yyy. But also you need to understand how the sparse matrix routines are done – data types Type(CONFIGPTR) and Type(SPARSEPTR).

The key variables are

ууу

yyy is type xarr that actions.f90 and prop.f90 some other things can use via xxxmod. It contains the wave function and other things, for the current time step (time step 0) and the previous one (time step 1).

yyy%cmfpsivec(:,0)

The wave function (orbital coefficients and A-vector) at the present time step. yyy%cmfpsivec(:,1) is the one previous.

spfstart, astart(mcscfnum)

For psitype=1, a CI wavefunction, these are the indices at which the SPF-vector and A-vector start. Thus, to pass a subroutine the current spfs, use call mysub(yyy%cmfpsivec(spfstart,0)).

spftotdim

This is the size of the SPF-vector.

totadim

This is the size of the A-vector.

...not finished

#### 8.5 Project directories

#### 8.5.1 Functions/subroutines that are necessary to define