progress

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## **Chapter 1**

## **README**

## A library for quantum chemistry solvers.

PROGRESS: Parallel, Rapid O(N) and Graph-based Recursive Electronic Structure Solver. LA-CC-16-068

- This library is focused on the development of general solvers that are commonly used in *quantum chemistry* packages.
- This library has to be compiled with the <code>Basic Matrix Library</code> (BML).
- Our webpage can be found at https://lanl.github.io/qmd-progress/

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(in alphabetical order)

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### **Build Dependencies**

- >=OpenMP-3.1
- >=metis-5.0 if building with PROGRESS\_GRAPHLIB

(On some distributions, metis is available as a package. Make sure you install the -dev package. For example, Ubuntu requires libmetis-dev.)

#### **Build and Install Instructions**

#### How to build

```
$ CMAKE_PREFIX_PATH=<BML install path> ./build.sh
```

#### How to install

```
$ cd build
$ sudo make install
```

#### To specify the Intel Fortran compiler:

```
$ FC=ifort PKG_CONFIG_PATH=<BML install path>/lib/pkgconfig ./build.sh
```

#### To build with the gfortran compiler and OpenMP:

```
$ CC=gcc FC=gfortran \
    CMAKE_BUILD_TYPE=Release \
    PROGRESS_OPENMP=yes \
    CMAKE_PREFIX_PATH=<BML install path> \
    CMAKE_INSTALL_PREFIX=<PROGRESS install path> \
    ./build.sh configure
```

#### To build with OpenMP, MPI and testing enabled:

```
$ CC=mpicc FC=mpif90 \
    CMAKE_BUILD_TYPE=Release \
    PROGRESS_OPENMP=yes \
    PROGRESS_MPI=yes \
    PROGRESS_TESTING=yes \
    CMAKE_PREFIX_PATH=<BML install path> \
    CMAKE_INSTALL_PREFIX=<PROGRESS install path> \
    ./build.sh configure
```

#### To build with OpenMP, MPI, testing enabled and example programs built:

```
$ CC=mpicc FC=mpif90 \
    CMAKE_BUILD_TYPE=Release \
    PROGRESS_OPENMP=yes \
    PROGRESS_MPI=yes \
    PROGRESS_TESTING=yes \
    PROGRESS_EXAMPLES=yes \
    CMAKE_PREFIX_PATH=<BML install path> \
    CMAKE_INSTALL_PREFIX=<PROGRESS install path> \
    ./build.sh configure
```

## To build with OpenMP and MPI and testing enabled and example programs built and the METIS graph partitioning library:

```
$ CC=mpicc FC=mpif90 \
    CMAKE_BUILD_TYPE=Release \
    PROGRESS_OPENMP=yes \
    PROGRESS_MPI=yes \
    PROGRESS_GRAPHLIB=yes \
    PROGRESS_TESTING=yes \
    PROGRESS_EXAMPLES=yes \
    CMAKE_PREFIX_PATH=<BML install path> \
    CMAKE_INSTALL_PREFIX=<PROGRESS install path> \
    ./build.sh configure
```

## Citing

```
@misc{2016progress,
    title={\textrm{PROGRESS} Version 1.0},
    author={Niklasson, Anders M. and Mniszewski, Susan M and Negre, Christian F. A. and Wall, Michael E. and O year={2016},
    url = {https://github.com/lanl/qmd-progress},
    institution={Los Alamos National Laboratory (LANL), Los Alamos, NM (United States)}
}
```

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## **Chapter 2**

# **Testing the Progress library**

## Testing program for the progress library

#### To run the tests:

Go into the build folder and type:

```
make test
```

To run the tests in verbose mode:

```
make test ARGS="-V"
```

### To run a single test:

To run a test on its own (in build) we just need to type:

```
/qmd-progress/build/main <test_name>
```

- , where "test\_name" is the name of the test we want to run. Right now the keywords (test\_name) we can pass are the following:
  - density: Tests the diagonalization routine to build the density.
  - sp2\_short : Tests the first version of sp2
  - sp2\_alg1 : Algorithm 1 for sp2
  - sp2\_alg2 : Algorithm 2 for sp2
  - sp2\_alg2\_ellpack : Algorithm 2 for sp2 with ellpack
  - sp2\_alg1\_seq : See sp2\_mod.F90 source file
  - sp2\_alg2\_seq : See sp2\_mod.F90 source file
  - deorthogonalize\_dense: See nonortho.F90 source file
  - orthogonalize\_dense: See nonortho.F90 source file
  - buildzdiag: See genz\_mod.F90 source file

### To add a test:

- add the corresponding name of the test in /progress/tests/CMakeLists.txt
- add the corresponding keyword and test in /progress/tests/src/main.F90
- Copy any file that is necessary to run (data) in /progress/tests/tests\_data/
- reconfigure and recompile

# **Chapter 3**

# **Todo List**

# Module prg\_dos\_mod

Add LDOS.

Subprogram prg\_pulaycomponent\_mod::prg\_pulaycomponent0 (rho\_bml, ham\_bml, pcm\_bml, threshold, M, bml\_type, verbose)

M and bml\_type will have to be removed from the input parameter.

Subprogram prg\_pulaycomponent\_mod::prg\_pulaycomponentt (rho\_bml, ham\_bml, zmat\_bml, pcm\_bml, threshold, M, bml\_type, verbose)

M and bml type will have to be removed from the input parameter.

## Module prg\_pulaymixer\_mod

add the density matrix mixer.

# Module prg\_response\_mod

Add the response scf

Change name response\_SP2 to dm\_prt\_response

Change name response\_rs to rs\_prt\_response

Subprogram prg\_response\_mod::prg\_pert\_from\_file (prt\_bml, norb)

Add read perturbation from file

Subprogram prg\_system\_mod::prg\_parse\_system (system, filename, extin)

Integrate this loop in the loop for building the splist.

8 Todo List

# **Chapter 4**

# **Module Index**

# 4.1 Modules

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(LATTE related routines)	17
(PROGRESS related routines)	18
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(High-level codes using PROGRESS/LATTE modules)	21

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# **Chapter 5**

# **Modules Index**

# 5.1 Modules List

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Module to obtain the density matrix by applying a Chebyshev polynomial expansion	25
prg_densitymatrix_mod	
Module to obtain the density matrix by diagonalizing an orthogonalized Hamiltonian	33
prg_dos_mod	
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# Chapter 7

# File Index

# 7.1 File List

Here is a list of all files with brief descriptions:

/home/christian/qmd-progress/src/prg_charges_mod.F90
/home/christian/qmd-progress/src/prg_chebyshev_mod.F90
/home/christian/qmd-progress/src/prg_densitymatrix_mod.F90
/home/christian/qmd-progress/src/prg_dos_mod.F90
/home/christian/qmd-progress/src/prg_doxy_mod.F90
/home/christian/qmd-progress/src/prg_extras_mod.F90
/home/christian/qmd-progress/src/prg_genz_mod.F90
$/home/christian/qmd-progress/src/prg\_graph\_mod.F90 \\ \ldots \\ \ldots \\ 23$
$/home/christian/qmd-progress/src/prg\_graphsp2parser\_mod. F90 \\ \ldots \\ 2322 \\ \ldots \\ 2322 \\ \ldots \\ 2322 \\ \ldots \\$
/home/christian/qmd-progress/src/prg_homolumo_mod.F90
$/home/christian/qmd-progress/src/prg\_implicit\_fermi\_mod.F90 \\ \ldots \\ \ldots \\ 23333333333333333333333333333$
/home/christian/qmd-progress/src/prg_initmatrices_mod.F90
/home/christian/qmd-progress/src/prg_kernelparser_mod.F90
/home/christian/qmd-progress/src/prg_nonortho_mod.F90
/home/christian/qmd-progress/src/prg_normalize_mod.F90
/home/christian/qmd-progress/src/prg_openfiles_mod.F90
/home/christian/qmd-progress/src/prg_parallel_mod.F90
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/home/christian/qmd-progress/src/prg_ptable_mod.F90
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/home/christian/qmd-progress/src/prg_quantumdynamics_mod.F90
/home/christian/qmd-progress/src/prg_response_mod.F90
$/home/christian/qmd-progress/src/prg\_sp2\_fermi\_mod.F90 \\ \ldots \\ \ldots \\ 24e^{-1}$
/home/christian/qmd-progress/src/prg_sp2_mod.F9024
/home/christian/qmd-progress/src/prg_sp2parser_mod.F90
$/home/christian/qmd-progress/src/prg\_subgraphloop\_mod. F90 \\ \\ 240$
/home/christian/qmd-progress/src/prg_syrotation_mod.F90
/home/christian/qmd-progress/src/prg_system_mod.F90
/home/christian/qmd-progress/src/prg_timer_mod.F90
/home/christian/qmd-progress/src/prg_xlbo_mod.F90
/home/christian/gmd-progress/src/prg_xlkernel_mod.F90

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# **Chapter 8**

# **Module Documentation**

8.1 (LATTE related routines)

# 8.2 (PROGRESS related routines)

#### **Modules**

· module prg\_charges\_mod

A module to compute the Mulliken charges of a chemical system.

· module prg\_chebyshev\_mod

Module to obtain the density matrix by applying a Chebyshev polynomial expansion.

module prg\_densitymatrix\_mod

Module to obtain the density matrix by diagonalizing an orthogonalized Hamiltonian.

• module prg\_extras\_mod

Extra routines.

module prg\_genz\_mod

To produce a matrix Z which is needed to orthogonalize H.

module prg\_graph\_mod

The graph module.

module prg\_graphsp2parser\_mod

Graph partitioning SP2 parser.

· module prg\_homolumo\_mod

The homolumo module.

· module prg\_initmatrices\_mod

Initialization module.

· module prg\_kernelparser\_mod

Some general parsing functions.

module prg\_nonortho\_mod

Module to prg\_orthogonalize and prg\_deorthogonalize any operator.

• module prg\_openfiles\_mod

Module to handle input output files for the PROGRESS lib.

module prg\_parallel\_mod

The parallel module.

• module prg\_progress\_mod

The progress module.

module prg\_ptable\_mod

Periodic table of elements.

· module prg\_pulaycomponent\_mod

Produces a matrix to get the Pulay Component of the forces.

• module prg\_pulaymixer\_mod

Pulay mixer mode.

• module prg\_quantumdynamics\_mod

A module to add in common quantum dynamical operations.

• module prg\_response\_mod

Module to compute the density matrix response and related quantities.

· module prg\_sp2\_fermi\_mod

The SP2 Fermi module.

module prg\_sp2\_mod

The SP2 module.

module prg\_sp2parser\_mod

SP2 parser.

· module prg syrotation mod

A module to rotate the coordinates of a sybsystem in chemical systems.

module prg\_system\_mod

A module to read and handle chemical systems.

• module prg\_timer\_mod

The timer module.

module prg\_xlbo\_mod

A module to perform XLBO integration.

• module prg\_xlkernel\_mod

Add name.

# 8.2.1 Detailed Description

8.3 (EXTERNAL related routines)

8.4 (High-level codes using PROGRESS/LATTE modules)

# **Chapter 9**

# **Module Documentation**

# 9.1 prg\_charges\_mod Module Reference

A module to compute the Mulliken charges of a chemical system.

## **Functions/Subroutines**

- subroutine, public prg\_get\_charges (rho\_bml, over\_bml, hindex, charges, numel, spindex, mdimin, threshold)

  Constructs the charges from the density matrix.
- subroutine, public prg\_get\_hscf (ham0\_bml, over\_bml, ham\_bml, spindex, hindex, hubbardu, charges, coulomb\_pot\_r, coulomb\_pot\_k, mdimin, threshold)

Constructs the SCF Hamiltonian given H0, HubbardU and charges. This routine does:  $H = \sum_i U_i q_i + V_i$ ;, where U is the Hubbard parameter for every atom i. V is the coulombic potential for every atom i.

## **Variables**

• integer, parameter dp = kind(1.0d0)

# 9.1.1 Detailed Description

A module to compute the Mulliken charges of a chemical system.

This module contains routines that compute properties related to charges.

# 9.1.2 Function/Subroutine Documentation

9.1.2.1 subroutine, public prg\_charges\_mod::prg\_get\_charges ( type(bml\_matrix\_t), intent(inout) rho\_bml, type(bml\_matrix\_t), intent(inout) over\_bml, integer, dimension(:,:), intent(in) hindex, real(dp), dimension(:), intent(inout), allocatable charges, real(dp), dimension(:), intent(in) numel, integer, dimension(:), intent(in) spindex, integer, intent(in) mdimin, real(dp), intent(in) threshold)

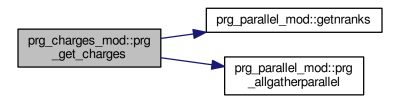
Constructs the charges from the density matrix.

## **Parameters**

rho_bml	Density matrix in bml format.
over_bml	Overlap matrix in bml format.
hindex	Start and end index for every atom in the system.
charges	Output parameter that gives the vectorized charges.
threshold	Threshold value for matrix elements.

Definition at line 31 of file prg\_charges\_mod.F90.

Here is the call graph for this function:



9.1.2.2 subroutine, public prg\_charges\_mod::prg\_get\_hscf ( type(bml\_matrix\_t), intent(in) ham0\_bml, type(bml\_matrix\_t), intent(in) over\_bml, type(bml\_matrix\_t), intent(inout) ham\_bml, integer, dimension(:), intent(in) spindex, integer, dimension(:), intent(in) hindex, real(dp), dimension(:), intent(in) hubbardu, real(dp), dimension(:), intent(in) coulomb\_pot\_r, real(dp), dimension(:), intent(in) coulomb\_pot\_k, integer, intent(in) mdimin, real(dp), intent(in) threshold)

Constructs the SCF Hamiltonian given H0, HubbardU and charges. This routine does:  $H = \sum_i U_i q_i + V_i$ ;, where U is the Hubbard parameter for every atom i. V is the coulombic potential for every atom i.

### **Parameters**

ham_bml	Hamiltonian in bml format.
over_bml	Overlap in bml format.
hindex	Start and end index for every atom in the system.
hubbardu	Hubbard parameter for every atom.
charges	Charges for every atom.
coulomb_pot←	Coulombic potential (r contribution)
_r	
coulomb_pot←	Coulombic potential (k contribution)
_k	
mdim	Maximum nonzeroes elements per row for every row.
threshold	Threshold value for matrix elements.

Definition at line 101 of file prg\_charges\_mod.F90.

#### 9.1.3 Variable Documentation

9.1.3.1 integer, parameter prg\_charges\_mod::dp = kind(1.0d0) [private]

Definition at line 17 of file prg charges mod.F90.

# 9.2 prg chebyshev mod Module Reference

Module to obtain the density matrix by applying a Chebyshev polynomial expansion.

## **Data Types**

type chebdata type

General Cheb solver type.

#### **Functions/Subroutines**

• subroutine, public prg parse cheb (chebdata, filename)

Chebyshev parser. This module is used to parse all the input variables for the cheb electronic structure solver. Adding a new input keyword to the parser:

• subroutine, public prg\_build\_density\_cheb (ham\_bml, rho\_bml, athr, threshold, ncoeffs, kbt, ef, bndfil, jon, verbose)

Builds the density matrix from  $H_0$  for a Fermi function approximated with a Chebyshev polynomial expansion.

• subroutine, public prg\_build\_density\_cheb\_fermi (ham\_bml, rho\_bml, athr, threshold, ncoeffs, kbt, ef, bndfil, getef, fermitol, jon, npts, trkfunc, verbose)

Builds the density matrix from  $H_0$  for a Fermi function approximated with a Chebyshev polynomial expansion. In this case the self-consistent recursion is applied to converge to the correct number of electrons and obtain the Fermi level.

• real(dp) function jackson (ncoeffs, i, jon)

Evaluates the Jackson Kernel Coefficients.

• subroutine <a href="mailto:prg\_get\_chebcoeffs">prg\_get\_chebcoeffs</a> (npts, kbt, ef, ncoeffs, coeffs, emin, emax)

Gets the coefficients of the Chebyshev expansion.

subroutine prg\_get\_chebcoeffs\_fermi\_bs (npts, kbt, ef, tracesT, ncoeffs, coeffs, emin, emax, bndfil, norb, tol, jon, verbose)

Gets the coefficients of the Chebyshev expansion with Ef computation.

• subroutine <a href="mailto:prg\_get\_chebcoeffs\_fermi\_nt">prg\_get\_chebcoeffs\_fermi\_nt</a> (npts, kbt, ef, tracesT, ncoeffs, coeffs, emin, emax, bndfil, norb, tol, jon, verbose)

Gets the coefficients of the Chebyshev expansion with Ef computation.

real(dp) function tr (r, x)

Chebyshev polynomial obtained by recursion.

• real(dp) function fermi (e, ef, kbt)

Gives the Fermi distribution value for energy e.

• real(dp) function absmaxderivative (func, de)

Gets the absolute maximum of the derivative of a function.

#### **Variables**

- integer, parameter dp = kind(1.0d0)
- real(dp), parameter pi = 3.14159265358979323846264338327950\_dp

# 9.2.1 Detailed Description

Module to obtain the density matrix by applying a Chebyshev polynomial expansion.

See Amparo Gil 2007 [Amparo2007] , See Silver et al [Silver1996] , See Weisse et al [Weisse2006]

## 9.2.2 Function/Subroutine Documentation

9.2.2.1 real(dp) function prg\_chebyshev\_mod::absmaxderivative ( real(dp), dimension(:), intent(in) func, real(dp), intent(in) de
) [private]

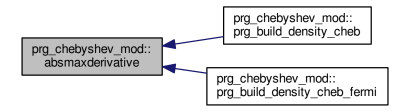
Gets the absolute maximum of the derivative of a function.

#### **Parameters**

func.	
de	Energy step.

Definition at line 802 of file prg\_chebyshev\_mod.F90.

Here is the caller graph for this function:



9.2.2.2 real(dp) function prg\_chebyshev\_mod::fermi ( real(dp), intent(in) e, real(dp), intent(in) ef, real(dp), intent(in) kbt )

[private]

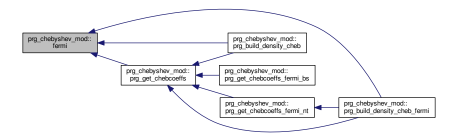
Gives the Fermi distribution value for energy e.

## **Parameters**

е	Energy.
ef	Fermi energy.

Definition at line 790 of file prg\_chebyshev\_mod.F90.

Here is the caller graph for this function:



9.2.2.3 real(dp) function prg\_chebyshev\_mod::jackson ( integer, intent(in) ncoeffs, integer, intent(in) i, logical, intent(in) jon )

[private]

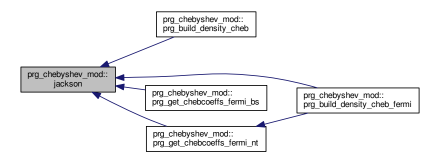
Evaluates the Jackson Kernel Coefficients.

#### **Parameters**

ncoeffs	Number of Chebyshev polynomial.
i	Coefficient number i.

Definition at line 532 of file prg\_chebyshev\_mod.F90.

Here is the caller graph for this function:



9.2.2.4 subroutine, public prg\_chebyshev\_mod::prg\_build\_density\_cheb ( type(bml\_matrix\_t), intent(in) ham\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp), intent(in) athr, real(dp), intent(in) threshold, integer, intent(in) ncoeffs, real(dp), intent(in) kbt, real(dp), intent(in) ef, real(dp), intent(in) bndfil, logical, intent(in) jon, integer, intent(in) verbose)

Builds the density matrix from  $H_0$  for a Fermi function approximated with a Chebyshev polynomial expansion.

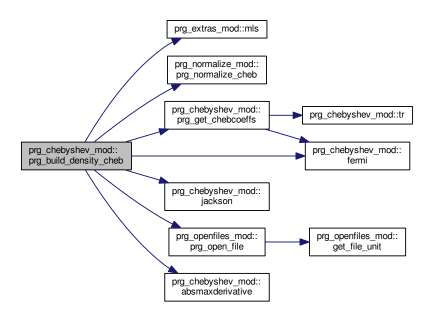
 $ho_{n+1}=b_{n+1}T_{n+1}+
ho_n$  Where,  $T_n$  is the nth Chebyshev polynomial and  $b_n$  is the nth coefficient of the expansion for the Fermi function. In the sparse version (when ellpack is used) the threshold can be varied linearly with the polynomial degree. The function is the following:  $Thresh(n)=Thresh_0[a_{thr}(n-1)+(1-a_{thr})]$ 

#### **Parameters**

ham_bml	Input Orthogonalized Hamiltonian matrix.
rho_bml	Output density matrix.
athr	Threshold linear increasing constant.
threshold	Threshold for sparse matrix algebra.
ncoeffs	Number of Chebyshev coefficients.
kbt	Electronic temperature in the energy units of the Hamiltonian.
ef	Fermi level in the energy units of the Hamiltonian.
bndfil	Band filing factor.
verbose	Verbosity level.

Definition at line 143 of file prg\_chebyshev\_mod.F90.

Here is the call graph for this function:



9.2.2.5 subroutine, public prg\_chebyshev\_mod::prg\_build\_density\_cheb\_fermi ( type(bml\_matrix\_t), intent(in) ham\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp), intent(in) athr, real(dp), intent(in) threshold, integer, intent(in) ncoeffs, real(dp), intent(in) kbt, real(dp), intent(inout) ef, real(dp), intent(in) bndfil, logical, intent(in) getef, real(dp) fermitol, logical, intent(in) jon, integer npts, logical, intent(in) trkfunc, integer, intent(in) verbose)

Builds the density matrix from  $H_0$  for a Fermi function approximated with a Chebyshev polynomial expansion. In this case the self-consistent recursion is applied to converge to the correct number of electrons and obtain the Fermi level.

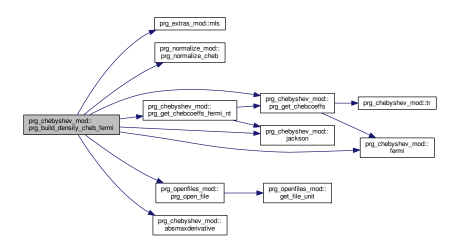
 $ho_{n+1}=b_{n+1}T_{n+1}+
ho_n$  Where,  $T_n$  is the nth Chebyshev polynomial and  $b_n$  is the nth coefficient of the expansion for the Fermi function. In the sparse version (when ellpack is used) the threshold can be varied linearly with the polynomial degree. The function is the following:  $Thresh_0[a_{thr}(n-1)+(1-a_{thr})]$ 

#### **Parameters**

ham_bml	Input Orthogonalized Hamiltonian matrix.
rho_bml	Output density matrix.
athr	Threshold linear increasing constant.
threshold	Threshold for sparse matrix algebra.
ncoeffs	Number of Chebyshev coefficients.
kbt	Electronic temperature in the energy units of the Hamiltonian.
ef	Fermi level in the energy units of the Hamiltonian.
bndfil	Band filing factor.
npts	Number of energy points to compute the coefficients
verbose	Verbosity level.

Definition at line 310 of file prg\_chebyshev\_mod.F90.

Here is the call graph for this function:



9.2.2.6 subroutine prg\_chebyshev\_mod::prg\_get\_chebcoeffs ( integer, intent(in) npts, real(dp), intent(in) kbt, real(dp), intent(in) ef, integer, intent(in) ncoeffs, real(dp), dimension(:), intent(inout) coeffs, real(dp), intent(in) emin, real(dp), intent(in) emax ) [private]

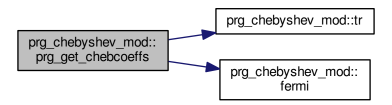
Gets the coefficients of the Chebyshev expansion.

#### **Parameters**

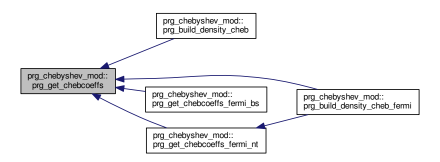
npts	Number of points for discretization.
kbt	Electronic temperature.
ef	Fermi level.
ncoeffs	Number of Chebyshev coefficients.
coeffs	Output vector for the Chebyshev coefficients.
emin	lowest boundary for the eigenvalues of H.
emax	highest boundary for the eigenvalues of H.

Definition at line 568 of file prg\_chebyshev\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



9.2.2.7 subroutine prg\_chebyshev\_mod::prg\_get\_chebcoeffs\_fermi\_bs ( integer, intent(in) npts, real(dp), intent(in) kbt, real(dp), intent(inout) ef, real(dp), dimension(:), intent(in) tracesT, integer, intent(in) ncoeffs, real(dp), dimension(:), intent(inout) coeffs, real(dp), intent(in) emin, real(dp), intent(in) emax, real(dp), intent(in) bndfil, integer, intent(in) norb, real(dp), intent(in) tol, logical, intent(in) jon, integer, intent(in) verbose) [private]

Gets the coefficients of the Chebyshev expansion with Ef computation.

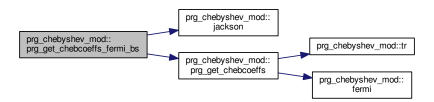
In this case we are applying the bisection method to find the root.

#### **Parameters**

npts	Number of points for the discretization.
kbt	Electronic temperature.
ef	Fermi level.
tracesT	Input traces for matrix polynomials.
ncoeffs	Number of Chebyshev coefficients.
coeffs	Output vector for the Chebyshev coefficients.
emin	lowest boundary for the eigenvalues of H.
emax	highest boundary for the eigenvalues of H.
tol	Tolerance for the bisection method.
verbose	Verbosity level.

Definition at line 620 of file prg\_chebyshev\_mod.F90.

Here is the call graph for this function:



9.2.2.8 subroutine prg\_chebyshev\_mod::prg\_get\_chebcoeffs\_fermi\_nt ( integer, intent(in) npts, real(dp), intent(in) kbt, real(dp), intent(inout) ef, real(dp), dimension(:), intent(in) tracesT, integer, intent(in) ncoeffs, real(dp), dimension(:), intent(inout) coeffs, real(dp), intent(in) emin, real(dp), intent(in) emax, real(dp), intent(in) bndfil, integer, intent(in) norb, real(dp), intent(in) tol, logical, intent(in) jon, integer, intent(in) verbose) [private]

Gets the coefficients of the Chebyshev expansion with Ef computation.

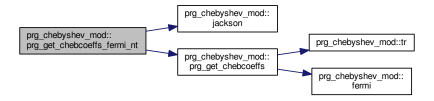
In this case the Newton-Raphson method is applied to find the root.

#### **Parameters**

npst	Number of points for the discretization.
kbt	Electronic temperature.
ef	Fermi level.
tracesT	Input traces for matrix polynomials.
ncoeffs	Number of Chebyshev coefficients.
coeffs	Output vector for the Chebyshev coefficients.
emin	lowest boundary for the eigenvalues of H.
emax	highest boundary for the eigenvalues of H.
bndfil	Band filing factor.
norb	Number of orbitals.
tol	Tolerance for NR method.
verbose	Verbosity level.

Definition at line 697 of file prg\_chebyshev\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



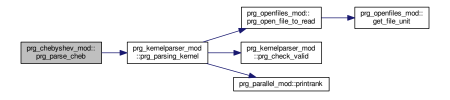
9.2.2.9 subroutine, public prg\_chebyshev\_mod::prg\_parse\_cheb ( type(chebdata\_type), intent(inout) chebdata, character(len=\*) filename )

Chebyshev parser. This module is used to parse all the input variables for the cheb electronic structure solver. Adding a new input keyword to the parser:

- If the variable is real, we have to increase nkey\_re.
- · Add the keyword (character type) in the keyvector re vector.
- Add a default value (real type) in the valvector\_re.
- Define a new variable and pass the value through valvector\_re(num) where num is the position of the new keyword in the vector.

Definition at line 54 of file prg\_chebyshev\_mod.F90.

Here is the call graph for this function:



9.2.2.10 real(dp) function prg\_chebyshev\_mod::tr ( integer, intent(in) r, real(dp), intent(in) x ) [private]

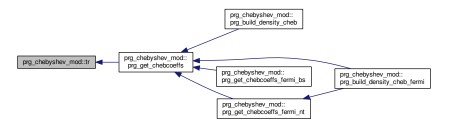
Chebyshev polynomial obtained by recursion.

#### **Parameters**

r	rth polynomial.
X	argument the evaluate the polynomial.

Definition at line 777 of file prg\_chebyshev\_mod.F90.

Here is the caller graph for this function:



## 9.2.3 Variable Documentation

**9.2.3.1** integer, parameter prg\_chebyshev\_mod::dp = kind(1.0d0) [private]

Definition at line 23 of file prg\_chebyshev\_mod.F90.

**9.2.3.2** real(dp), parameter prg\_chebyshev\_mod::pi = 3.14159265358979323846264338327950\_dp [private]

Definition at line 24 of file prg\_chebyshev\_mod.F90.

# 9.3 prg\_densitymatrix\_mod Module Reference

Module to obtain the density matrix by diagonalizing an orthogonalized Hamiltonian.

#### **Functions/Subroutines**

• subroutine, public prg\_build\_density\_t0 (ham\_bml, rho\_bml, threshold, bndfil)

Builds the density matrix from  $H_0$  for zero electronic temperature.  $\rho = C\Theta(\mu I - \epsilon)C^{\dagger}$  Where, C is the matrix eigenvector and  $\epsilon$  is the matrix eigenvalue.  $\Theta()$  is the Heaviside function.

• subroutine, public prg\_build\_density\_t (ham\_bml, rho\_bml, threshold, bndfil, kbt, ef)

Builds the density matrix from  $H_0$  for electronic temperature T.  $\rho = C f(\mu I - \epsilon) C^{\dagger}$  Where, C is the matrix eigenvector and  $\epsilon$  is the matrix eigenvalue. f is the Fermi function.

• subroutine, public prg\_build\_density\_t\_fermi (ham\_bml, rho\_bml, threshold, kbt, ef, verbose)

Builds the density matrix from  $H_0$  for electronic temperature T.  $\rho = Cf(\mu I - \epsilon)C^{\dagger}$  Where, C is the matrix eigenvector and  $\epsilon$  is the matrix eigenvalue. f is the Fermi function. In this routine the Fermi level is passed as an argument.

subroutine, public prg\_build\_atomic\_density (rhoat\_bml, numel, hindex, spindex, norb, bml\_type)

Builds the atomic density matrix.  $ho_{ii} = mathcal Z_{ii}$  Where,  $mathcal Z_{ii}$  is the number of electrons for orbital i.

• subroutine, public prg\_get\_flevel (eigenvalues, kbt, bndfil, tol, Ef)

Routine to compute the Fermi level given a set of eigenvalues and a temperature. It applies the Bisection method over the function:  $g(\mu) = \sum_k 2f(\epsilon_k - \mu) - N = 0$  Where  $f(\epsilon_k - \mu) = \frac{1}{1 + \exp{(\epsilon_k - \mu)/(k_h T)}}$ .

subroutine, public prg\_get\_eigenvalues (ham\_bml, eigenvalues, verbose)

Gets the eigenvalues of the Orthogonalized Hamiltonian.

• subroutine, public prg\_check\_idempotency (mat\_bml, threshold, idempotency)

To check the idempotency error of a matrix. This is calculated as the Frobenius norm of  $(A - A^2)$ .

real(dp) function fermi (e, ef, kbt)

Gives the Fermi distribution value for energy e.

#### **Variables**

integer, parameter dp = kind(1.0d0)

## 9.3.1 Detailed Description

Module to obtain the density matrix by diagonalizing an orthogonalized Hamiltonian.

#### 9.3.2 Function/Subroutine Documentation

9.3.2.1 real(dp) function prg\_densitymatrix\_mod::fermi ( real(dp), intent(in) e, real(dp), intent(in) ef, real(dp), intent(in) kbt )

[private]

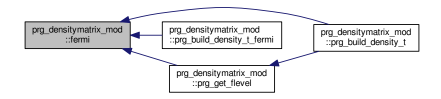
Gives the Fermi distribution value for energy e.

#### **Parameters**

е	Energy.
ef	Fermi energy.

Definition at line 413 of file prg\_densitymatrix\_mod.F90.

Here is the caller graph for this function:



9.3.2.2 subroutine, public prg\_densitymatrix\_mod::prg\_build\_atomic\_density ( type(bml\_matrix\_t), intent(inout) rhoat\_bml, real(dp), dimension(:), intent(in) numel, integer, dimension(:,:), intent(in) hindex, integer, dimension(:), intent(in) spindex, integer, intent(in) norb, character(len=\*), intent(in) bml\_type )

Builds the atomic density matrix.  $\rho_{ii} = mathcal Z_{ii}$  Where,  $mathcal Z_{ii}$  is the number of electrons for orbital i.

#### **Parameters**

rhoat	Output atomic diagonal density matrix,
hindex	Start and end index for every atom in the system.
numel	Number of electrons per specie. It runs over the specie index.
spindex	Specie index.
norbs	Number of orbitals.

Definition at line 215 of file prg\_densitymatrix\_mod.F90.

9.3.2.3 subroutine, public prg\_densitymatrix\_mod::prg\_build\_density\_t ( type(bml\_matrix\_t), intent(in) ham\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(8), intent(in) threshold, real(8), intent(in) bndfil, real(8), intent(inout) ef )

Builds the density matrix from  $H_0$  for electronic temperature T.  $\rho = C f(\mu I - \epsilon) C^{\dagger}$  Where, C is the matrix eigenvector and  $\epsilon$  is the matrix eigenvalue. f is the Fermi function.

## **Parameters**

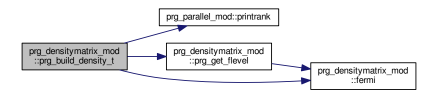
ham_bml	Input Orthogonalized Hamiltonian matrix.
rho_bml	Output density matrix,
threshold	Threshold for sparse matrix algebra.
bndfil	Filing factor.

#### Warning

This does not solve the generalized eigenvalue problem. The Hamiltonian that comes in has to be preorthogonalized.

Definition at line 94 of file prg\_densitymatrix\_mod.F90.

Here is the call graph for this function:



9.3.2.4 subroutine, public prg\_densitymatrix\_mod::prg\_build\_density\_t0 ( type(bml\_matrix\_t), intent(in) ham\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(8), intent(in) threshold, real(8), intent(in) bndfil )

Builds the density matrix from  $H_0$  for zero electronic temperature.  $\rho = C\Theta(\mu I - \epsilon)C^{\dagger}$  Where, C is the matrix eigenvector and  $\epsilon$  is the matrix eigenvalue.  $\Theta()$  is the Heaviside function.

#### **Parameters**

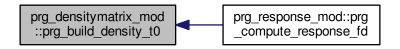
ham_bml	Input Orthogonalized Hamiltonian matrix.
rho_bml	Output density matrix,
threshold	Threshold for sparse matrix algebra.
bndfil	Filing factor.

# Warning

This does not solve the generalized eigenvalue problem. The Hamiltonian that comes in has to be preprg\_ orthogonalized.

Definition at line 33 of file prg\_densitymatrix\_mod.F90.

Here is the caller graph for this function:



9.3.2.5 subroutine, public prg\_densitymatrix\_mod::prg\_build\_density\_t\_fermi ( type(bml\_matrix\_t), intent(in) ham\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(8), intent(in) threshold, real(8), intent(in) kbt, real(8), intent(in) ef, integer, intent(in), optional verbose )

Builds the density matrix from  $H_0$  for electronic temperature T.  $\rho=Cf(\mu I-\epsilon)C^\dagger$  Where, C is the matrix eigenvector and  $\epsilon$  is the matrix eigenvalue. f is the Fermi function. In this routine the Fermi level is passed as an argument.

#### **Parameters**

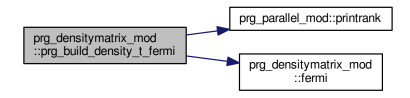
ham_bml	Input Orthogonalized Hamiltonian matrix.
rho_bml	Output density matrix,
threshold	Threshold for sparse matrix algebra.

## Warning

This does not solve the generalized eigenvalue problem. The Hamiltonian that comes in has to be preorthogonalized.

Definition at line 156 of file prg\_densitymatrix\_mod.F90.

Here is the call graph for this function:



9.3.2.6 subroutine, public prg\_densitymatrix\_mod::prg\_check\_idempotency ( type(bml\_matrix\_t), intent(in) *mat\_bml*, real(dp), intent(in) *threshold*, real(dp), intent(out) *idempotency* )

To check the idempotency error of a matrix. This is calculated as the Frobenius norm of  $(A-A^2)$ .

#### **Parameters**

mat_bml	Some bml matrix
idempotency	(Output value of the idempotency error)

Definition at line 389 of file prg\_densitymatrix\_mod.F90.

9.3.2.7 subroutine, public prg\_densitymatrix\_mod::prg\_get\_eigenvalues ( type(bml\_matrix\_t), intent(in) ham\_bml, real(dp), dimension(:), intent(inout), allocatable eigenvalues, integer, intent(in) verbose )

Gets the eigenvalues of the Orthogonalized Hamiltonian.

#### **Parameters**

ham_bml	Input Orthogonalized Hamiltonian matrix.
eigenvalues	Output eigenvalues of the system.
verbose	Verbosity level.

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Definition at line 344 of file prg\_densitymatrix\_mod.F90.

9.3.2.8 subroutine, public prg\_densitymatrix\_mod::prg\_get\_flevel ( real(dp), dimension(:), intent(in) eigenvalues, real(dp), intent(in) kbt, real(dp), intent(in) bndfil, real(dp) tol, real(dp), intent(inout) Ef )

Routine to compute the Fermi level given a set of eigenvalues and a temperature. It applies the Bisection method over the function:  $g(\mu) = \sum_k 2f(\epsilon_k - \mu) - N = 0$  Where  $f(\epsilon_k - \mu) = \frac{1}{1 + \exp{(\epsilon_k - \mu)/(k_b T)}}$ .

#### **Parameters**

eigenvalues	Eigenvalues of the system ( $\{\epsilon_k\}$ ).
kbt	Temperature times the Boltzmans's constant ( $k_bT$ ).
bndfil	Filing factor ( $N_{el}/(2*N_{orbs})$ ).
tol	Tolerance for the bisection method.
Ef	Fermi level ( $\mu$ ).

Definition at line 280 of file prg\_densitymatrix\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



# 9.3.3 Variable Documentation

**9.3.3.1** integer, parameter prg\_densitymatrix\_mod::dp = kind(1.0d0) [private]

Definition at line 14 of file prg\_densitymatrix\_mod.F90.

# 9.4 prg\_dos\_mod Module Reference

A module to compute the Density of state (DOS) and IDOS.

#### **Functions/Subroutines**

- subroutine, public prg\_write\_tdos (eigenvals, gamma, npts, emin, emax, filename) Writes the total DOS into a file.  $DOS(\epsilon) = \sum_k L(\epsilon \epsilon_k)$  Where  $\int_{-\infty}^{\infty} DOS(\epsilon) = Nstates$ .
- real(dp) function lorentz (energy, eigenvals, loads, Gamma)
   Lorentzian Function.

#### **Variables**

• integer, parameter dp = kind(1.0d0)

# 9.4.1 Detailed Description

A module to compute the Density of state (DOS) and IDOS.

This module will be used to compute DOS and IDOS.

Todo Add LDOS.

# 9.4.2 Function/Subroutine Documentation

9.4.2.1 real(dp) function prg\_dos\_mod::lorentz ( real(dp), intent(in) energy, real(dp), dimension(:), intent(in) eigenvals, real(dp), dimension(:), intent(in) loads, real(dp), intent(in) Gamma ) [private]

Lorentzian Function.

Computes: 
$$L(\epsilon) = \sum_k \frac{\omega(k)\Gamma}{2\pi} \frac{1}{(\epsilon - \epsilon_k)^2 + (\Gamma/2)^2}$$

# Parameters

energy	Energy point.
eigenvals	Eigenvalues of the system.
Gamma	Lorentz function broadening.

Definition at line 78 of file prg\_dos\_mod.F90.

Here is the caller graph for this function:



9.4.2.2 subroutine, public prg\_dos\_mod::prg\_write\_tdos ( real(dp), dimension(:), intent(in) eigenvals, real(dp), intent(in) gamma, integer, intent(in) npts, real(dp), intent(in) emin, real(dp), intent(in) emax, character(len=\*), intent(in) filename

Writes the total DOS into a file.  $DOS(\epsilon) = \sum_k L(\epsilon - \epsilon_k)$  Where  $\int_{-\infty}^{\infty} DOS(\epsilon) = Nstates$ .

Note

DOS is NOT shifted respect to Ef.

#### **Parameters**

eigenvals	Eigenvalues of the system. gamma Lorentzian width.
npts	Number of energy points.
emin	Minimum energy value.
emax	Maximum energy value.
filename	Filename to write the DOS.

Definition at line 36 of file prg\_dos\_mod.F90.

Here is the call graph for this function:



## 9.4.3 Variable Documentation

**9.4.3.1** integer, parameter prg\_dos\_mod::dp = kind(1.0d0) [private]

Definition at line 17 of file prg\_dos\_mod.F90.

# 9.5 prg\_extras\_mod Module Reference

Extra routines.

# **Data Types**

- interface prg\_memory\_consumption
- · interface to string

## **Functions/Subroutines**

```
• character(len=:) function, allocatable to_string_integer (i)
```

Convert integer to string.

• character(len=:) function, allocatable to\_string\_long\_long (i)

Convert integer to string.

• character(len=:) function, allocatable to\_string\_double (x)

Convert double to string.

• subroutine, public prg\_print\_matrix (matname, amat, i1, i2, j1, j2)

To write a dense matrix to screen.

• real(dp) function, public mls ()

To get the actual time in milliseconds.

• subroutine, public prg\_delta (x, s, nn, dta)

Delta function  $||X^{\wedge}tSX - I||$ .

• subroutine, public prg\_get\_mem (procname, tag)

Get proc memory.

• subroutine prg\_twonorm (a, nn, norm2)

Gets the norm2 of a square matrix.

#### **Variables**

• integer, parameter dp = kind(1.0d0)

## 9.5.1 Detailed Description

Extra routines.

A module to add any extra routine considered necessary but which is NOT essential for any other PROGRESS routine.

## 9.5.2 Function/Subroutine Documentation

9.5.2.1 real(dp) function, public prg\_extras\_mod::mls ( )

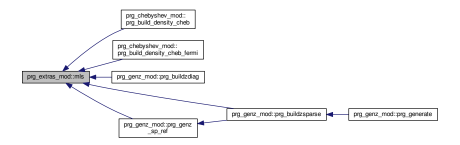
To get the actual time in milliseconds.

#### **Parameters**

Is Output value with the machine time in milliseconds.
--

Definition at line 139 of file prg\_extras\_mod.F90.

Here is the caller graph for this function:



9.5.2.2 subroutine, public prg\_extras\_mod::prg\_delta ( real(dp), dimension(nn,nn) x, real(dp), dimension(nn,nn) s, integer nn, real(dp) dta )

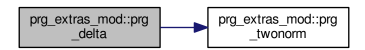
Delta function  $||X^{\hat{}}tSX - I||$ .

### **Parameters**

Х	input matrix.
s	overlap matrix.
dta	Delta output value.

Definition at line 155 of file prg\_extras\_mod.F90.

Here is the call graph for this function:



9.5.2.3 subroutine, public prg\_extras\_mod::prg\_get\_mem ( character(\*), intent(in) procname, character(\*), intent(in) tag )

Get proc memory.

#### **Parameters**

procname	Process name to get the mem usage.
tag	Tag to pprint the processor mem usage.

Definition at line 191 of file prg\_extras\_mod.F90.

9.5.2.4 subroutine, public prg\_extras\_mod::prg\_print\_matrix ( character(len=\*) matname, real(dp), dimension(:,:), intent(in) amat, integer, intent(in) i1, integer, intent(in) i2, integer, intent(in) j1, integer, intent(in) j2)

To write a dense matrix to screen.

#### **Parameters**

matname	Matrix name.
amat	Matrix to be printed.
i1	Print from row i1.
i2	Print up to from row i2.
j1	Print from column j1.
j2	Print up to column j2.

Definition at line 100 of file prg\_extras\_mod.F90.

9.5.2.5 subroutine prg\_extras\_mod::prg\_twonorm ( real(dp), dimension(nn,nn) a, integer nn, real(dp) norm2 ) [private]

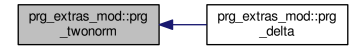
Gets the norm2 of a square matrix.

## **Parameters**

а	Square matrix.
nn	Matrix size.
norm2	Two-norm of matrix a.

Definition at line 215 of file prg\_extras\_mod.F90.

Here is the caller graph for this function:



9.5.2.6 character(len=:) function, allocatable prg\_extras\_mod::to\_string\_double ( double precision, intent(in) x ) [private] Convert double to string. **Parameters** The double Returns The string Definition at line 80 of file prg\_extras\_mod.F90. 9.5.2.7 character(len=:) function, allocatable prg\_extras\_mod::to\_string\_integer( integer, intent(in) i) [private] Convert integer to string. **Parameters** The integer Returns The string Definition at line 46 of file prg\_extras\_mod.F90. 9.5.2.8 character(len=:) function, allocatable prg\_extras\_mod::to\_string\_long\_long ( integer(kind=c\_long\_long), intent(in) i ) [private] Convert integer to string. **Parameters** The integer Returns The string Definition at line 62 of file prg\_extras\_mod.F90. 9.5.3 Variable Documentation 9.5.3.1 integer, parameter prg\_extras\_mod::dp = kind(1.0d0) [private] Definition at line 31 of file prg\_extras\_mod.F90.

# 9.6 prg\_genz\_mod Module Reference

To produce a matrix Z which is needed to orthogonalize H.

# **Data Types**

· type genzspdata

contains the data for the genZ driver.

type genzspinp

Input for the genz driver. This type controlls all the variables that are needed by genz.

#### **Functions/Subroutines**

• subroutine, public prg parse zsp (input, filename)

The parser for genz solver.

subroutine prg\_init (self, input)

Initializes the genz input variables.

Allocates the matrices for the XI integration of Z.

subroutine, public prg\_init\_zspmat (igenz, zk1\_bml, zk2\_bml, zk3\_bml, zk4\_bml, zk5\_bml, zk6\_bml, norb, bml\_type)

Initiates the matrices for the XI integration of Z.

• subroutine <a href="mailto:prg\_generate">prg\_generate</a> (self, over\_bml, zmat\_bml, igenz, mdim, bml\_type, zk1\_bml, zk2\_bml, zk3\_bml, zk4\_bml, zk5\_bml, zk6\_bml)

Generates the Z matrix.

• subroutine, public prg\_buildzdiag (smat\_bml, zmat\_bml, threshold, mdimin, bml\_type, verbose)

Usual subroutine involving diagonalization.  $Z=U\sqrt{s}U^\dagger$ , where U = eigenvectors and s = eigenvalues. The purpose of this subroutine is to have an exact way of computing z for comparing with the sparse approach.

subroutine, public prg\_buildzsparse (smat\_bml, zmat\_bml, igenz, mdim, bml\_type, zk1\_bml, zk2\_bml, zk3
 \_bml, zk4\_bml, zk6\_bml, rirst, nrefi, nreff, thresholdi, thresholdi, integration, verbose)

Inverse factorization using Niklasson's algorithm.

• subroutine, public prg\_genz\_sp\_initialz0 (smat\_bml, zmat\_bml, norb, mdim, bml\_type\_f, threshold)

Initial estimation of Z.

Initial estimation of Z.

• subroutine, public prg\_genz\_sp\_initial\_zmat (smat\_bml, zmat\_bml, norb, mdim, bml\_type\_f, threshold)

• subroutine <a href="mailto:prg\_genz\_sp\_int">prg\_genz\_sp\_int</a> (zmat\_bml, zk1\_bml, zk2\_bml, zk3\_bml, zk4\_bml, zk5\_bml, zk6\_bml, igenz, norb, bml\_type, threshold)

Inverse factorization using Niklasson's algorithm.

# **Variables**

integer, parameter dp = kind(1.0d0)

# 9.6.1 Detailed Description

To produce a matrix Z which is needed to orthogonalize H.

 $H_{orth} = Z^{\dagger}HZ$  See Negre 2016 [2]

# 9.6.2 Function/Subroutine Documentation

9.6.2.1 subroutine prg\_genz\_mod::prg\_allocatezspmat ( class(genzspdata), intent(in) self, type(bml\_matrix\_t) zk1\_bml, type(bml\_matrix\_t) zk2\_bml, type(bml\_matrix\_t) zk3\_bml, type(bml\_matrix\_t) zk4\_bml, type(bml\_matrix\_t) zk5\_bml, type(bml\_matrix\_t) zk6\_bml, integer norb, character(20) bml\_type ) [private]

Allocates the matrices for the XI integration of Z.

#### **Parameters**

self	input zsp variables
zk1_bml-zk6_bml	history record of the previous Z matrices.
norb	number of orbitals.
bml_type	the bml format we are passing.

Definition at line 170 of file prg\_genz\_mod.F90.

9.6.2.2 subroutine, public prg\_genz\_mod::prg\_buildzdiag ( type(bml\_matrix\_t), intent(inout) smat\_bml, type(bml\_matrix\_t) zmat\_bml, real(dp) threshold, integer, intent(in) mdimin, character(len=\*) bml\_type, integer, intent(in), optional verbose )

Usual subroutine involving diagonalization.  $Z=U\sqrt{s}U^{\dagger}$ , where U= eigenvectors and s= eigenvalues. The purpose of this subroutine is to have an exact way of computing z for comparing with the sparse approach.

#### **Parameters**

smat_bml	Overlap matrix in bml format.
zmat_bml	Congruence transform in bml format.
threshold	Threshold value to use, in this case, only in the backtransformation to ellpack format.
mdim	Maximun nonzero to use, in this case, only in the backtransformation to ellpack format.
bml_type	the bml type we are passing.

Definition at line 274 of file prg\_genz\_mod.F90.

Here is the call graph for this function:



9.6.2.3 subroutine, public prg\_genz\_mod::prg\_buildzsparse (\_type(bml\_matrix\_t) smat\_bml, \_type(bml\_matrix\_t) zmat\_bml, \_integer igenz, integer mdim, character(20) bml\_type, type(bml\_matrix\_t) zk1\_bml, type(bml\_matrix\_t) zk2\_bml, type(bml\_matrix\_t) zk3\_bml, type(bml\_matrix\_t) zk4\_bml, type(bml\_matrix\_t) zk5\_bml, type(bml\_matrix\_t) zk6\_bml, integer nfirst, integer nrefi, integer nrefi, real(dp) thresholdi, real(dp) thresholdi, logical integration, integer verbose)

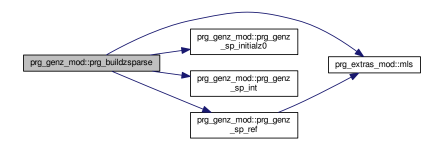
Inverse factorization using Niklasson's algorithm.

#### **Parameters**

smat_bml	overlap matrix
zmat_bml	congruence transform to be updated or computed. (bml format)
igenz	counter to keep track of the calls to this subroutine.
mdim	dimension of the maxnonzero per row.
zk1_bml-zk6_bml	history of the past congruence transforms.
nfirst	first pre septs with nrefi and thresholdi.
nrefi	number of refinement iterations for the firsts "nfirst" steps.
nreff	number of refinement iterations for the rest of the steps.
integration	if we want to apply xl integration scheme for z (default is always .true.)
verbose	to print extra information.

Definition at line 413 of file prg\_genz\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



9.6.2.4 subroutine prg\_genz\_mod::prg\_generate ( class(genzspdata), intent(in) self, type(bml\_matrix\_t), intent(in) over\_bml, type(bml\_matrix\_t), intent(inout) zmat\_bml, integer igenz, integer mdim, character(20) bml\_type, type(bml\_matrix\_t), intent(inout) zk1\_bml, type(bml\_matrix\_t), intent(inout) zk2\_bml, type(bml\_matrix\_t), intent(inout) zk3\_bml, type(bml\_matrix\_t), intent(inout) zk4\_bml, type(bml\_matrix\_t), intent(inout) zk6\_bml, type(bml\_matrix\_t), intent(inout) zk6\_bml) [private]

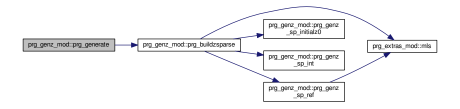
Generates the Z matrix.

#### **Parameters**

over_bml	Overlap matrix.
zmat_bml	Congruence transform to be computed. (bml format)
igenz	Counter to keep track of the calls to this subroutine.
mdim	dimension of the maxnonzero per row.
zk1_bml-zk6_bml	history of the past congruence transforms.

Definition at line 237 of file prg\_genz\_mod.F90.

Here is the call graph for this function:



9.6.2.5 subroutine, public prg\_genz\_mod::prg\_genz\_sp\_initial\_zmat ( type(bml\_matrix\_t), intent(in) smat\_bml, type(bml\_matrix\_t) zmat\_bml, integer norb, integer mdim, character(20) bml\_type\_f, real(dp), intent(in) threshold )

Initial estimation of Z.

#### Note

Most of the operations are done in pure dense format. The purpose of this subroutine is to have an exact way of computing z for comparing with the sparse approach.

#### **Parameters**

smat_bml	Overlap matrix in bml format.
zmat_bml	Congruence transform in bml format.
norb	Congruence transform in bml format.
mdim	Congruence transform in bml format.
bml_←	The bml final type of zmat_bml.
type_f	
threshold	Threshold value to use, in this case, only in the backtransformation to ellpack format.

Definition at line 611 of file prg\_genz\_mod.F90.

9.6.2.6 subroutine, public prg\_genz\_mod::prg\_genz\_sp\_initialz0 ( type(bml\_matrix\_t), intent(in) smat\_bml, type(bml\_matrix\_t) zmat\_bml, integer norb, integer mdim, character(20) bml\_type\_f, real(dp) threshold )

Initial estimation of Z.

#### Note

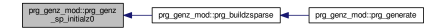
Most of the operations are done in pure dense format. The purpose of this subroutine is to have an exact way of computing z for comparing with the sparse approach.

#### **Parameters**

smat_bml	Overlap matrix in bml format.
zmat_bml	Congruence transform in bml format.
norb	Congruence transform in bml format.
mdim	Congruence transform in bml format.
bml_←	The bml final type of zmat_bml.
type_f	
threshold	Threshold value to use, in this case, only in the backtransformation to ellpack format.

Definition at line 477 of file prg\_genz\_mod.F90.

Here is the caller graph for this function:



9.6.2.7 subroutine prg\_genz\_mod::prg\_genz\_sp\_int ( type(bml\_matrix\_t) zmat\_bml, type(bml\_matrix\_t) zk1\_bml, type(bml\_matrix\_t) zk2\_bml, type(bml\_matrix\_t) zk3\_bml, type(bml\_matrix\_t) zk4\_bml, type(bml\_matrix\_t) zk5\_bml, type(bml\_matrix\_t) zk6\_bml, integer igenz, integer norb, character(20) bml\_type, real(dp) threshold ) [private]

Inverse factorization using Niklasson's algorithm.

# **Parameters**

smat_bml	overlap matrix
zmat_bml	congruence transform to be updated or computed. (bml format)
mdim	dimension of the maxnonzero per row.
zk1_bml-zk6_bml	history of the past congruence transforms.
igenz	counter to keep track of the calls to this subroutine.
norb	Congruence transform in bml format.
bml_type_f	The bml final type of zmat_bml.
threshold	Threshold value to use.

Definition at line 759 of file prg\_genz\_mod.F90.

Here is the caller graph for this function:



9.6.2.8 subroutine, public prg\_genz\_mod::prg\_genz\_sp\_ref ( type(bml\_matrix\_t), intent(in) smat\_bml, type(bml\_matrix\_t), intent(inout) zmat\_bml, integer, intent(in) nref, integer, intent(inout) norb, character(20), intent(in) bml\_type, real(dp), intent(in) threshold )

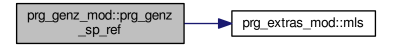
Iterative refinement.

### **Parameters**

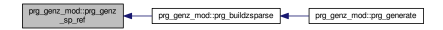
smat_bml	overlap matrix
zmat_bml	congruence transform to be updated or computed. (bml format)
nref	Number of refinement iterations.
bml_←	The bml final type of zmat_bml.
type_f	
threshold	Threshold value to use.
verbose	to print extra information.

Definition at line 833 of file prg\_genz\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



9.6.2.9 subroutine prg\_genz\_mod::prg\_init ( class(genzspdata), intent(out) self, type(genzspinp), intent(in) input ) [private]

Initializes the genz input variables.

#### **Parameters**

self	basic input parameters.
input	basic input parameters from the parser.

Definition at line 149 of file prg\_genz\_mod.F90.

9.6.2.10 subroutine, public prg\_genz\_mod::prg\_init\_zspmat ( integer *igenz*, type(bml\_matrix\_t) *zk1\_bml*, type(bml\_matrix\_t) *zk2\_bml*, type(bml\_matrix\_t) *zk3\_bml*, type(bml\_matrix\_t) *zk4\_bml*, type(bml\_matrix\_t) *zk5\_bml*, type(bml\_matrix\_t) *zk6\_bml*, integer *norb*, character(20) *bml\_type* )

Initiates the matrices for the XI integration of Z.

# **Parameters**

self	input zsp variables
zk1_bml-zk6_bml	history record of the previous Z matrices.
norb	number of orbitals.
bml_type	the bml format we are passing.

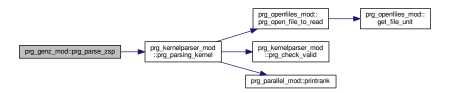
Definition at line 202 of file prg\_genz\_mod.F90.

9.6.2.11 subroutine, public prg\_genz\_mod::prg\_parse\_zsp ( type(genzspinp), intent(inout) input, character(len=\*) filename )

The parser for genz solver.

Definition at line 86 of file prg\_genz\_mod.F90.

Here is the call graph for this function:



#### 9.6.3 Variable Documentation

9.6.3.1 integer, parameter prg\_genz\_mod::dp = kind(1.0d0) [private]

Definition at line 18 of file prg\_genz\_mod.F90.

# 9.7 prg\_graph\_mod Module Reference

The graph module.

# **Data Types**

- · type graph\_partitioning\_t
  - Trace per iteration.
- type subgraph\_t

Subgraph type.

### **Functions/Subroutines**

- subroutine, public prg\_initsubgraph (sg, pnum, hsize)
  - Initialize subgraph.
- subroutine, public prg\_destroysubgraph (sg)

Destroy subgraph.

• subroutine, public prg\_initgraphpartitioning (gp, pname, np, nnodes, nnodes2)

Initialize graph partitioning.

- subroutine, public prg\_destroygraphpartitioning (gp)
  - Destroy graph partitioning.
- subroutine, public prg\_printgraphpartitioning (gp)

Print graph partitioning structure data.

• subroutine, public prg\_equalpartition (gp, nodesPerPart, nnodes)

Create equal graph partitions, based on number of rows/orbitals.

• subroutine, public prg\_equalgrouppartition (gp, hindex, ngroup, nodesPerPart, nnodes)

Create equal group graph partitions, based on number of atoms/groups.

• subroutine, public <a href="mailto:prg\_filepartition">prg\_filepartition</a> (gp, partFile)

Read graph partitions from a file, based on number of rows/orbitals.

• subroutine <a href="mailto:prg\_readpart">prg\_readpart</a> (gp, partFile)

Read parts (core) from part file.

• subroutine, public prg\_fnormgraph (gp)

Accumulate trace norm across all subgraphs.

# **Variables**

• integer, parameter dp = kind(1.0d0)

# 9.7.1 Detailed Description

The graph module.

### 9.7.2 Function/Subroutine Documentation

9.7.2.1 subroutine, public prg\_graph\_mod::prg\_destroygraphpartitioning ( type (graph\_partitioning\_t), intent(inout) gp )

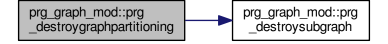
Destroy graph partitioning.

#### **Parameters**

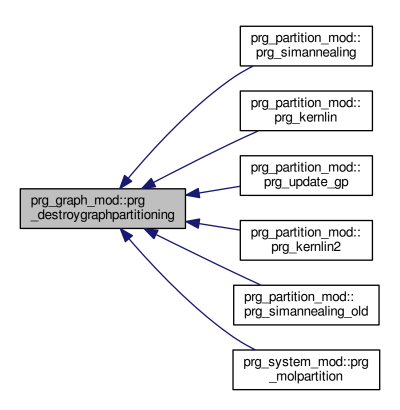
sg Subgraph

Definition at line 263 of file prg\_graph\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



9.7.2.2 subroutine, public prg\_graph\_mod::prg\_destroysubgraph ( type (subgraph\_t), intent(inout) sg )

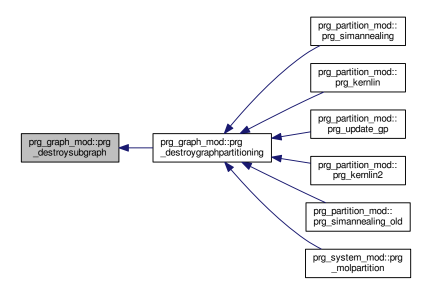
Destroy subgraph.

**Parameters** 

sg Subgraph

Definition at line 159 of file prg\_graph\_mod.F90.

Here is the caller graph for this function:



9.7.2.3 subroutine, public prg\_graph\_mod::prg\_equalgrouppartition ( type (graph\_partitioning\_t), intent(inout) *gp*, integer, dimension(2,ngroup), intent(in) *hindex*, integer, intent(in) *ngroup*, integer, intent(in) *nnodes*)

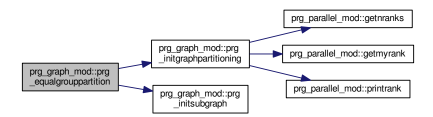
Create equal group graph partitions, based on number of atoms/groups.

# **Parameters**

gp	Graph partitioning
hindex	Node indeces that represent ranges of atoms/groups
ngroup	Number of group nodes
nodesPerPart	Number of core nodes per partition
nnodes	Total nodes in Hamiltonian matrix

Definition at line 402 of file prg\_graph\_mod.F90.

Here is the call graph for this function:



9.7.2.4 subroutine, public prg\_graph\_mod::prg\_equalpartition ( type (graph\_partitioning\_t), intent(inout) *gp*, integer, intent(in) *nodesPerPart*, integer, intent(in) *nnodes* )

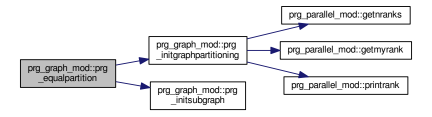
Create equal graph partitions, based on number of rows/orbitals.

#### **Parameters**

gp	Graph partitioning	
nodesPerPart	Number of core nodes per partition	
nnodes	Total nodes in Hamiltonian matrix	

Definition at line 355 of file prg\_graph\_mod.F90.

Here is the call graph for this function:



9.7.2.5 subroutine, public prg\_graph\_mod::prg\_filepartition ( type (graph\_partitioning\_t), intent(inout) *gp*, character(len=\*), intent(in) *partFile* )

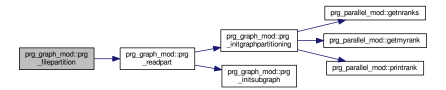
Read graph partitions from a file, based on number of rows/orbitals.

#### **Parameters**

partFile	File containing core nodes for each partition
gp	Graph partitioning

Definition at line 463 of file prg\_graph\_mod.F90.

Here is the call graph for this function:



9.7.2.6 subroutine, public prg\_graph\_mod::prg\_fnormgraph ( type(graph\_partitioning\_t), intent(inout) gp )

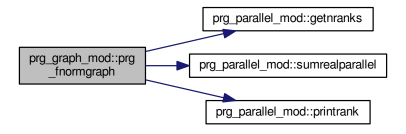
Accumulate trace norm across all subgraphs.

#### **Parameters**

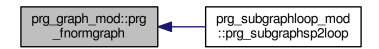
gp	Graph partitioning
----	--------------------

Definition at line 516 of file prg\_graph\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



9.7.2.7 subroutine, public prg\_graph\_mod::prg\_initgraphpartitioning ( type (graph\_partitioning\_t), intent(inout) *gp,* character(len=\*), intent(in) *pname,* integer, intent(in) *np,* integer, intent(in) *nnodes,* integer, intent(in) *nnodes2*)

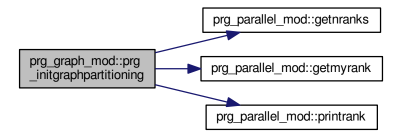
Initialize graph partitioning.

# **Parameters**

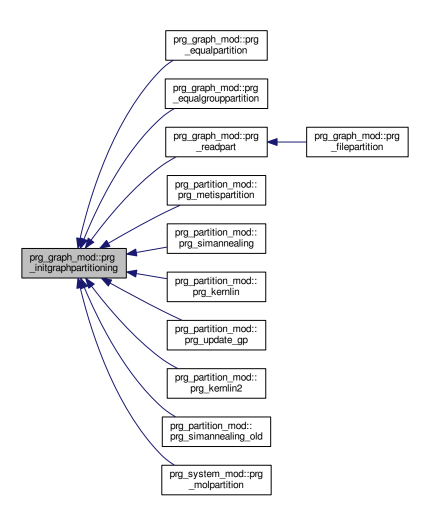
gp	Graph partitioning
pname	Partitioning name
np	Number of partitions
nnodes	Number of groups/nodes
nnodes2	Number of nodes

Definition at line 175 of file prg\_graph\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



9.7.2.8 subroutine, public prg\_graph\_mod::prg\_initsubgraph ( type (subgraph\_t), intent(inout) sg, integer, intent(in) pnum, integer, intent(in) hsize )

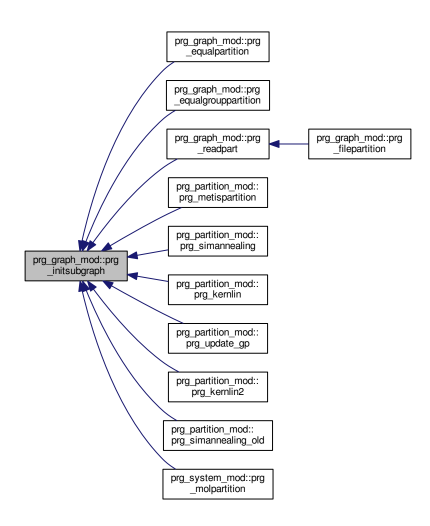
Initialize subgraph.

# **Parameters**

sg	Subgraph
pnum	Part number
hsize	Size of full matrix

Definition at line 143 of file prg\_graph\_mod.F90.

Here is the caller graph for this function:



9.7.2.9 subroutine, public prg\_graph\_mod::prg\_printgraphpartitioning ( type (graph\_partitioning\_t), intent(in) gp )

Print graph partitioning structure data.

### **Parameters**

gp Graph partitioning

Definition at line 292 of file prg\_graph\_mod.F90.

9.7.2.10 subroutine prg\_graph\_mod::prg\_readpart ( type (graph\_partitioning\_t), intent(inout) *gp,* character(len=\*), intent(in) *partFile* ) [private]

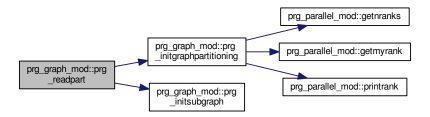
Read parts (core) from part file.

#### **Parameters**

gp	Graph partitioning
partFile	Partition file

Definition at line 475 of file prg\_graph\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



# 9.7.3 Variable Documentation

9.7.3.1 integer, parameter prg\_graph\_mod::dp = kind(1.0d0) [private]

Definition at line 16 of file prg\_graph\_mod.F90.

# 9.8 prg\_graphsp2parser\_mod Module Reference

Graph partitioning SP2 parser.

# **Data Types**

type gsp2data\_type

General SP2 solver type.

# **Functions/Subroutines**

subroutine, public prg\_parse\_gsp2 (gsp2data, filename)
 The parser for SP2 solver.

#### **Variables**

integer, parameter dp = kind(1.0d0)

# 9.8.1 Detailed Description

Graph partitioning SP2 parser.

This module is used to parse all the neccesary input variables for graph-based SP2 electronic structure solver. Adding a new input keyword to the parser:

- If the variable is real, we have to increase nkey\_re.
- · Add the keyword (character type) in the keyvector re vector.
- · Add a default value (real type) in the valvector re.
- Define a new variable and pass the value through valvector\_re(num) where num is the position of the new keyword in the vector.

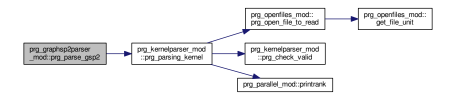
### 9.8.2 Function/Subroutine Documentation

9.8.2.1 subroutine, public prg\_graphsp2parser\_mod::prg\_parse\_gsp2 ( type(gsp2data\_type), intent(inout) gsp2data, character(len=\*) filename )

The parser for SP2 solver.

Definition at line 62 of file prg\_graphsp2parser\_mod.F90.

Here is the call graph for this function:



#### 9.8.3 Variable Documentation

**9.8.3.1** integer, parameter prg\_graphsp2parser\_mod::dp = kind(1.0d0) [private]

Definition at line 22 of file prg\_graphsp2parser\_mod.F90.

# 9.9 prg\_homolumo\_mod Module Reference

The homolumo module.

# **Functions/Subroutines**

- subroutine, public prg\_homolumogap (vv, imax, pp, mineval, maxeval, ehomo, elumo, egap, verbose)
- subroutine, public prg\_sp2sequence (pp, imax, mineval, maxeval, ehomo, elumo, errlimit, verbose)

#### **Variables**

• integer, parameter dp = kind(1.0d0)

# 9.9.1 Detailed Description

The homolumo module.

#### 9.9.2 Function/Subroutine Documentation

9.9.2.1 subroutine, public prg\_homolumo\_mod::prg\_homolumogap ( real(dp), dimension(:), intent(in) vv, integer, intent(in) imax, integer, dimension(:), intent(in) pp, real(dp), intent(in) mineval, real(dp), intent(in) maxeval, real(dp), intent(inout) ehomo, real(dp), intent(inout) elumo, real(dp), intent(inout) egap, integer, intent(in), optional verbose)

Definition at line 24 of file prg homolumo mod.F90.

9.9.2.2 subroutine, public prg\_homolumo\_mod::prg\_sp2sequence ( integer, dimension(:), intent(inout) pp, integer, intent(inout) imax, real(dp), intent(in) mineval, real(dp), intent(in) maxeval, real(dp), intent(in) ehomo, real(dp), intent(in) elumo, real(dp), intent(in) errlimit, integer, intent(in), optional verbose )

Definition at line 94 of file prg\_homolumo\_mod.F90.

#### 9.9.3 Variable Documentation

9.9.3.1 integer, parameter prg\_homolumo\_mod::dp = kind(1.0d0) [private]

Definition at line 14 of file prg\_homolumo\_mod.F90.

# 9.10 prg\_implicit\_fermi\_mod Module Reference

# **Functions/Subroutines**

• subroutine, public prg\_implicit\_fermi (h\_bml, xi0\_bml, p\_bml, nsteps, nocc, mu, beta, osteps, occErrLimit, threshold)

Recursive Implicit Fermi Dirac.

# Variables

• integer, parameter dp = kind(1.0d0)

#### 9.10.1 Function/Subroutine Documentation

9.10.1.1 subroutine, public prg\_implicit\_fermi\_mod::prg\_implicit\_fermi ( type(bml\_matrix\_t), intent(in) h\_bml, type(bml\_matrix\_t), intent(inout) xi0\_bml, type(bml\_matrix\_t), intent(inout) p\_bml, integer, intent(in) nsteps, real(dp), intent(in) nocc, real(dp), intent(inout) mu, real(dp), intent(in) beta, integer, intent(in) osteps, real(dp), intent(in) occErrLimit, real(dp), intent(in) threshold)

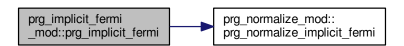
Recursive Implicit Fermi Dirac.

#### **Parameters**

h_bml	Input Hamiltonian matrix.
xi0_bml	Initial guess of first inverse.
p_bml	Output density matrix.
nsteps	Number of sp2 iterations.
посс	Number of occupied states.
mu	Shifted chemical potential
beta	Input inverse temperature.
osteps	Outer loop steps to converge chemical potential
occErrLimit	Occupation error limit.
threshold	Threshold for multiplication.

Definition at line 37 of file prg\_implicit\_fermi\_mod.F90.

Here is the call graph for this function:



# 9.10.2 Variable Documentation

9.10.2.1 integer, parameter prg\_implicit\_fermi\_mod::dp = kind(1.0d0) [private]

Definition at line 17 of file prg\_implicit\_fermi\_mod.F90.

# 9.11 prg\_initmatrices\_mod Module Reference

Initialization module.

#### **Functions/Subroutines**

- subroutine, public prg\_init\_hsmat (ham\_bml, over\_bml, bml\_type, mdim, norb)

  Initialize Hamiltonian and Overlap Matrix.
- subroutine, public prg\_init\_pzmat (rho\_bml, zmat\_bml, bml\_type, mdim, norb)

  Initialize Density matrix and Inverse square root Overlap.
- subroutine, public prg\_init\_ortho (orthoh\_bml, orthop\_bml, bml\_type, mdim, norb)

  Initialize The orthogonal versions of Hamiltonian and Density Matrix.

#### **Variables**

integer, parameter dp = kind(1.0d0)

# 9.11.1 Detailed Description

Initialization module.

Routines in this module are used to initialize several matrices that will be used in the code.

# 9.11.2 Function/Subroutine Documentation

9.11.2.1 subroutine, public prg\_initmatrices\_mod::prg\_init\_hsmat ( type(bml\_matrix\_t), intent(inout) ham\_bml, type(bml\_matrix\_t), intent(inout) over\_bml, character(20) bml\_type, integer, intent(inout) mdim, integer, intent(in) norb

Initialize Hamiltonian and Overlap Matrix.

Allocation of the Hamiltonian and Overlap matrix into bml formats.

#### **Parameters**

ham_bml	Hamiltonian in bml format.
over_bml	Overlap in bml format.
threshold	Threshold value for matrix elements.
mdim	Max nonzero elements per row for every row see [1] .
norb	Total number of orbitals.

Definition at line 29 of file prg\_initmatrices\_mod.F90.

9.11.2.2 subroutine, public prg\_initmatrices\_mod::prg\_init\_ortho ( type(bml\_matrix\_t), intent(inout) orthoh\_bml, type(bml\_matrix\_t), intent(inout) orthop\_bml, character(20) bml\_type, integer, intent(inout) mdim, integer, intent(inout) norb )

Initialize The orthogonal versions of Hamiltonian and Density Matrix.

Allocation of the orthogonal Hamiltonian and Density matrix into bml formats.

#### **Parameters**

orthoh_bml	Orthogonal Hamiltonian in bml format.
orthop_bml	Orthogonal Density Matrix in bml format.
threshold	Threshold value for matrix elements.
mdim	Max nonzero elements per row for every row see [1] .
norb	Total number of orbitals.

Definition at line 73 of file prg\_initmatrices\_mod.F90.

9.11.2.3 subroutine, public prg\_initmatrices\_mod::prg\_init\_pzmat ( type(bml\_matrix\_t), intent(inout) rho\_bml, type(bml\_matrix\_t), intent(inout) zmat\_bml, character(20) bml\_type, integer, intent(inout) mdim, integer, intent(in) norb )

Initialize Density matrix and Inverse square root Overlap.

Allocation of the Density matrix and Inverse square root Overlap matrix into bml formats.

#### **Parameters**

rho_bml	Density matrix in bml format.
zmat_bml	Inverse square root Overlap in bml format.
threshold	Threshold value for matrix elements.
mdim	Max nonzero elements per row for every row see [1] .
norb	Total number of orbitals.

Definition at line 51 of file prg\_initmatrices\_mod.F90.

# 9.11.3 Variable Documentation

9.11.3.1 integer, parameter prg\_initmatrices\_mod::dp = kind(1.0d0) [private]

Definition at line 14 of file prg\_initmatrices\_mod.F90.

# 9.12 prg\_kernelparser\_mod Module Reference

Some general parsing functions.

# **Functions/Subroutines**

• subroutine, public prg\_parsing\_kernel (keyvector\_char, valvector\_char, keyvector\_int, valvector\_int, keyvector\_re, valvector\_re, keyvector\_log, valvector\_log, filename, startstop)

The general parsing function. It is used to vectorize a set of "keywords" "value" pairs as included in a general input

• subroutine <a href="mailto:prg\_check\_valid">prg\_check\_valid</a> (invalidc)

Check for valid keywords (checks for an = sign)

# **Variables**

• integer, parameter dp = kind(1.0d0)

# 9.12.1 Detailed Description

Some general parsing functions.

# 9.12.2 Function/Subroutine Documentation

9.12.2.1 subroutine prg\_kernelparser\_mod::prg\_check\_valid ( character(len=\*), intent(in) invalidc ) [private]

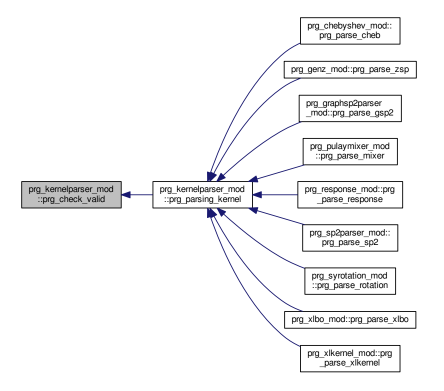
Check for valid keywords (checks for an = sign)

#### **Parameters**

invalidc	Keyword to check.
----------	-------------------

Definition at line 393 of file prg\_kernelparser\_mod.F90.

Here is the caller graph for this function:



9.12.2.2 subroutine, public prg\_kernelparser\_mod::prg\_parsing\_kernel ( character(50), dimension(:) keyvector\_char, character(100), dimension(:) valvector\_char, character(50), dimension(:) keyvector\_int, integer, dimension(:) valvector\_int, character(50), dimension(:) keyvector\_re, real(dp), dimension(:) valvector\_re, character(50), dimension(:) keyvector\_log, logical, dimension(:) valvector\_log, character(len=\*) filename, character(len=\*), dimension(2), intent(in), optional startstop)

The general parsing function. It is used to vectorize a set of "keywords" "value" pairs as included in a general input file.

#### Note

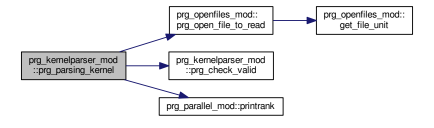
This parsing strategy can only parse a file of 500 lines by 500 words.

# Warning

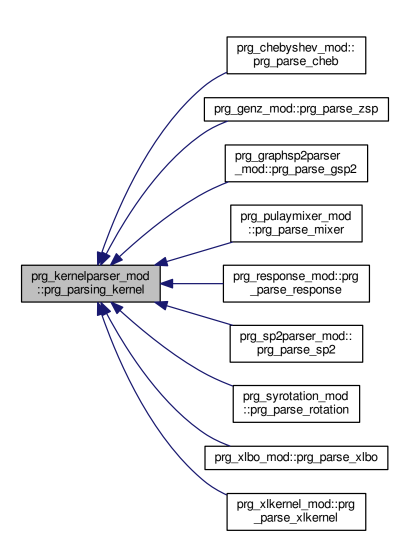
If the length of variable vect is changed, this could produce a segmentation fault.

Definition at line 30 of file prg\_kernelparser\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



# 9.12.3 Variable Documentation

9.12.3.1 integer, parameter prg\_kernelparser\_mod::dp = kind(1.0d0) [private]

Definition at line 13 of file prg\_kernelparser\_mod.F90.

# 9.13 prg\_nonortho\_mod Module Reference

Module to prg\_orthogonalize and prg\_deorthogonalize any operator.

# **Functions/Subroutines**

• subroutine, public prg\_orthogonalize (A\_bml, zmat\_bml, orthoA\_bml, threshold, bml\_type, verbose) This routine performs:  $A_{ortho} = Z^{\dagger}AZ$ .

• subroutine, public prg\_deorthogonalize (orthoA\_bml, zmat\_bml, a\_bml, threshold, bml\_type, verbose) This routine performs:  $A = ZA_{ortho}Z^{\dagger}$ .

# **Variables**

• integer, parameter dp = kind(1.0d0)

# 9.13.1 Detailed Description

Module to prg\_orthogonalize and prg\_deorthogonalize any operator.

Typically the Hamiltonin needs to be prg\_orthogonalized:  $H_{
m ortho}=Z^\dagger H Z$ 

Also, if the density matrix was obtained from the prg\_orthogonalized Hamiltonian, it can be prg\_deorthogonalized as:  $\rho = Z \rho_{\rm ortho} Z^\dagger$ 

# 9.13.2 Function/Subroutine Documentation

9.13.2.1 subroutine, public prg\_nonortho\_mod::prg\_deorthogonalize ( type(bml\_matrix\_t), intent(in) orthoA\_bml, type(bml\_matrix\_t), intent(in) unat\_bml, type(bml\_matrix\_t), intent(inout) a\_bml, real(dp) threshold, character(len=\*) bml\_type, integer verbose )

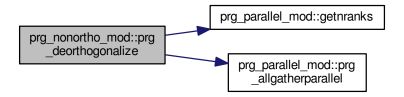
This routine performs:  $A = ZA_{ortho}Z^{\dagger}$ .

#### **Parameters**

orthoA_bml	Matrix to be prg_deorthogonalized.
zmat_bml	Congruence transform to be used.
A_bml	Matrix resulting from the prg_deorthogonalized in bml format.
threshold	Threshold value to be used in the matrix-matrix operations.
bml_type	bml format to be used.
verbose	Verbosity level.

Definition at line 82 of file prg\_nonortho\_mod.F90.

Here is the call graph for this function:



9.13.2.2 subroutine, public prg\_nonortho\_mod::prg\_orthogonalize ( type(bml\_matrix\_t), intent(inout) *A\_bml*, type(bml\_matrix\_t), intent(inout) *zmat\_bml*, type(bml\_matrix\_t), intent(inout) *orthoA\_bml*, real(dp), intent(in) *threshold*, character(len=\*), intent(in) *bml\_type*, integer, intent(in) *verbose* )

This routine performs:  $A_{ortho} = Z^{\dagger}AZ$ .

#### **Parameters**

A_bml	Matrix to be prg_orthogonalized in bml format.
zmat_bml	Congruence transform to be used.
orthoA_bml	Matrix resulting from the orthogonalization.
threshold	Threshold value to be used in the matrix-matrix operations.
bml_type	bml format to be used.
verbose	Verbosity level.

Definition at line 36 of file prg\_nonortho\_mod.F90.

# 9.13.3 Variable Documentation

9.13.3.1 integer, parameter prg\_nonortho\_mod::dp = kind(1.0d0) [private]

Definition at line 19 of file prg\_nonortho\_mod.F90.

# 9.14 prg\_normalize\_mod Module Reference

The prg\_normalize module.

#### **Functions/Subroutines**

• subroutine, public prg\_normalize (h\_bml)

Normalize a Hamiltonian matrix prior to running the SP2 algorithm.

• subroutine, public prg\_normalize\_fermi (h\_bml, h1, hN, mu)

Normalize a Hamiltonian matrix prior to running the truncated SP2 algorithm.

subroutine, public prg\_normalize\_implicit\_fermi (h\_bml, cnst, mu)

Normalize a Hamiltonian matrix prior to running the implicit fermi dirac algorithm.

• subroutine, public prg\_gershgorinreduction (gp)

Determine gershgorin bounds across all parts, local and distributed.

• subroutine, public prg\_normalize\_cheb (h\_bml, mu, emin, emax, alpha, scaledmu)

Normalize a Hamiltonian matrix prior to running the Chebyshev algorithm.

#### **Variables**

integer, parameter dp = kind(1.0d0)

# 9.14.1 Detailed Description

The prg normalize module.

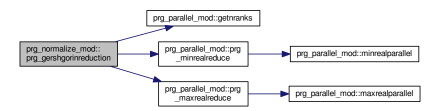
# 9.14.2 Function/Subroutine Documentation

9.14.2.1 subroutine, public prg\_normalize\_mod::prg\_gershgorinreduction (type(graph\_partitioning\_t), intent(inout) gp )

Determine gershgorin bounds across all parts, local and distributed.

Definition at line 101 of file prg normalize mod.F90.

Here is the call graph for this function:



9.14.2.2 subroutine, public prg\_normalize\_mod::prg\_normalize ( type(bml\_matrix\_t), intent(inout) h\_bml )

Normalize a Hamiltonian matrix prior to running the SP2 algorithm.

$$X0 = (e_max * I - H) / (e_max - e_min)$$

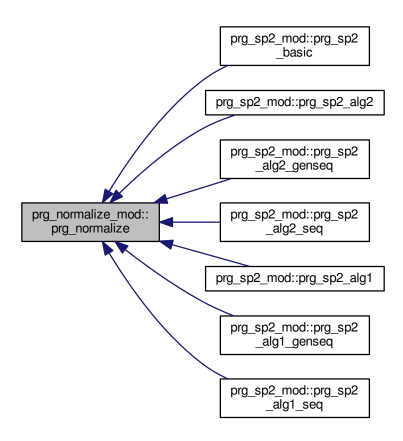
where e\_max and e\_min are obtained sing the Gershgorin circle theorem.

#### **Parameters**

h_bml Input/Output Hamiltonian matrix	X
---------------------------------------	---

Definition at line 33 of file prg\_normalize\_mod.F90.

Here is the caller graph for this function:



9.14.2.3 subroutine, public prg\_normalize\_mod::prg\_normalize\_cheb ( type(bml\_matrix\_t), intent(inout) h\_bml, real(dp), intent(in) mu, real(dp), intent(in) emin, real(dp), intent(in) emax, real(dp), intent(inout) alpha, real(dp), intent(inout) scaledmu )

Normalize a Hamiltonian matrix prior to running the Chebyshev algorithm.

$$X0 = 2*(H - e min*I) / (e max - e min) - I$$

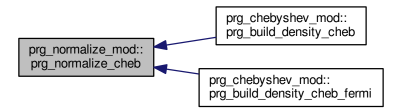
where e\_max and e\_min are obtained sing the Gershgorin circle theorem.

#### **Parameters**

h_bml	Input/Output Hamiltonian matrix
-------	---------------------------------

Definition at line 130 of file prg\_normalize\_mod.F90.

Here is the caller graph for this function:



9.14.2.4 subroutine, public prg\_normalize\_mod::prg\_normalize\_fermi ( type(bml\_matrix\_t), intent(inout) h\_bml, real(dp), intent(in) h1, real(dp), intent(in) hN, real(dp), intent(in) mu )

Normalize a Hamiltonian matrix prior to running the truncated SP2 algorithm.

$$X0 = ((hN-mu) * I - H) / (hN - h1) \text{ or } X0 = (hN*I-H0-mu*I)/(hN-h1)$$

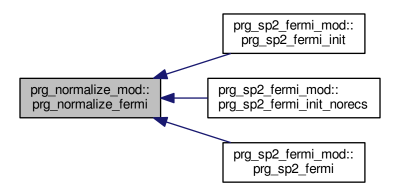
where h1 and hN are scaled Gershgorin bounds.

# Parameters

H_bml	Hamiltonian matrix
h1	Scaled minimum Gershgorin bound.
hN	Scaled maximum Gershgorin bound.
mu	Chemical potential

Definition at line 63 of file prg\_normalize\_mod.F90.

Here is the caller graph for this function:



9.14.2.5 subroutine, public prg\_normalize\_mod::prg\_normalize\_implicit\_fermi ( type(bml\_matrix\_t), intent(inout) h\_bml, real(dp), intent(in) cnst, real(dp), intent(in) mu )

Normalize a Hamiltonian matrix prior to running the implicit fermi dirac algorithm.

$$X0 = 0.5*II - cnst*(H0-mu0*II) \text{ or } X0 = (0.5 + cnst * mu0)*II - cnst* H0$$

#### **Parameters**

H_bml	Hamiltonian matrix
cnst	Constant based on beta and steps
ти	Chemical potential

Definition at line 87 of file prg\_normalize\_mod.F90.

Here is the caller graph for this function:



# 9.14.3 Variable Documentation

**9.14.3.1** integer, parameter prg\_normalize\_mod::dp = kind(1.0d0) [private]

Definition at line 15 of file prg\_normalize\_mod.F90.

# 9.15 prg\_openfiles\_mod Module Reference

Module to handle input output files for the PROGRESS lib.

# **Functions/Subroutines**

• integer function, public get\_file\_unit (io\_max)

Returns a unit number that is not in use.

• subroutine, public prg\_open\_file (io, name)

Opens a file to write.

• subroutine, public prg\_open\_file\_to\_read (io, name)

Opens a file to read.

# 9.15.1 Detailed Description

Module to handle input output files for the PROGRESS lib.

# 9.15.2 Function/Subroutine Documentation

9.15.2.1 integer function, public prg\_openfiles\_mod::get\_file\_unit ( integer io\_max )

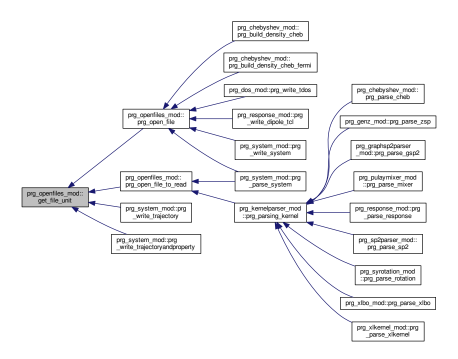
Returns a unit number that is not in use.

#### **Parameters**

io_max	Maximum units to search.
get_file_unit	Unit return to use for the file.

Definition at line 19 of file prg\_openfiles\_mod.F90.

Here is the caller graph for this function:



9.15.2.2 subroutine, public prg\_openfiles\_mod::prg\_open\_file ( integer io, character(len=\*) name )

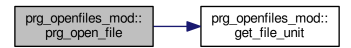
Opens a file to write.

#### **Parameters**

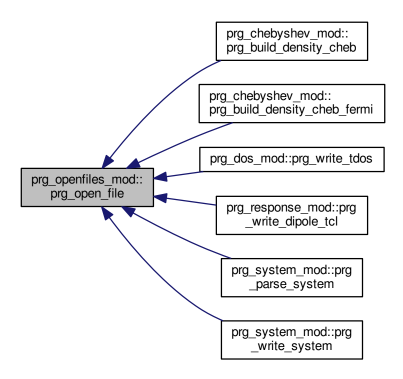
io	Unit for the file.
name	Name of the file.

Definition at line 38 of file prg\_openfiles\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



9.15.2.3 subroutine, public prg\_openfiles\_mod::prg\_open\_file\_to\_read ( integer io, character(len=\*) name )

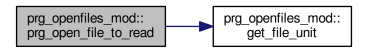
Opens a file to read.

### **Parameters**

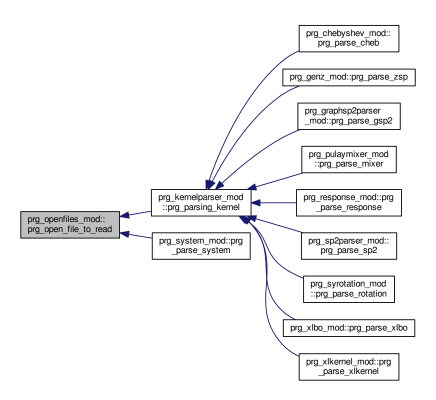
io	Unit for the file.
name	Name of the file.

Definition at line 54 of file prg\_openfiles\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



# 9.16 prg\_parallel\_mod Module Reference

The parallel module.

# **Data Types**

• type rankreducedata\_t

Data structure for rection over MPI ranks.

## **Functions/Subroutines**

- integer function, public getnranks ()
- integer function, public getmyrank ()
- integer function, public printrank ()
- subroutine, public prg\_initparallel ()
- subroutine, public prg\_shutdownparallel ()
- integer function saverequest (irequest)
- subroutine, public prg\_barrierparallel ()
- subroutine, public sendreceiveparallel (sendBuf, sendLen, dest, recvBuf, recvLen, source, nreceived)
- subroutine, public isendparallel (sendBuf, sendLen, dest)
- subroutine, public sendparallel (sendBuf, sendLen, dest)

- subroutine, public prg\_iprg\_recvparallel (recvBuf, recvLen, rind)
- subroutine, public <a href="mailto:prg\_recvparallel">prg\_recvparallel</a> (recvBuf, recvLen)
- subroutine, public sumintparallel (sendBuf, recvBuf, icount)
- subroutine, public sumrealparallel (sendBuf, recvBuf, icount)
- subroutine, public maxintparallel (sendBuf, recvBuf, icount)
- subroutine, public maxrealparallel (sendBuf, recvBuf, icount)
- subroutine, public minintparallel (sendBuf, recvBuf, icount)
- subroutine, public minrealparallel (sendBuf, recvBuf, icount)
- subroutine, public prg minrealreduce (rvalue)
- subroutine, public prg\_maxrealreduce (rvalue)
- subroutine, public prg\_maxintreduce2 (value1, value2)
- subroutine, public prg\_sumintreduce2 (value1, value2)
- subroutine, public prg\_sumrealreduce (value1)
- subroutine, public prg\_sumrealreduce2 (value1, value2)
- subroutine, public prg\_sumrealreduce3 (value1, value2, value3)
- subroutine, public prg\_sumrealreducen (valueVec, N)
- subroutine, public prg\_sumintreducen (valueVec, N)
- subroutine, public minrankrealparallel (sendBuf, recvBuf, icount)
- subroutine, public maxrankrealparallel (sendBuf, recvBuf, icount)
- subroutine, public prg bcastparallel (buf, blen, root)
- subroutine, public allgatherrealparallel (sendBuf, sendLen, recvBuf, recvLen)
- subroutine, public allgatherintparallel (sendBuf, sendLen, recvBuf, recvLen)
- subroutine, public allgathervrealparallel (sendBuf, sendLen, recvBuf, recvLen, recvDispl)
- subroutine, public allgathervintparallel (sendBuf, sendLen, recvBuf, recvLen, recvDispl)
- subroutine, public prg\_allsumrealreduceparallel (buf, buflen)
- subroutine, public prg\_allsumintreduceparallel (buf, buflen)
- subroutine, public prg\_allgatherparallel (a)
- subroutine, public prg wait ()

# Variables

- integer, parameter dp = kind(1.0d0)
- · integer myrank
- · integer nranks
- · integer ierr
- · integer reqcount
- integer, dimension(:), allocatable requestlist
- integer, dimension(:), allocatable rused

# 9.16.1 Detailed Description

The parallel module.

#### 9.16.2 Function/Subroutine Documentation

9.16.2.1 subroutine, public prg\_parallel\_mod::allgatherintparallel ( integer, dimension(\*), intent(in) sendBuf, integer, intent(in) sendLen, integer, dimension(\*), intent(out) recvBuf, integer, intent(in) recvLen )

Definition at line 660 of file prg parallel mod.F90.

9.16.2.2 subroutine, public prg\_parallel\_mod::allgatherrealparallel ( real(dp), dimension(\*), intent(in) sendBuf, integer, intent(in) sendLen, real(dp), dimension(\*), intent(out) recvBuf, integer, intent(in) recvLen )

Definition at line 644 of file prg\_parallel\_mod.F90.

9.16.2.3 subroutine, public prg\_parallel\_mod::allgathervintparallel ( integer, dimension(\*), intent(in) sendBuf, integer, intent(in) sendLen, integer, dimension(\*), intent(in) recvLen, integer, dimension(\*), intent(in) recvDispl )

Definition at line 696 of file prg\_parallel\_mod.F90.

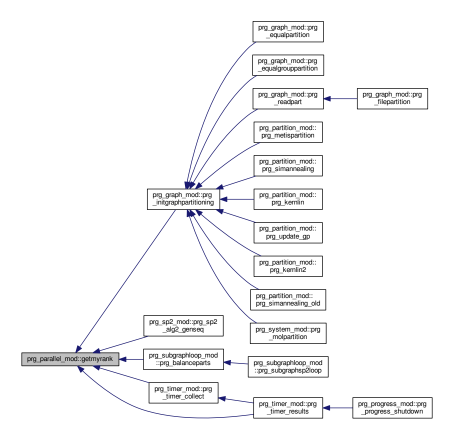
9.16.2.4 subroutine, public prg\_parallel\_mod::allgathervrealparallel ( real(dp), dimension(\*), intent(in) sendBuf, integer, intent(in) sendLen, real(dp), dimension(\*), intent(out) recvBuf, integer, dimension(\*), intent(in) recvLen, integer, dimension(\*), intent(in) recvDispl )

Definition at line 676 of file prg parallel mod.F90.

9.16.2.5 integer function, public prg\_parallel\_mod::getmyrank()

Definition at line 99 of file prg parallel mod.F90.

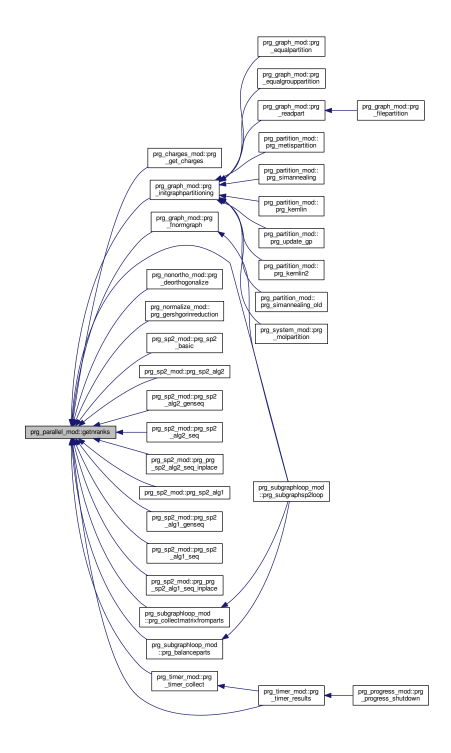
Here is the caller graph for this function:



### 9.16.2.6 integer function, public prg\_parallel\_mod::getnranks ( )

Definition at line 88 of file prg\_parallel\_mod.F90.

Here is the caller graph for this function:



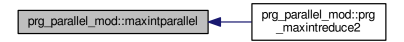
9.16.2.7 subroutine, public prg\_parallel\_mod::isendparallel ( real(dp), dimension(\*), intent(in) sendBuf, integer, intent(in) sendLen, integer, intent(in) dest )

Definition at line 230 of file prg\_parallel\_mod.F90.

9.16.2.8 subroutine, public prg\_parallel\_mod::maxintparallel ( integer, dimension(\*), intent(in) sendBuf, integer, dimension(\*), intent(out) recvBuf, integer, intent(in) icount )

Definition at line 337 of file prg\_parallel\_mod.F90.

Here is the caller graph for this function:



9.16.2.9 subroutine, public prg\_parallel\_mod::maxrankrealparallel ( type(rankreducedata\_t), dimension(\*), intent(in) sendBuf, type(rankreducedata\_t), dimension(\*), intent(out) recvBuf, integer, intent(in) icount )

Definition at line 607 of file prg\_parallel\_mod.F90.

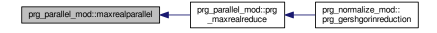
Here is the caller graph for this function:



9.16.2.10 subroutine, public prg\_parallel\_mod::maxrealparallel ( real(dp), dimension(\*), intent(in) sendBuf, real(dp), dimension(\*), intent(out) recvBuf, integer, intent(in) icount )

Definition at line 358 of file prg\_parallel\_mod.F90.

Here is the caller graph for this function:



9.16.2.11 subroutine, public prg\_parallel\_mod::minintparallel ( integer, dimension(\*), intent(in) sendBuf, integer, dimension(\*), intent(out) recvBuf, integer, intent(in) icount )

Definition at line 379 of file prg\_parallel\_mod.F90.

9.16.2.12 subroutine, public prg\_parallel\_mod::minrankrealparallel ( type(rankreducedata\_t), dimension(\*), intent(in) sendBuf, type(rankreducedata\_t), dimension(\*), intent(out) recvBuf, integer, intent(in) icount )

Definition at line 584 of file prg\_parallel\_mod.F90.

Here is the caller graph for this function:



9.16.2.13 subroutine, public prg\_parallel\_mod::minrealparallel ( real(dp), dimension(\*), intent(in) sendBuf, real(dp), dimension(\*), intent(out) recvBuf, integer, intent(in) icount )

Definition at line 400 of file prg\_parallel\_mod.F90.

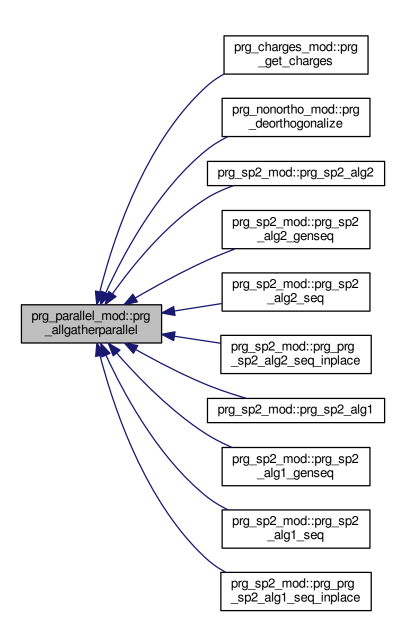
Here is the caller graph for this function:



9.16.2.14 subroutine, public prg\_parallel\_mod::prg\_allgatherparallel (type (bml\_matrix\_t), intent(inout) a)

Definition at line 744 of file prg\_parallel\_mod.F90.

Here is the caller graph for this function:



9.16.2.15 subroutine, public prg\_parallel\_mod::prg\_allsumintreduceparallel ( integer, dimension(\*), intent(inout) *buf*, integer, intent(in) *buflen* )

Definition at line 729 of file prg\_parallel\_mod.F90.

9.16.2.16 subroutine, public prg\_parallel\_mod::prg\_allsumrealreduceparallel ( real(dp), dimension(\*), intent(inout) *buf*, integer, intent(in) *buflen* )

Definition at line 714 of file prg\_parallel\_mod.F90.

9.16.2.17 subroutine, public prg\_parallel\_mod::prg\_barrierparallel ( )

Definition at line 196 of file prg parallel mod.F90.

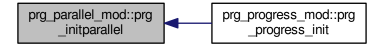
9.16.2.18 subroutine, public prg\_parallel\_mod::prg\_bcastparallel ( character, dimension(\*), intent(in) *buf*, integer, intent(in) *blen*, integer, intent(in) *root* )

Definition at line 630 of file prg\_parallel\_mod.F90.

9.16.2.19 subroutine, public prg\_parallel\_mod::prg\_initparallel ( )

Definition at line 127 of file prg\_parallel\_mod.F90.

Here is the caller graph for this function:



9.16.2.20 subroutine, public prg\_parallel\_mod::prg\_iprg\_recvparallel ( real(dp), dimension(\*) recvBuf, integer, intent(in) recvLen, integer rind )

Definition at line 261 of file prg\_parallel\_mod.F90.

Here is the call graph for this function:



9.16.2.21 subroutine, public prg\_parallel\_mod::prg\_maxintreduce2 ( integer, intent(inout) value1, integer, intent(inout) value2 )

Definition at line 453 of file prg\_parallel\_mod.F90.

Here is the call graph for this function:



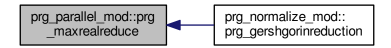
9.16.2.22 subroutine, public prg\_parallel\_mod::prg\_maxrealreduce ( real(dp), intent(inout) rvalue )

Definition at line 437 of file prg\_parallel\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



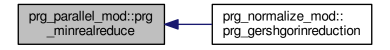
9.16.2.23 subroutine, public prg\_parallel\_mod::prg\_minrealreduce ( real(dp), intent(inout) rvalue )

Definition at line 421 of file prg\_parallel\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



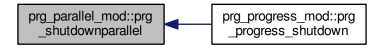
9.16.2.24 subroutine, public prg\_parallel\_mod::prg\_recvparallel ( real(dp), dimension(\*) recvBuf, integer, intent(in) recvLen )

Definition at line 279 of file prg\_parallel\_mod.F90.

9.16.2.25 subroutine, public prg\_parallel\_mod::prg\_shutdownparallel ( )

Definition at line 154 of file prg\_parallel\_mod.F90.

Here is the caller graph for this function:



9.16.2.26 subroutine, public prg\_parallel\_mod::prg\_sumintreduce2 ( integer, intent(inout) value1, integer, intent(inout) value2 )

Definition at line 471 of file prg\_parallel\_mod.F90.

Here is the call graph for this function:



9.16.2.27 subroutine, public prg\_parallel\_mod::prg\_sumintreducen ( integer, dimension(n), intent(inout) valueVec, integer, intent(in) N )

Definition at line 564 of file prg\_parallel\_mod.F90.

Here is the call graph for this function:



9.16.2.28 subroutine, public prg\_parallel\_mod::prg\_sumrealreduce ( real(dp), intent(inout) value1 )

Definition at line 489 of file prg\_parallel\_mod.F90.

Here is the call graph for this function:



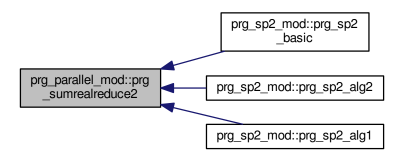
9.16.2.29 subroutine, public prg\_parallel\_mod::prg\_sumrealreduce2 ( real(dp), intent(inout) value1, real(dp), intent(inout) value2 )

Definition at line 505 of file prg\_parallel\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



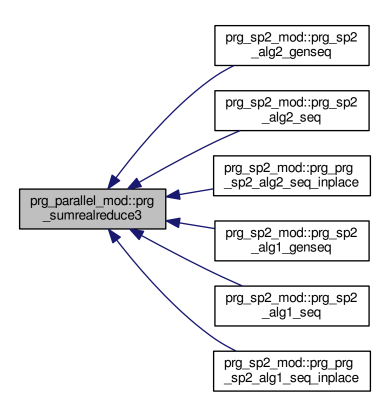
9.16.2.30 subroutine, public prg\_parallel\_mod::prg\_sumrealreduce3 ( real(dp), intent(inout) value1, real(dp), intent(inout) value2, real(dp), intent(inout) value3 )

Definition at line 523 of file prg\_parallel\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



9.16.2.31 subroutine, public prg\_parallel\_mod::prg\_sumrealreducen ( real(dp), dimension(n), intent(inout) valueVec, integer, intent(in) N )

Definition at line 543 of file prg\_parallel\_mod.F90.

Here is the call graph for this function:



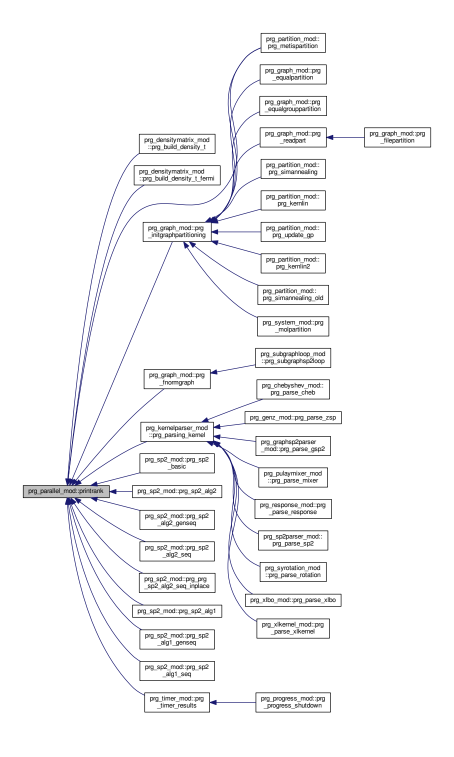
9.16.2.32 subroutine, public prg\_parallel\_mod::prg\_wait ( )

Definition at line 758 of file prg\_parallel\_mod.F90.

9.16.2.33 integer function, public prg\_parallel\_mod::printrank()

Definition at line 111 of file prg\_parallel\_mod.F90.

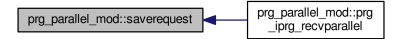
Here is the caller graph for this function:



9.16.2.34 integer function prg\_parallel\_mod::saverequest ( integer, intent(in) irequest ) [private]

Definition at line 170 of file prg\_parallel\_mod.F90.

Here is the caller graph for this function:



9.16.2.35 subroutine, public prg\_parallel\_mod::sendparallel ( real(dp), dimension(\*), intent(in) sendBuf, integer, intent(in) sendLen, integer, intent(in) dest )

Definition at line 246 of file prg\_parallel\_mod.F90.

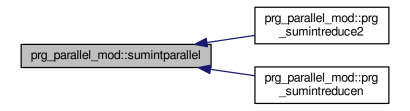
9.16.2.36 subroutine, public prg\_parallel\_mod::sendreceiveparallel ( real(dp), dimension(\*), intent(in) sendBuf, integer, intent(in) sendLen, integer, intent(in) dest, real(dp), dimension(\*), intent(out) recvBuf, integer, intent(in) recvLen, integer, intent(in) source, integer, intent(out) nreceived )

Definition at line 207 of file prg\_parallel\_mod.F90.

9.16.2.37 subroutine, public prg\_parallel\_mod::sumintparallel ( integer, dimension(\*), intent(in) sendBuf, integer, dimension(\*) recvBuf, integer, intent(in) icount )

Definition at line 295 of file prg\_parallel\_mod.F90.

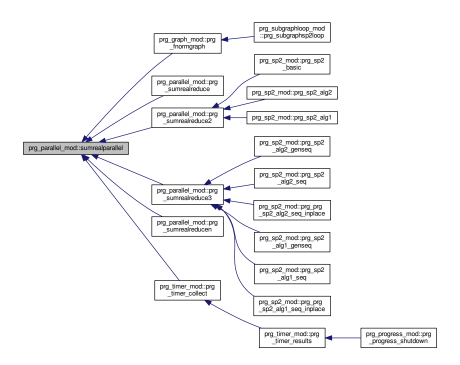
Here is the caller graph for this function:



9.16.2.38 subroutine, public prg\_parallel\_mod::sumrealparallel ( real(dp), dimension(\*), intent(in) sendBuf, real(dp), dimension(\*), intent(out) recvBuf, integer, intent(in) icount )

Definition at line 316 of file prg\_parallel\_mod.F90.

Here is the caller graph for this function:



### 9.16.3 Variable Documentation

9.16.3.1 integer, parameter prg\_parallel\_mod::dp = kind(1.0d0) [private]

Definition at line 26 of file prg\_parallel\_mod.F90.

9.16.3.2 integer prg\_parallel\_mod::ierr [private]

Definition at line 29 of file prg\_parallel\_mod.F90.

9.16.3.3 integer prg\_parallel\_mod::myrank [private]

Definition at line 28 of file prg\_parallel\_mod.F90.

9.16.3.4 integer prg\_parallel\_mod::nranks [private]

Definition at line 28 of file prg\_parallel\_mod.F90.

9.16.3.5 integer prg\_parallel\_mod::reqcount [private]

Definition at line 29 of file prg parallel mod.F90.

9.16.3.6 integer, dimension(:), allocatable prg\_parallel\_mod::requestlist [private]

Definition at line 30 of file prg\_parallel\_mod.F90.

**9.16.3.7** integer, dimension(:), allocatable prg\_parallel\_mod::rused [private]

Definition at line 30 of file prg parallel mod.F90.

# 9.17 prg\_partition\_mod Module Reference

The partition module.

### **Functions/Subroutines**

• subroutine, public prg\_metispartition (gp, ngroups, nnodes, xadj, adjncy, nparts, part, core\_count, CH\_count, Halo count, sumCubes, maxCH, smooth maxCH, pnorm)

Create graph partitions minizing number of cut edges.

subroutine, public prg\_costpartition (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_count, sum
 —
 Cubes, maxCH, smooth\_maxCH, pnorm)

Compute cost of a partition.

• subroutine, public update\_prg\_costpartition (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_← count, sumCubes, maxCH, smooth\_maxCH, pnorm, node, new\_part)

Update cost of partition and the different parameters node is moves into new\_part For each neighbor of node, the following cases hold: Case 1: neighbor is in old\_part Case 2: neighbor is in new\_part Case 3: neighbor is neither in old\_ or new\_part.

subroutine prg\_accept\_prob (it, prg\_delta, r)

Compute acceptance probability for simulated annealing.

• subroutine prg costindex (cost, sumCubes, maxCH, smooth maxCH, obj fun)

Choose objective function to work with.

• subroutine <a href="mailto:prg\_rand\_node">prg\_rand\_node</a> (gp, node, seed)

Pick a random node.

• subroutine, public prg\_simannealing (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm, niter, seed)

Graph partitioning based on Simulated Annealing.

• subroutine, public <a href="mailto:prg\_kernlin">prg\_kernlin</a> (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm, nconverg, seed)

Graph partitioning based on inspired by Kernighan-Lin Review METiS manual for description of k-way implementation of KL Pick a core together with its halos Place free vertices on a priority queue with (key, value) =(prg\_delta, best\_\circ part), with prg\_delta = change in obj\_value Dequeue and allow hill climbing.

- subroutine, public <a href="mailto:prg\_update\_gp">prg\_update\_gp</a> (gp, partNumber, core\_count)
- subroutine prg rand shuffle (array, seed)

Randomly shuffle array.

• subroutine, public prg\_check\_arrays (gp, core\_count, CH\_count, Halo\_count)

Error checking Checking that core\_count, CH\_count, Halo\_count match.

• subroutine, public prg\_kernlin\_queue (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm)

Greedy algorithm. At each step it chooses the (vertex, new\_part) pair with highest gain Currently implementation is very slow.

subroutine prg\_find\_best\_move (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_count, sum
 Cubes, maxCH, smooth\_maxCH, pnorm, best\_node, best\_part)

For kerlin\_queue to find (vertex, new\_part) pair with highest gain.

- subroutine, public prg\_kernlin2 (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_count, sum
   — Cubes, maxCH, smooth\_maxCH, pnorm)
- subroutine prg\_get\_largest\_hedge\_in\_part (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_← count, sumCubes, maxCH, smooth\_maxCH, pnorm, search\_part, largest\_Hedge)
- subroutine, public prg\_simannealing\_old (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm, niter, seed)

## **Variables**

- integer, parameter dp = kind(1.0d0)
- integer, parameter metis\_index\_kind = METIS\_INDEX\_KIND

From /usr/include/metis.h.

integer, parameter metis\_real\_kind = kind(METIS\_REAL\_KIND)

From /usr/include/metis.h.

# 9.17.1 Detailed Description

The partition module.

Contains different partitioning algorihms such as Metis, Simulated Annealing etc. Also contains optimization routines to improve upon existing partitioning, such as simulated annealing, etc.

## 9.17.2 Function/Subroutine Documentation

9.17.2.1 subroutine prg\_partition\_mod::prg\_accept\_prob ( integer, intent(in) it, real(dp), intent(in) prg\_delta, real, intent(inout) r ) [private]

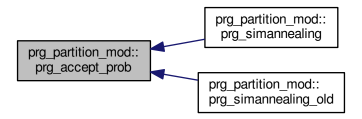
Compute acceptance probability for simulated annealing.

# **Parameters**

it	iteration
prg_delta	(new_obj_value - old_obj_value)
r	acceptance probability

Definition at line 489 of file prg\_partition\_mod.F90.

Here is the caller graph for this function:



9.17.2.2 subroutine, public prg\_partition\_mod::prg\_check\_arrays ( type (graph\_partitioning\_t), intent(inout) *gp,* integer, dimension(:), intent(inout), allocatable *CH\_count,* integer, dimension(:), intent(inout), allocatable *CH\_count,* integer, dimension(:,:), intent(inout), allocatable *Halo\_count* )

Error checking Checking that core\_count, CH\_count, Halo\_count match.

Definition at line 1146 of file prg\_partition\_mod.F90.

9.17.2.3 subroutine prg\_partition\_mod::prg\_costindex ( real(dp), intent(inout) cost, real(dp), intent(inout) sumCubes, real(dp), intent(inout) maxCH, real(dp), intent(inout) smooth\_maxCH, integer, intent(inout) obj\_fun ) [private]

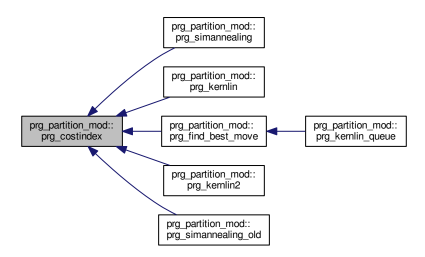
Choose objective function to work with.

## **Parameters**

cost	output according to chosen obj_fun
sumCubes	Sum of cubes obj value
maxCH	maximum core-halo part size obective value
obj_fun	0=sumcubes, 1=maxCH

Definition at line 507 of file prg\_partition\_mod.F90.

Here is the caller graph for this function:



9.17.2.4 subroutine, public prg\_partition\_mod::prg\_costpartition ( type (graph\_partitioning\_t), intent(inout) *gp*, integer, dimension(:), intent(inout), allocatable *xadj*, integer, dimension(:), intent(inout), allocatable *adjncy*, integer, dimension(:), intent(inout), allocatable *partNumber*, integer, dimension(:), intent(inout), allocatable *core\_count*, integer, dimension(:), intent(inout), allocatable *Halo\_count*, real(dp), intent(inout) *sumCubes*, real(dp), intent(inout) *maxCH*, real(dp), intent(inout) *smooth\_maxCH*, real(dp), intent(inout) *pnorm* )

Compute cost of a partition.

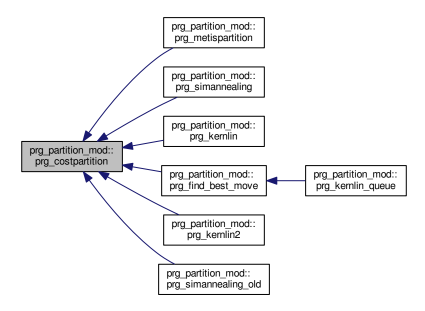
## **Parameters**

gp	Graph partitioning
xadj	CSR array of graph nodes
adjncy	CSR array of graph neighbors
nparts	Number of Parts
partNumber	Partition vector
core_count	Array: number of core vertices in each part
CH_count	Array: number of core+halo vertices in each part
Halo_count	2D Array of size nparts by totalNodes: Halo_count(i,j) = k, node j is a halo of part i with k connections
sumCubes	Sum of cubes objective value
maxCh	maximum core-halo part size obective value

prg initialize

Definition at line 327 of file prg\_partition\_mod.F90.

Here is the caller graph for this function:

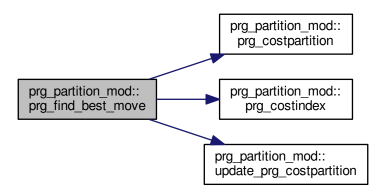


9.17.2.5 subroutine prg\_partition\_mod::prg\_find\_best\_move ( type (graph\_partitioning\_t), intent(inout) *gp*, integer, dimension(:), intent(inout), allocatable *xadj*, integer, dimension(:), intent(inout), allocatable *adjncy*, integer, dimension(:), intent(inout), allocatable *partNumber*, integer, dimension(:), intent(inout), allocatable *core\_count*, integer, dimension(:), intent(inout), allocatable *Halo\_count*, real(dp), intent(inout) *sumCubes*, real(dp), intent(inout) *maxCH*, real(dp), intent(inout) *smooth\_maxCH*, real(dp), intent(inout) *pnorm*, integer, intent(inout) *best\_node*, integer, intent(inout) *best\_part* ) [private]

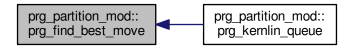
For kerlin\_queue to find (vertex, new\_part) pair with highest gain.

Definition at line 1209 of file prg\_partition\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:

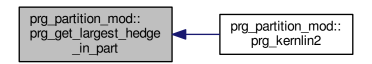


9.17.2.6 subroutine prg\_partition\_mod::prg\_get\_largest\_hedge\_in\_part ( type (graph\_partitioning\_t), intent(inout) *gp*, integer, dimension(:), intent(inout), allocatable *xadj*, integer, dimension(:), intent(inout), allocatable *adjncy*, integer, dimension(:), intent(inout), allocatable *core\_count*, integer, dimension(:), intent(inout), allocatable *core\_count*, integer, dimension(:), intent(inout), allocatable *Halo\_count*, real(dp), intent(inout) *sumCubes*, real(dp), intent(inout) *maxCH*, real(dp), intent(inout) *smooth\_maxCH*, real(dp), intent(inout) *pnorm*, integer, intent(inout) *search\_part*, integer, intent(inout) *largest\_Hedge*) [private]

i can be viewed as a hyperedge for all hyperedges in search part, pick the one with largest size

Definition at line 1420 of file prg partition mod.F90.

Here is the caller graph for this function:



9.17.2.7 subroutine, public prg\_partition\_mod::prg\_kernlin ( type (graph\_partitioning\_t), intent(inout) gp, integer, dimension(:), intent(inout), allocatable xadj, integer, dimension(:), intent(inout), allocatable adjncy, integer, dimension(:), intent(inout), allocatable partNumber, integer, dimension(:), intent(inout), allocatable core\_count, integer, dimension(:), intent(inout), allocatable Halo\_count, real(dp), intent(inout) sumCubes, real(dp), intent(inout) maxCH, real(dp), intent(inout) smooth\_maxCH, real(dp), intent(inout) pnorm, integer, intent(in) nconverg, integer, intent(inout) seed)

Graph partitioning based on inspired by Kernighan-Lin Review METiS manual for description of k-way implementation of KL Pick a core together with its halos Place free vertices on a priority queue with (key, value) =(prg\_delta, best\_part),with prg\_delta = change in obj\_value Dequeue and allow hill climbing.

#### **Parameters**

gp	Graph partitioning
----	--------------------

#### **Parameters**

xadj	CSR array of graph nodes
adjncy	CSR array of graph neighbors
nparts	Number of Parts
partNumber	Partition vector
core_count	Array: number of core vertices in each part
CH_count	Array: number of core+halo vertices in each part
Halo_count	2D Array of size nparts by totalNodes: Halo_count(i,j) = k, node j is a halo of part i with k connections
sumCubes	Sum of cubes objective value
maxCh	maximum core-halo part size obective value
nconverg	number of before convergence
seed	random number generator seed

Allocate arrays

Initialize variables

Initialize array of nodes

Randomize nodes

Compute current cost of partition

Choose objective function to minimize

iterate over the columns of the matrix, ie the hyperedges

KL iteration

let min\_part be the smallest CH\_part

Try and move free nodes to min\_part

lock vertices (climb\_counter) vertices have been accepted need to lock (climb\_counter) vertices Last vertex to be moved is node\_backup(climb\_counter)

reset

If all vertices locked, go to next iteration

If empty parts exit, place a vertex in max\_part there

Place j and it's neighbors that are in the max part into the empty part

Check Convergence

Check empty part exist move nodes from maxpart to empty part

move it neighbor in the max parts to the newpart

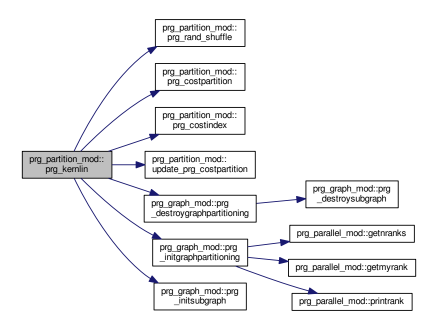
Update graph structure

Allocate subgraph structure

Assign node ids to sgraph

Definition at line 758 of file prg partition mod.F90.

Here is the call graph for this function:



9.17.2.8 subroutine, public prg\_partition\_mod::prg\_kernlin2 ( type (graph\_partitioning\_t), intent(inout) *gp,* integer, dimension(:), intent(inout), allocatable *xadj,* integer, dimension(:), intent(inout), allocatable *adjncy,* integer, dimension(:), intent(inout), allocatable *partNumber,* integer, dimension(:), intent(inout), allocatable *core\_count,* integer, dimension(:), intent(inout), allocatable *Halo\_count,* real(dp), intent(inout) *sumCubes,* real(dp), intent(inout) *maxCH,* real(dp), intent(inout) *smooth\_maxCH,* real(dp), intent(inout) *pnorm* )

# Allocate arrays

Pick hyperedge with largest size or random hyperedge with probability 0.5 We will change it to pick hyperedge with highest priority, where priority will be defined according to different metrics

Find part with smalles size (should be included in update\_prg\_costPartition

if current part is max, move to min\_part then move subsets (neighbors)

Move hyperedges to minCH part

Try and move intersecting hyperedges

Move k number of vertices. k should be small i.e k <=20, k set in prg\_Kernlin\_queue Only use this for small systems

Check empty part exist move nodes from maxpart to empty part

move it neighbor in the max parts to the newpart

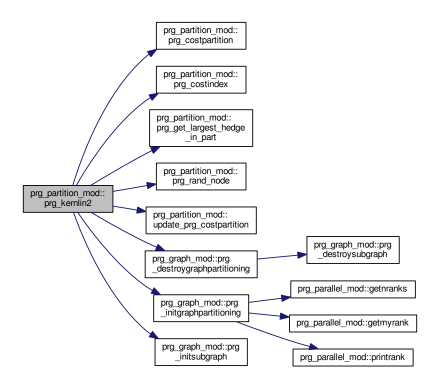
Update graph structure

Allocate subgraph structure

Assign node ids to sgraph

Definition at line 1257 of file prg\_partition\_mod.F90.

Here is the call graph for this function:

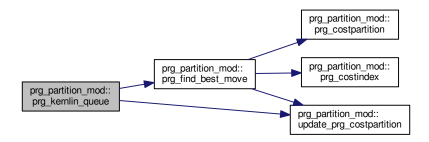


9.17.2.9 subroutine, public prg\_partition\_mod::prg\_kernlin\_queue ( type (graph\_partitioning\_t), intent(inout) *gp*, integer, dimension(:), intent(inout), allocatable *xadj*, integer, dimension(:), intent(inout), allocatable *adjncy*, integer, dimension(:), intent(inout), allocatable *partNumber*, integer, dimension(:), intent(inout), allocatable *core\_count*, integer, dimension(:), intent(inout), allocatable *Halo\_count*, real(dp), intent(inout) *sumCubes*, real(dp), intent(inout) *maxCH*, real(dp), intent(inout) *smooth\_maxCH*, real(dp), intent(inout) *pnorm* )

Greedy algorithm. At each step it chooses the (vertex, new\_part) pair with highest gain Currently implementation is very slow.

Definition at line 1173 of file prg\_partition\_mod.F90.

Here is the call graph for this function:



9.17.2.10 subroutine, public prg\_partition\_mod::prg\_metispartition ( type (graph\_partitioning\_t), intent(inout) *gp*, integer, intent(in) *ngroups*, integer, intent(in) *nnodes*, integer, dimension(:), intent(inout), allocatable *xadj*, integer, dimension(:), intent(inout), allocatable *adjncy*, integer, intent(inout) *nparts*, integer, dimension(:), intent(inout), allocatable *part*, integer, dimension(:), intent(inout), allocatable *core\_count*, integer, dimension(:), intent(inout), allocatable *Halo\_count*, real(dp), intent(inout) *sumCubes*, real(dp), intent(inout) *maxCH*, real(dp), intent(inout) *pnorm*)

Create graph partitions minizing number of cut edges.

#### **Parameters**

gp	Graph partitioning
ngroups	Number of groups/nodes
nnodes	Number of nodes
xadj	CSR array of graph nodes
adjncy	CSR array of graph neighbors
nparts	Number of Parts
part	Partition vector
core_count	Array: number of core vertices in each part
CH_count	Array: number of core+halo vertices in each part
Halo_count	2D Array of size nparts by totalNodes: Halo_count(i,j) = k, node j is a halo of part i with k connections
sumCubes	Sum of cubes objective value
maxCh	maximum core-halo part size obective value

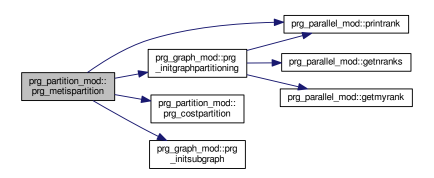
prg\_initialize

Partition graph into nparts'

Compute cost of partition

Definition at line 217 of file prg\_partition\_mod.F90.

Here is the call graph for this function:



9.17.2.11 subroutine prg\_partition\_mod::prg\_rand\_node ( type (graph\_partitioning\_t), intent(inout) *gp,* integer, intent(inout) *node,* integer, intent(inout) *seed* ) [private]

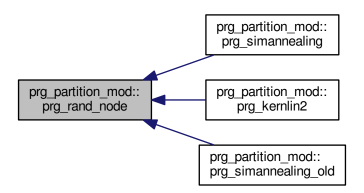
Pick a random node.

### **Parameters**

gp	graph partitioning structure
node	output node
seed	random seed

Definition at line 527 of file prg\_partition\_mod.F90.

Here is the caller graph for this function:



9.17.2.12 subroutine prg\_partition\_mod::prg\_rand\_shuffle ( integer, dimension(:), intent(inout) array, integer, intent(inout) seed
) [private]

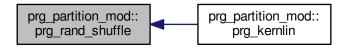
Randomly shuffle array.

Random seed

Shuffle array

Definition at line 1123 of file prg\_partition\_mod.F90.

Here is the caller graph for this function:



9.17.2.13 subroutine, public prg\_partition\_mod::prg\_simannealing ( type (graph\_partitioning\_t), intent(inout) gp, integer, dimension(:), intent(inout), allocatable xadj, integer, dimension(:), intent(inout), allocatable partNumber, integer, dimension(:), intent(inout), allocatable core\_count, integer, dimension(:), intent(inout), allocatable CH\_count, integer, dimension(:,:), intent(inout), allocatable Halo\_count, real(dp), intent(inout) sumCubes, real(dp), intent(inout) maxCH, real(dp), intent(inout) smooth\_maxCH, real(dp), intent(inout) pnorm, integer, intent(in) niter, integer, intent(inout) seed )

Graph partitioning based on Simulated Annealing.

## **Parameters**

gp	Graph partitioning
xadj	CSR array of graph nodes
adjncy	CSR array of graph neighbors
nparts	Number of Parts
partNumber	Partition vector
core_count	Array: number of core vertices in each part
CH_count	Array: number of core+halo vertices in each part
Halo_count	2D Array of size nparts by totalNodes: Halo_count(i,j) = k, node j is a halo of part i with k connections
sumCubes	Sum of cubes objective value
maxCh	maximum core-halo part size obective value
niter	Number of iterations
seed	Random seed

Compute current cost of partition

Choose objective function to minimize

#### Perform SA

Find part with smalles size (should be included in update\_prg\_costPartition

if part(node) == max\_ch\_part, try to move node and it's neighbors to min\_ch\_part else move neighbors to part(node)

Check empty part exist move nodes from maxpart to empty part

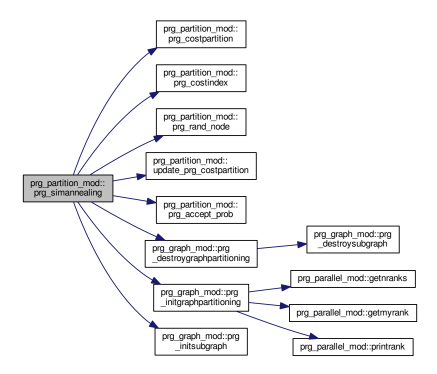
move it neighbor in the max parts to the newpart

Update graph structure

For debuging

Definition at line 552 of file prg partition mod.F90.

Here is the call graph for this function:



9.17.2.14 subroutine, public prg\_partition\_mod::prg\_simannealing\_old ( type (graph\_partitioning\_t), intent(inout) gp, integer, dimension(:), intent(inout), allocatable xadj, integer, dimension(:), intent(inout), allocatable adjncy, integer, dimension(:), intent(inout), allocatable partNumber, integer, dimension(:), intent(inout), allocatable core\_count, integer, dimension(:), intent(inout), allocatable Halo\_count, real(dp), intent(inout) sumCubes, real(dp), intent(inout) maxCH, real(dp), intent(inout) smooth\_maxCH, real(dp), intent(inout) pnorm, integer, intent(in) niter, integer, intent(inout) seed )

Compute current cost of partition

Choose objective function to minimize

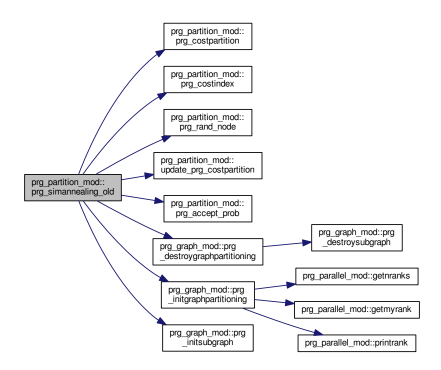
Perform SA

Update graph structure

For debuging

Definition at line 1453 of file prg\_partition\_mod.F90.

Here is the call graph for this function:



9.17.2.15 subroutine, public prg\_partition\_mod::prg\_update\_gp ( type (graph\_partitioning\_t), intent(inout) *gp*, integer, dimension(:), intent(inout), allocatable *partNumber*, integer, dimension(:), intent(inout), allocatable *core\_count* )

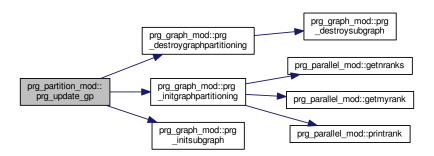
Update graph structure

Allocate subgraph structure

Assign node ids to sgraph

Definition at line 1082 of file prg\_partition\_mod.F90.

Here is the call graph for this function:



9.17.2.16 subroutine, public prg\_partition\_mod::update\_prg\_costpartition ( type (graph\_partitioning\_t), intent(inout) gp, integer, dimension(:), intent(inout), allocatable xadj, integer, dimension(:), intent(inout), allocatable adjncy, integer, dimension(:), intent(inout), allocatable partNumber, integer, dimension(:), intent(inout), allocatable core\_count, integer, dimension(:,:), intent(inout), allocatable Halo\_count, real(dp), intent(inout) sumCubes, real(dp), intent(inout) maxCH, real(dp), intent(inout) smooth\_maxCH, real(dp), intent(inout) pnorm, integer, intent(in) node, integer, intent(in) new\_part)

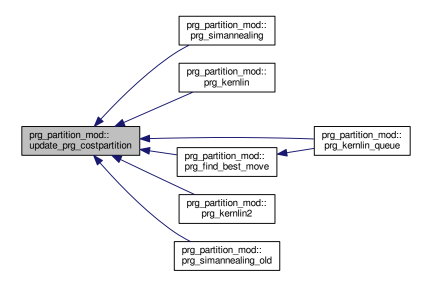
Update cost of partition and the different parameters node is moves into new\_part For each neighbor of node, the following cases hold: Case 1: neighbor is in old\_part Case 2: neighbor is in new\_part Case 3: neighbor is neither in old\_ or new\_part.

## **Parameters**

gp	Graph partitioning
xadj	CSR array of graph nodes
adjncy	CSR array of 1043365660.0000000graph neighbors
nparts	Number of Parts
partNumber	Partition vector
core_count	Array: number of core vertices in each part
CH_count	Array: number of core+halo vertices in each part
Halo_count	2D Array of size nparts by totalNodes: Halo_count(i,j) = k, node j is a halo of part i with k connections
sumCubes	Sum of cubes objective value
maxCh	maximum core-halo part size obective value
node	Vertex that has moved to new_part
new_part	new part that node has moved to

Definition at line 401 of file prg\_partition\_mod.F90.

Here is the caller graph for this function:



## 9.17.3 Variable Documentation

**9.17.3.1** integer, parameter prg\_partition\_mod::dp = kind(1.0d0) [private]

Definition at line 18 of file prg\_partition\_mod.F90.

**9.17.3.2** integer, parameter prg\_partition\_mod::metis\_index\_kind = METIS\_INDEX\_KIND [private]

From /usr/include/metis.h.

 $IDXTYPEWIDTH = 32 -> metis\_index\_kind = 4 \ IDXTYPEWIDTH = 64 -> metis\_index\_kind = 8$ 

Definition at line 24 of file prg\_partition\_mod.F90.

9.17.3.3 integer, parameter prg\_partition\_mod::metis\_real\_kind = kind(METIS\_REAL\_KIND) [private]

From /usr/include/metis.h.

REALTYPEWIDTH = 32 -> metis\_real\_kind = kind(0e0) REALTYPEWIDTH = 64 -> metis\_real\_kind = kind(0d0)

Definition at line 30 of file prg\_partition\_mod.F90.

# 9.18 prg\_progress\_mod Module Reference

The progress module.

# **Functions/Subroutines**

• subroutine, public prg\_progress\_init ()

Initialize progress.

• subroutine, public prg\_progress\_shutdown ()

Shutdown progress.

# Variables

• integer, parameter dp = kind(1.0d0)

# 9.18.1 Detailed Description

The progress module.

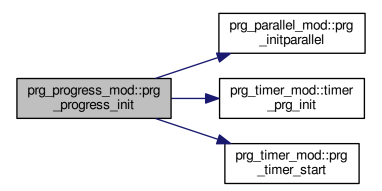
# 9.18.2 Function/Subroutine Documentation

9.18.2.1 subroutine, public prg\_progress\_mod::prg\_progress\_init ( )

Initialize progress.

Definition at line 25 of file prg\_progress\_mod.F90.

Here is the call graph for this function:

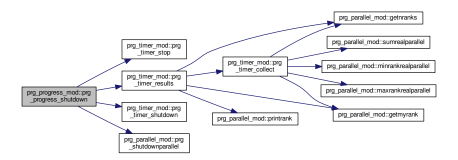


9.18.2.2 subroutine, public prg\_progress\_mod::prg\_progress\_shutdown ( )

Shutdown progress.

Definition at line 37 of file prg\_progress\_mod.F90.

Here is the call graph for this function:



# 9.18.3 Variable Documentation

9.18.3.1 integer, parameter prg\_progress\_mod::dp = kind(1.0d0) [private]

Definition at line 16 of file prg\_progress\_mod.F90.

# 9.19 prg\_ptable\_mod Module Reference

Periodic table of elements.

## **Functions/Subroutines**

- integer function, public element atomic number (symbol)
- integer function element atomic number upper (symbol)

### **Variables**

- integer, parameter nz = 103
- integer, parameter, private dp = kind(1.0d0)
- character(2), dimension(nz), parameter element\_symbol = [character(2) :: "H", "He", "Li", "Be", "B", "C", "N", "O", "F", "Ne", "Na", "Mg", "Al", "Si", "P", "S", "Cl", "Ar", "K", "Ca", "Sc", "Ti", "V", "Cr", "Mn", "Fe", "Co", "Ni", "Cu", "Zn", "Ga", "Ge", "As", "Se", "Br", "Kr", "Rb", "Sr", "Y", "Zr", "Nb", "Mo", "Tc", "Ru", "Rh", "Pd", "Ag", "Cd", "In", "Sn", "Sb", "Te", "I", "Xe", "Cs", "Ba", "La", "Ce", "Pr", "Nd", "Pm", "Sm", "Eu", "Gd", "Tb", "Dy", "Ho", "Er", "Tm", "Yb", "Lu", "Hf", "Ta", "W", "Re", "Os", "Ir", "Pt", "Au", "Hg", "TI", "Pb", "Bi", "Po", "At", "Rn", "Fr", "Ra", "Ac", "Th", "Pa", "U", "Np", "Pu", "Am", "Cm", "Bk", "Cf", "Es", "Fm", "Md", "No", "Lr"]

Element symbol.

character(2), dimension(nz), parameter element\_symbol\_upper = [character(2) :: "H", "HE", "LI", "BE", "B", "C", "N", "O", "F", "NE", "NA", "MG", "AL", "SI", "P", "S", "CL", "AR", "K", "CA", "SC", "TI", "V", "CR", "MN", "FE", "CO", "NI", "CU", "ZN", "GA", "GE", "AS", "SE", "BR", "KR", "RB", "SR", "Y", "ZR", "NB", "MO", "TC", "RU", "RH", "PD", "AG", "CD", "IN", "SN", "SB", "TE", "I", "XE", "CS", "BA", "LA", "CE", "PR", "ND", "PM", "SM", "EU", "GD", "TB", "DY", "HO", "ER", "TM", "YB", "LU", "HF", "TA", "W", "RE", "OS", "IR", "PT", "AU", "HG", "TL", "PB", "BI", "PO", "AT", "RN", "FR", "RA", "AC", "TH", "PA", "U", "NP", "PU", "AM", "CM", "BK", "CF", "ES", "FM", "MD", "NO", "LR"]

Element symbol upper.

character(20), dimension(nz), parameter element\_name = [character(20) :: "Hydrogen", "Helium", "Lithium", "Beryllium", "Boron", "Carbon", "Nitrogen", "Oxygen", "Fluorine", "Neon", "Sodium", "Magnesium", "Aluminium", "Silicon", "Phosphorus", "Sulfur", "Chlorine", "Argon", "Potassium", "Calcium", "Scandium", "Titanium", "Vanadium", "Chromium", "Manganese", "Iron", "Cobalt", "Nickel", "Copper", "Zinc", "Gallium", "Germanium", "Arsenic", "Selenium", "Bromine", "Krypton", "Rubidium", "Strontium", "Yttrium", "Zirconium", "Niobium", "Molybdenum", "Technetium", "Ruthenium", "Rhodium", "Palladium", "Silver", "Cadmium", "Indium", "Tin", "Antimony", "Tellurium", "Iodine", "Xenon", "Caesium", "Barium", "Lanthanum", "Cerium", "Praseodymium", "Neodymium", "Promethium", "Samarium", "Europium", "Gadolinium", "Terbium", "Dysprosium", "Holmium", "Erbium", "Thulium", "Ytterbium", "Lutetium", "Hafnium", "Tantalum", "Tungsten", "Rhenium", "Osmium", "Iridium", "Platinum", "Gold", "Mercury", "Thallium", "Lead", "Bismuth", "Polonium", "Astatine", "Radon", "Francium", "Radium", "Actinium", "Thorium", "Protactinium", "Uranium", "Neptunium", "Plutonium", "Americium", "Curium", "Berkelium", "Californium", "Einsteinium", "Fermium", "Mendelevium", "Nobelium", "Lawrencium"]

Element name.

• real(dp), dimension(nz), parameter element\_mass = (/ 1.007825032 , 4.002603254 , 7.01600455 , 9.0121822 , 11.0093054 , 12.0 , 14.003074005 , 15.99491462 , 18.99840322 , 19.992440175 , 22.989769281 , 23. 
9850417 , 26.98153863 , 27.976926532 , 30.97376163 , 31.972071 , 34.96885268 , 39.962383123 , 38. 
96370668 , 39.96259098 , 44.9559119 , 47.9479463 , 50.9439595 , 51.9405075 , 54.9380451 , 55.9349375 , 58.933195 , 57.9353429 , 62.9295975 , 63.929142 , 68.925573 , 73.921177 , 74.921596 , 79.916521 , 78. 
918337 , 83.911507 , 84.911789 , 87.905612 , 88.905848 , 89.904704 , 92.906378 , 97.905408 , 97.907216 , 101.904349 , 102.905504 , 105.903486 , 106.905097 , 113.903358 , 114.903878 , 119.902194 , 120. 
903815 , 129.906224 , 126.904473 , 131.904153 , 132.905451 , 137.905247 , 138.906353 , 139.905438 , 140.907652 , 141.907723 , 144.912749 , 151.919732 , 152.92123 , 157.924103 , 158.925346 , 163.929174 , 164.930322 , 165.930293 , 168.934213 , 173.938862 , 174.940771 , 179.94655 , 180.947995 , 183.950931 , 186.955753 , 191.96148 , 192.962926 , 194.964791 , 196.966568 , 201.970643 , 204.974427 , 207.976652 , 208.980398 , 208.98243 , 209.987148 , 222.017577 , 223.019735 , 226.025409 , 227.027752 , 232.038055 , 231.035884 , 238.050788 , 237.048173 , 244.064204 , 243.061381 , 247.070354 , 247.070307 , 251.079587 , 252.08298 , 257.095105 , 258.098431 , 259.10103 , 262.10963 /)

Element mass in atomic mass units (1.66 x 10-27 kg)

• real(dp), dimension(nz), parameter element\_vdwr = (/ 1.1 , 1.4 , 1.81 , 1.53 , 1.92 , 1.7 , 1.55 , 1.52 , 1.47 , 1.54 , 2.27 , 1.73 , 1.84 , 2.1 , 1.8 , 1.8 , 1.75 , 1.88 , 2.75 , 2.31 , 2.3 , 2.15 , 2.05 , 2.05 , 2.05 , 2.05 , 2.0 , 2.0 , 2.0 , 2.0 , 2.1 , 1.87 , 2.11 , 1.85 , 1.9 , 1.83 , 2.02 , 3.03 , 2.49 , 2.4 , 2.3 , 2.15 , 2.1 , 2.05 , 2.05 , 2.0 , 2.05 , 2.1 , 2.2 , 2.2 , 1.93 , 2.17 , 2.06 , 1.98 , 2.16 , 3.43 , 2.68 , 2.5 , 2.48 , 2.47 , 2.45 , 2.43 , 2.42 , 2.4 , 2.38 , 2.37 , 2.35 , 2.33 , 2.32 , 2.3 , 2.28 , 2.27 , 2.25 , 2.2 , 2.1 , 2.05 , 2.0 , 2.0 , 2.0 , 2.0 , 2.0 , 1.96 , 2.02 , 2.07 , 1.97 , 2.02 , 2.2 , 3.48 , 2.83 , 2.0 , 2.4 , 2.0 , 2.3 , 2.0 , 2

van der Waals radius (in Angstroms)

• real(dp), dimension(nz), parameter element\_covr = (/ 0.31 , 0.28 , 1.28 , 0.96 , 0.84 , 0.76 , 0.71 , 0.66 , 0.57 , 0.58 , 1.66 , 1.41 , 1.21 , 1.11 , 1.07 , 1.05 , 1.02 , 1.06 , 2.03 , 1.76 , 1.7 , 1.6 , 1.53 , 1.39 , 1.39 , 1.32 , 1.26 , 1.24 , 1.32 , 1.22 , 1.22 , 1.2 , 1.19 , 1.2 , 1.2 , 1.16 , 2.2 , 1.95 , 1.9 , 1.75 , 1.64 , 1.54 , 1.47 , 1.46 , 1.42 , 1.39 , 1.45 , 1.44 , 1.42 , 1.39 , 1.39 , 1.38 , 1.39 , 1.4 , 2.44 , 2.15 , 2.07 , 2.04 , 2.03 , 2.01 , 1.99 , 1.98 , 1.98 , 1.96 , 1.94 , 1.92 , 1.92 , 1.89 , 1.9 , 1.87 , 1.87 , 1.75 , 1.7 , 1.62 , 1.51 , 1.44 , 1.41 , 1.36 , 1.36 , 1.32 , 1.45 , 1.46 , 1.48 , 1.4 , 1.5 , 1.5 , 2.6 , 2.21 , 2.15 , 2.06 , 2.0 , 1.96 , 1.9 , 1.87 , 1.8 , 1.69 , 1.6 , 1.6 , 1.6 , 1.6 , 1.6 , 1.6 , 1.6 , 1.6 , 1.6 , 1.6 , 1.6 , 1.6 , 1.6 )

Covalent radius (in Angstroms)

real(dp), dimension(nz), parameter element\_ip = (/ 13.5984 , 24.5874 , 5.3917 , 9.3227 , 8.298 , 11.2603 , 14.5341 , 13.6181 , 17.4228 , 21.5645 , 5.1391 , 7.6462 , 5.9858 , 8.1517 , 10.4867 , 10.36 , 12.9676 , 15.7596 , 4.3407 , 6.1132 , 6.5615 , 6.8281 , 6.7462 , 6.7665 , 7.434 , 7.9024 , 7.881 , 7.6398 , 7.7264 , 9.3942 , 5.9993 , 7.8994 , 9.7886 , 9.7524 , 11.8138 , 13.9996 , 4.1771 , 5.6949 , 6.2173 , 6.6339 , 6.7589

 $, 7.0924 \;, 7.28 \;, 7.3605 \;, 7.4589 \;, 8.3369 \;, 7.5762 \;, 8.9938 \;, 5.7864 \;, 7.3439 \;, 8.6084 \;, 9.0096 \;, 10.4513 \;, \\ 12.1298 \;, 3.8939 \;, 5.2117 \;, 5.5769 \;, 5.5387 \;, 5.473 \;, 5.525 \;, 5.582 \;, 5.6437 \;, 5.6704 \;, 6.1498 \;, 5.8638 \;, 5.9389 \;, \\ 6.0215 \;, 6.1077 \;, 6.1843 \;, 6.2542 \;, 5.4259 \;, 6.8251 \;, 7.5496 \;, 7.864 \;, 7.8335 \;, 8.4382 \;, 8.967 \;, 8.9588 \;, \\ 9.2255 \;, 10.4375 \;, 6.1082 \;, 7.4167 \;, 7.2855 \;, 8.414 \;, 0.0 \;, 10.7485 \;, 4.0727 \;, 5.2784 \;, 5.17 \;, 6.3067 \;, 5.89 \;, \\ 6.1941 \;, 6.2657 \;, 6.026 \;, 5.9738 \;, 5.9914 \;, 6.1979 \;, 6.2817 \;, 6.42 \;, 6.5 \;, 6.58 \;, 6.65 \;, 4.9 /)$ 

Ionization energy (in eV)

• real(dp), dimension(nz), parameter element\_ea = (/ 0.75420375 , 0.0 , 0.618049 , 0.0 , 0.279723 , 1.262118 , -0.07 , 1.461112 , 3.4011887 , 0.0 , 0.547926 , 0.0 , 0.43283 , 1.389521 , 0.7465 , 2.0771029 , 3.612724 , 0.0 , 0.501459 , 0.02455 , 0.188 , 0.084 , 0.525 , 0.67584 , 0.0 , 0.151 , 0.6633 , 1.15716 , 1.23578 , 0.0 , 0.41 , 1.232712 , 0.814 , 2.02067 , 3.363588 , 0.0 , 0.485916 , 0.05206 , 0.307 , 0.426 , 0.893 , 0.7472 , 0.55 , 1.04638 , 1.14289 , 0.56214 , 1.30447 , 0.0 , 0.404 , 1.112066 , 1.047401 , 1.970875 , 3.059038 , 0.0 , 0.471626 , 0.14462 , 0.47 , 0.5 ,

Electron affprg inity (in eV)

• real(dp), dimension(nz), parameter atom\_en = (/ 2.2 , 0.0 , 0.98 , 1.57 , 2.04 , 2.55 , 3.04 , 3.44 , 3.98 , 0.0 , 0.93 , 1.31 , 1.61 , 1.9 , 2.19 , 2.58 , 3.16 , 0.0 , 0.82 , 1.0 , 1.36 , 1.54 , 1.63 , 1.66 , 1.55 , 1.83 , 1.88 , 1.91 , 1.9 , 1.65 , 1.81 , 2.01 , 2.18 , 2.55 , 2.96 , 3.0 , 0.82 , 0.95 , 1.22 , 1.33 , 1.6 , 2.16 , 1.9 , 2.2 , 2.28 , 2.2 , 1.93 , 1.69 , 1.78 , 1.96 , 2.05 , 2.1 , 2.66 , 2.6 , 0.79 , 0.89 , 1.1 , 1.12 , 1.13 , 1.14 , 0.0 , 1.17 , 0.0 , 1.2 , 0.0 , 1.22 , 1.23 , 1.24 , 1.25 , 0.0 , 1.27 , 1.3 , 1.5 , 2.36 , 1.9 , 2.2 , 2.2 , 2.28 , 2.54 , 2.0 , 1.62 , 2.33 , 2.02 , 2.0 , 2.2 , 0.0 , 0.7 , 0.9 , 1.1 , 1.3 , 1.5 , 1.38 , 1.36 , 1.28 , 1.3 , 1.3 , 1.3 , 1.3 , 1.3 , 1.3 , 1.3 , 1.3 , 1.3 , 0.0 /)

The Pauling electronegativity for this element.

The maximum expected number of bonds to this element.

- integer, dimension(nz), parameter element\_numel = (/1,2,1,2,3,4,5,6,7,8,1,2,3,4,5,6,7,8,1,2,3,4,5,6,7,8,1,2,3,4,5,6,7,8,9,10,11,12,13,14,15,16,17,18,1,2,3,4,5,6,7,8,9,10,11,12,13,14,15,16,17,18,1,2,3,4,5,6,7,8,9,10,11,12,13,14,15,16,17,18,19,20,21,22,23,24,25,26,27,28,29,30,31,32,1,2,3,4,5,6,7,8,9,10,11,12,13,14,15,16,17/)

  Last shell number of electrons.
- character(50), dimension(nz), parameter element\_econf = [character(50) :: "1s" , "1s2" , "1s22s" , "1s22s" , "1s22s2" , "1s22s22p2" , "1s22s22p3" , "1s22s22p3" , "1s22s22p4" , "1s22s22p5" , "1s22s22p6" , "[Ne]3s" , "[Ne]3s2" , "[Ne]3s23p2" , "[Ne]3s23p2" , "[Ne]3s23p3" , "[Ne]3s23p4" , "[Ne]3s23p5" , "[Ne]3s23p6" , "[Ar]3d54s2" , "[Ar]3d54s2" , "[Ar]3d64s2" , "[Ar]3d104s24p2" , "[Ar]3d104s24p3" , "[\leftarrow Ar]3d104s24p4" , "[Ar]3d104s24p5" , "[Ar]3d104s24p6" , "[Kr]5s" , "[Kr]5s2" , "[Kr]4d5s2" , "[Kr]4d5s2" , "[Kr]4d5s2" , "[Kr]4d5s2" , "[Kr]4d105s" , "[Cd]5p7 , "[Cd]5p7 , "[Cd]5p2" , "[Cd]5p4" , "[Cd]5p5" , "[Cd]5p6" , "[Xe]6s7 , "[Xe]6s2" , "[Xe]6s2" , "[Xe]4f166s2" , "[Xe]4f145d6s2" , "[Xe]4f145d7s2" , "[Xe]4f145d7s2" , "[Rn]6p7s2" , "[Rn]6p7s2" , "[Rn]5f167s2" , "[Rn]5f167s2"

The electronic configuration.

## 9.19.1 Detailed Description

Periodic table of elements.

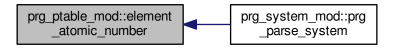
This data was prg\_generated with pybabel and openbable packages Openbabel:  $http://openbabel. \leftarrow org/dev-api/index.shtml Pybel: <math>https://openbabel.org/docs/dev/UseTheLibrary/\leftarrow Python_Pybel.html# Other sources includes NIST: <math>http://www.nist.gov/pml/data/ion\_\leftarrow energy.cfm$ 

### 9.19.2 Function/Subroutine Documentation

9.19.2.1 integer function, public prg\_ptable\_mod::element\_atomic\_number ( character(len=\*) symbol )

Definition at line 394 of file prg\_ptable\_mod.F90.

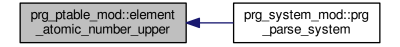
Here is the caller graph for this function:



9.19.2.2 integer function prg\_ptable\_mod::element\_atomic\_number\_upper ( character(len=\*) symbol )

Definition at line 408 of file prg\_ptable\_mod.F90.

Here is the caller graph for this function:



### 9.19.3 Variable Documentation

9.19.3.1 real(dp), dimension(nz), parameter prg\_ptable\_mod::atom\_en = (/ 2.2 , 0.0 , 0.98 , 1.57 , 2.04 , 2.55 , 3.04 , 3.44 , 3.98 , 0.0 , 0.93 , 1.31 , 1.61 , 1.9 , 2.19 , 2.58 , 3.16 , 0.0 , 0.82 , 1.0 , 1.36 , 1.54 , 1.63 , 1.66 , 1.55 , 1.83 , 1.88 , 1.91 , 1.9 , 1.65 , 1.81 , 2.01 , 2.18 , 2.55 , 2.96 , 3.0 , 0.82 , 0.95 , 1.22 , 1.33 , 1.6 , 2.16 , 1.9 , 2.2 , 2.28 , 2.2 , 1.93 , 1.69 , 1.78 , 1.96 , 2.05 , 2.1 , 2.66 , 2.6 , 0.79 , 0.89 , 1.1 , 1.12 , 1.13 , 1.14 , 0.0 , 1.17 , 0.0 , 1.2 , 0.0 , 1.22 , 1.23 , 1.24 , 1.25 , 0.0 , 1.27 , 1.3 , 1.5 , 2.36 , 1.9 , 2.2 , 2.2 , 2.28 , 2.54 , 2.0 , 1.62 , 2.33 , 2.02 , 2.0 , 2.2 , 0.0 , 0.7 , 0.9 , 1.1 , 1.3 , 1.5 , 1.38 , 1.36 , 1.28 , 1.3

The Pauling electronegativity for this element.

Definition at line 266 of file prg\_ptable\_mod.F90.

9.19.3.2 integer, parameter, private prg\_ptable\_mod::dp = kind(1.0d0) [private]

Definition at line 13 of file prg ptable mod.F90.

9.19.3.3 real(dp), dimension(nz), parameter prg\_ptable\_mod::element\_covr = (/ 0.31 , 0.28 , 1.28 , 0.96 , 0.84 , 0.76 , 0.71 , 0.66 , 0.57 , 0.58 , 1.66 , 1.41 , 1.21 , 1.11 , 1.07 , 1.05 , 1.02 , 1.06 , 2.03 , 1.76 , 1.7 , 1.6 , 1.53 , 1.39 , 1.39 , 1.32 , 1.26 , 1.24 , 1.32 , 1.22 , 1.22 , 1.2 , 1.19 , 1.2 , 1.2 , 1.16 , 2.2 , 1.95 , 1.9 , 1.75 , 1.64 , 1.54 , 1.47 , 1.46 , 1.42 , 1.39 , 1.45 , 1.44 , 1.42 , 1.39 , 1.39 , 1.38 , 1.39 , 1.4 , 2.44 , 2.15 , 2.07 , 2.04 , 2.03 , 2.01 , 1.99 , 1.98 , 1.98 , 1.96 , 1.94 , 1.92 , 1.92 , 1.89 , 1.9 , 1.87 , 1.87 , 1.75 , 1.7 , 1.62 , 1.51 , 1.44 , 1.41 , 1.36 , 1.36 , 1.32 , 1.45 , 1.46 , 1.48 , 1.4 , 1.5 , 1.5 , 2.6 , 2.21 , 2.15 , 2.06 , 2.0 , 1.96 , 1.9 , 1.87 , 1.8 , 1.69 , 1.6 ,

Covalent radius (in Angstroms)

Definition at line 173 of file prg\_ptable\_mod.F90.

 $9.19.3.4 \quad \text{real(dp), dimension(nz), parameter prg\_ptable\_mod::element\_ea} = (/\ 0.75420375\ ,\ 0.0\ ,\ 0.618049\ ,\ 0.0\ ,\ 0.279723\ ,\ 1.262118\ ,\ -0.07\ ,\ 1.461112\ ,\ 3.4011887\ ,\ 0.0\ ,\ 0.547926\ ,\ 0.0\ ,\ 0.43283\ ,\ 1.389521\ ,\ 0.7465\ ,\ 2.0771029\ ,\ 3.612724\ ,\ 0.0\ ,\ 0.501459\ ,\ 0.02455\ ,\ 0.188\ ,\ 0.084\ ,\ 0.525\ ,\ 0.67584\ ,\ 0.0\ ,\ 0.151\ ,\ 0.6633\ ,\ 1.15716\ ,\ 1.23578\ ,\ 0.0\ ,\ 0.41\ ,\ 1.232712\ ,\ 0.814\ ,\ 2.02067\ ,\ 3.363588\ ,\ 0.0\ ,\ 0.485916\ ,\ 0.05206\ ,\ 0.307\ ,\ 0.426\ ,\ 0.893\ ,\ 0.7472\ ,\ 0.55\ ,\ 1.04638\ ,\ 1.14289\ ,\ 0.56214\ ,\ 1.30447\ ,\ 0.0\ ,\ 0.404\ ,\ 1.112066\ ,\ 1.047401\ ,\ 1.970875\ ,\ 3.059038\ ,\ 0.0\ ,\ 0.471626\ ,\ 0.14462\ ,\ 0.47\ ,\ 0.5\ ,\$ 

Electron affprg\_inity (in eV)

Definition at line 235 of file prg ptable mod.F90.

9.19.3.5 character(50), dimension(nz), parameter prg\_ptable\_mod::element\_econf = [character(50) :: "1s", "1s2", "1s22s", "1s22s2", "1s22s22p", "1s22s22p2", "1s22s22p3", "1s22s22p4", "1s22s22p5", "1s22s22p6", "[Ne]3s2", "[Ne]3s2", "[Ne]3s23p6", "[Ne]3s23p2", "[Ne]3s23p3", "[Ne]3s23p4", "[Ne]3s23p5", "[Ne]3s23p6", "[Ar]4s4", "[Ar]4s2", "[Ar]3d4s2", "[Ar]3d24s2", "[Ar]3d34s2", "[Ar]3d54s4", "[Ar]3d54s2", "[Ar]3d64s2", "[Ar]3d74s2", "[Ar]3d74s2", "[Ar]3d104s24p4", "[Ar]3d104s24p5", "[Ar]3d104s24p6", "[Ar]3d104s24p6", "[Kr]5s2", "[Kr]4d5s2", "[Kr]4d5s2", "[Kr]4d25s2", "[Kr]4d25s2", "[Kr]4d45s", "[Kr]4d5s2", "[Kr]4d5s2", "[Kr]4d5s2", "[Kr]4d5s2", "[Kr]4d5s2", "[Kr]4d5s2", "[Kr]4d5s2", "[Kr]4d6s2", "[Xe]4f36s2", "[Xe]4f36s2", "[Xe]4f36s2", "[Xe]4f36s2", "[Xe]4f16s2", "[Xe]4f16s2", "[Xe]4f16s2", "[Xe]4f145d6s2", "[Xe]4f145d6s2", "[Xe]4f145d6s2", "[Xe]4f145d6s2", "[Xe]4f145d6s2", "[Xe]4f145d6s2", "[Xe]4f145d6s2", "[Xe]4f145d6s2", "[Xe]4f145d6s2", "[Xe]4f145d106s", "[Xe]4f145d106s2", "[Xe]4f145d1

The electronic configuration.

Definition at line 360 of file prg\_ptable\_mod.F90.

 $9.19.3.6 \quad real(dp), \\ dimension(nz), \\ parameter \\ prg\_ptable\_mod::element\_ip = (/ 13.5984, 24.5874, 5.3917, 9.3227, 8.298, 11.2603, 14.5341, 13.6181, 17.4228, 21.5645, 5.1391, 7.6462, 5.9858, 8.1517, 10.4867, 10.36, 12.9676, 15.7596, 4.3407, 6.1132, 6.5615, 6.8281, 6.7462, 6.7665, 7.434, 7.9024, 7.881, 7.6398, 7.7264, 9.3942, 5.9993, 7.8994, 9.7886, 9.7524, 11.8138, 13.9996, 4.1771, 5.6949, 6.2173, 6.6339, 6.7589, 7.0924, 7.28, 7.3605, 7.4589, 8.3369, 7.5762, 8.9938, 5.7864, 7.3439, 8.6084, 9.0096, 10.4513, 12.1298, 3.8939, 5.2117, 5.5769, 5.5387, 5.473, 5.525, 5.582, 5.6437, 5.6704, 6.1498, 5.8638, 5.9389, 6.0215, 6.1077, 6.1843, 6.2542, 5.4259, 6.8251, 7.5496, 7.864, 7.8335, 8.4382, 8.967, 8.9588, 9.2255, 10.4375, 6.1082, 7.4167, 7.2855, 8.414, 0.0, 10.7485, 4.0727, 5.2784, 5.17, 6.3067, 5.89, 6.1941, 6.2657, 6.026, 5.9738, 5.9914, 6.1979, 6.2817, 6.42, 6.5, 6.58, 6.65, 4.9 /)$ 

Ionization energy (in eV)

Definition at line 204 of file prg\_ptable\_mod.F90.

9.19.3.7 real(dp), dimension(nz), parameter prg\_ptable\_mod::element\_mass = (/ 1.007825032 , 4.002603254 , 7.01600455 , 9.0121822 , 11.0093054 , 12.0 , 14.003074005 , 15.99491462 , 18.99840322 , 19.992440175 , 22.989769281 , 23.9850417 , 26.98153863 , 27.976926532 , 30.97376163 , 31.972071 , 34.96885268 , 39.962383123 , 38.96370668 , 39.96259098 , 44.9559119 , 47.9479463 , 50.9439595 , 51.9405075 , 54.9380451 , 55.9349375 , 58.933195 , 57.9353429 , 62.9295975 , 63.929142 , 68.925573 , 73.921177 , 74.921596 , 79.916521 , 78.918337 , 83.911507 , 84.911789 , 87.905612 , 88.905848 , 89.904704 , 92.906378 , 97.905408 , 97.907216 , 101.904349 , 102.905504 , 105.903486 , 106.905097 , 113.903358 , 114.903878 , 119.902194 , 120.903815 , 129.906224 , 126.904473 , 131.904153 , 132.905451 , 137.905247 , 138.906353 , 139.905438 , 140.907652 , 141.907723 , 144.912749 , 151.919732 , 152.92123 , 157.924103 , 158.925346 , 163.929174 , 164.930322 , 165.930293 , 168.934213 , 173.938862 , 174.940771 , 179.94655 , 180.947995 , 183.950931 , 186.955753 , 191.96148 , 192.962926 , 194.964791 , 196.966568 , 201.970643 , 204.974427 , 207.976652 , 208.980398 , 208.98243 , 209.987148 , 222.017577 , 223.019735 , 226.025409 , 227.027752 , 232.038055 , 231.035884 , 238.050788 , 237.048173 , 244.064204 , 243.061381 , 247.070354 , 247.070307 , 251.079587 , 252.08298 , 257.095105 , 258.098431 , 259.10103 , 262.10963 /)

Element mass in atomic mass units (1.66 x 10-27 kg)

Definition at line 110 of file prg ptable mod.F90.

The maximum expected number of bonds to this element.

Definition at line 297 of file prg ptable mod.F90.

9.19.3.9 character(20), dimension(nz), parameter prg\_ptable\_mod::element\_name = [character(20) :: "Hydrogen", "Helium", "Lithium", "Beryllium", "Boron", "Carbon", "Nitrogen", "Oxygen", "Fluorine", "Neon", "Sodium", "Magnesium", "Aluminium", "Silicon", "Phosphorus", "Sulfur", "Chlorine", "Argon", "Potassium", "Calcium", "Scandium", "Titanium", "Vanadium", "Chromium", "Manganese", "Iron", "Cobalt", "Nickel", "Copper", "Zinc", "Gallium", "Germanium", "Arsenic", "Selenium", "Bromine", "Krypton", "Rubidium", "Strontium", "Yttrium", "Zirconium", "Niobium", "Molybdenum", "Technetium", "Ruthenium", "Rhodium", "Palladium", "Silver", "Cadmium", "Indium", "Tin", "Antimony", "Tellurium", "Iodine", "Xenon", "Caesium", "Barium", "Lanthanum", "Cerium", "Praseodymium", "Neodymium", "Promethium", "Samarium", "Europium", "Gadolinium", "Terbium", "Dysprosium", "Holmium", "Erbium", "Thulium", "Ytterbium", "Lutetium", "Hafnium", "Tantalum", "Tungsten", "Rhenium", "Osmium", "Iridium", "Platinum", "Gold", "Mercury", "Thallium", "Lead", "Bismuth", "Polonium", "Astatine", "Radon", "Francium", "Radium", "Actinium", "Thorium", "Protactinium", "Uranium", "Neptunium", "Plutonium", "Americium", "Curium", "Berkelium", "Californium", "Einsteinium", "Fermium", "Mendelevium", "Nobelium", "Lawrencium"]

Element name.

Definition at line 79 of file prg\_ptable\_mod.F90.

9.19.3.10 integer, dimension(nz), parameter prg\_ptable\_mod::element\_numel = (/ 1, 2, 1, 2, 3, 4, 5, 6, 7, 8, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17/)

Last shell number of electrons.

Definition at line 329 of file prg ptable mod.F90.

9.19.3.11 character(2), dimension(nz), parameter prg\_ptable\_mod::element\_symbol = [character(2) :: "H" , "He" , "Li" , "Be" , "B" , "C" , "N" , "O" , "F" , "Ne" , "Na" , "Mg" , "AI" , "Si" , "P" , "S" , "CI" , "Ar" , "K" , "Ca" , "Sc" , "Ti" , "V" , "Cr" , "Mn" , "Fe" , "Co" , "Ni" , "Cu" , "Zn" , "Ga" , "Ge" , "As" , "Se" , "Br" , "Kr" , "Rb" , "Sr" , "Y" , "Zr" , "Nb" , "Mo" , "Tc" , "Ru" , "Rh" , "Pd" , "Ag" , "Cd" , "In" , "Sn" , "Sb" , "Te" , "I" , "Xe" , "Cs" , "Ba" , "La" , "Ce" , "Pr" , "Nd" , "Pm" , "Sm" , "Eu" , "Gd" , "Tb" , "Dy" , "Ho" , "Er" , "Tm" , "Yb" , "Lu" , "Hf" , "Ta" , "W" , "Re" , "Os" , "Ir" , "Pt" , "Au" , "Hg" , "TI" , "Pb" , "Bi" , "Po" , "At" , "Rn" , "Fr" , "Ra" , "Ac" , "Th" , "Pa" , "U" , "Np" , "Pu" , "Am" , "Cm" , "Bk" , "Cf" , "Es" , "Fm" , "Md" , "No" , "Lr" ]

Element symbol.

Definition at line 17 of file prg ptable mod.F90.

9.19.3.12 character(2), dimension(nz), parameter prg\_ptable\_mod::element\_symbol\_upper = [character(2) :: "H" , "HE" , "LI" , "BE" , "B" , "C" , "N" , "O" , "F" , "NE" , "NA" , "MG" , "AL" , "SI" , "P" , "S" , "CL" , "AR" , "K" , "CA" , "SC" , "TI" , "V" , "CR" , "MN" , "FE" , "CO" , "NI" , "CU" , "ZN" , "GA" , "GE" , "AS" , "SE" , "BR" , "KR" , "RB" , "SR" , "Y" , "ZR" , "NB" , "MO" , "TC" , "RU" , "RH" , "PD" , "AG" , "CD" , "IN" , "SN" , "SB" , "TE" , "I" , "XE" , "CS" , "BA" , "LA" , "CE" , "PR" , "ND" , "PM" , "SM" , "EU" , "GD" , "TB" , "DY" , "HO" , "ER" , "TM" , "YB" , "LU" , "HF" , "TA" , "W" , "RE" , "OS" , "IR" , "PT" , "AU" , "HG" , "TL" , "PB" , "BI" , "PO" , "AT" , "RN" , "FR" , "RA" , "AC" , "TH" , "PA" , "U" , "NP" , "PU" , "AM" , "CM" , "BK" , "CF" , "ES" , "FM" , "MD" , "NO" , "LR" ]

Element symbol upper.

Definition at line 48 of file prg ptable mod.F90.

9.19.3.13 real(dp), dimension(nz), parameter prg\_ptable\_mod::element\_vdwr = (/ 1.1 , 1.4 , 1.81 , 1.53 , 1.92 , 1.7 , 1.55 , 1.52 , 1.47 , 1.54 , 2.27 , 1.73 , 1.84 , 2.1 , 1.8 , 1.8 , 1.75 , 1.88 , 2.75 , 2.31 , 2.3 , 2.15 , 2.05 , 2.05 , 2.05 , 2.05 , 2.05 , 2.0 , 2.0 , 2.0 , 2.1 , 1.87 , 2.11 , 1.85 , 1.9 , 1.83 , 2.02 , 3.03 , 2.49 , 2.4 , 2.3 , 2.15 , 2.1 , 2.05 , 2.05 , 2.0 , 2.05 , 2.1 , 2.2 , 2.2 , 1.93 , 2.17 , 2.06 , 1.98 , 2.16 , 3.43 , 2.68 , 2.5 , 2.48 , 2.47 , 2.45 , 2.43 , 2.42 , 2.4 , 2.38 , 2.37 , 2.35 , 2.33 , 2.32 , 2.3 , 2.28 , 2.27 , 2.25 , 2.2 , 2.1 , 2.05 , 2.0 ,

van der Waals radius (in Angstroms)

Definition at line 141 of file prg ptable mod.F90.

9.19.3.14 integer, parameter prg\_ptable\_mod::nz = 103

Definition at line 12 of file prg\_ptable\_mod.F90.

# 9.20 prg\_pulaycomponent\_mod Module Reference

Produces a matrix to get the Pulay Component of the forces.

### **Functions/Subroutines**

- subroutine, public prg\_pulaycomponent0 (rho\_bml, ham\_bml, pcm\_bml, threshold, M, bml\_type, verbose)  $At T = 0K, P = \rho H \rho.$
- subroutine, public prg\_pulaycomponentt (rho\_bml, ham\_bml, zmat\_bml, pcm\_bml, threshold, M, bml\_type, verbose)

At 
$$T > 0K$$
,  $P = \rho H S^{-1} + S^{-1} H \rho$ .

• subroutine, public prg\_get\_pulayforce (nats, zmat\_bml, ham\_bml, rho\_bml, dSx\_bml, dSy\_bml, dSz\_bml, hindex, FPUL, threshold)

Pulay Force FPUL from  $2Tr[ZZ'HD\frac{dS}{dR}]$ .

### **Variables**

integer, parameter dp = kind(1.0d0)

# 9.20.1 Detailed Description

Produces a matrix to get the Pulay Component of the forces.

For a further explanation please see Niklasson 2008 [3]

### 9.20.2 Function/Subroutine Documentation

9.20.2.1 subroutine, public prg\_pulaycomponent\_mod::prg\_get\_pulayforce ( integer, intent(in) nats, type(bml\_matrix\_t), intent(in) zmat\_bml, type(bml\_matrix\_t), intent(in) ham\_bml, type(bml\_matrix\_t), intent(in) rho\_bml, type(bml\_matrix\_t), intent(in) dSx\_bml, type(bml\_matrix\_t), intent(in) dSy\_bml, type(bml\_matrix\_t), intent(in) dSz\_bml, integer, dimension(:,:), intent(in) hindex, real(dp), dimension(:,:), intent(inout), allocatable FPUL, real(dp), intent(in) threshold)

Pulay Force FPUL from  $2Tr[ZZ'HD\frac{dS}{dR}]$ .

#### **Parameters**

nats	Number of atoms.
zmat_bml	Congruence transform in bml format.
rho_bml	Density matrix.
dSx_bml	x derivative of S.
dSy_bml	y derivative of S.
dSz_bml	z derivative of S.
hindex	Contains the Hamiltonian indices for every atom (see get_hindex).

Definition at line 152 of file prg\_pulaycomponent\_mod.F90.

9.20.2.2 subroutine, public prg\_pulaycomponent\_mod::prg\_pulaycomponent0 ( type(bml\_matrix\_t), intent(in) *rho\_bml*, type(bml\_matrix\_t), intent(in) *ham\_bml*, type(bml\_matrix\_t), intent(inout) *pcm\_bml*, real(dp), intent(in) *threshold*, integer, intent(in) *M*, character(20), intent(in) *bml\_type*, integer *verbose* )

At 
$$T = 0K$$
,  $P = \rho H \rho$ .

#### **Parameters**

rho_bml	Density matrix in bml format.
ham_bml	Hamiltonian matrix in bml format.
pcm_bml	Pulay matix output in bml format.
threshold	Threshold for the matrix elements.
М	Maximum nonzero values per row.
bml_type	Bml format type.
verbose	Verbosity level.

**Todo** M and bml type will have to be removed from the input parameter.

Definition at line 32 of file prg\_pulaycomponent\_mod.F90.

9.20.2.3 subroutine, public prg\_pulaycomponent\_mod::prg\_pulaycomponentt ( type(bml\_matrix\_t), intent(in) *rho\_bml*, type(bml\_matrix\_t), intent(in) *type*(bml\_matrix\_t), intent(in) *zmat\_bml*, type(bml\_matrix\_t), intent(inout) *pcm\_bml*, real(dp), intent(in) *threshold*, integer, intent(in) *M*, character(20), intent(in) *bml\_type*, integer *verbose* )

At 
$$T > 0K$$
,  $P = \rho H S^{-1} + S^{-1} H \rho$ .

#### **Parameters**

rho_bml	Density matrix in bml format.
ham_bml	Hamiltonian matrix in bml format.
Z_bml	Congruence transform in bml format.
pcm_bml	Pulay matrix output in bml format.
threshold	Threshold for the matrix elements.
М	Maximum nonzero values per row.
bml_type	Bml format type.
verbose	Verbosity level.

**Todo** M and bml\_type will have to be removed from the input parameter.

Definition at line 83 of file prg\_pulaycomponent\_mod.F90.

# 9.20.3 Variable Documentation

9.20.3.1 integer, parameter prg\_pulaycomponent\_mod::dp = kind(1.0d0) [private]

Definition at line 13 of file prg\_pulaycomponent\_mod.F90.

# 9.21 prg\_pulaymixer\_mod Module Reference

Pulay mixer mode.

# **Data Types**

• type mx\_type

### **Functions/Subroutines**

• subroutine, public prg\_parse\_mixer (input, filename)

The parser for the mixer routines.

- subroutine, public prg\_qmixer (charges, oldcharges, dqin, dqout, scferror, piter, pulaycoef, mpulay, verbose)

  Mixing the charges to acelerate scf convergence.
- subroutine, public prg\_linearmixer (charges, oldcharges, scferror, linmixcoef, verbose)

  Routine to perform linear mixing.

### **Variables**

• integer, parameter dp = kind(1.0d0)

# 9.21.1 Detailed Description

Pulay mixer mode.

Gets the best coefficient for mixing the charges during scf.

**Todo** add the density matrix mixer.

# 9.21.2 Function/Subroutine Documentation

9.21.2.1 subroutine, public prg\_pulaymixer\_mod::prg\_linearmixer ( real(dp), dimension(:), intent(inout), allocatable *charges*, real(dp), dimension(:), intent(inout), allocatable *oldcharges*, real(dp), intent(inout) *scferror*, real(dp), intent(in) *linmixcoef*, integer, intent(in) *verbose* )

Routine to perform linear mixing.

#### **Parameters**

charges	Actual charges of the system.
oldcharges	Previous scf charges.
scferror	SCF error.
linmixcoef	Mixing coefficient.
verbose	Verbosity level.

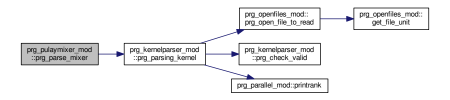
Definition at line 237 of file prg\_pulaymixer\_mod.F90.

9.21.2.2 subroutine, public prg\_pulaymixer\_mod::prg\_parse\_mixer ( type(mx\_type), intent(inout) input, character(len=\*) filename )

The parser for the mixer routines.

Definition at line 42 of file prg\_pulaymixer\_mod.F90.

Here is the call graph for this function:



9.21.2.3 subroutine, public prg\_pulaymixer\_mod::prg\_qmixer ( real(dp), dimension(:), intent(inout) charges, real(dp), dimension(:), intent(inout), allocatable oldcharges, real(dp), dimension(:,:), intent(inout), allocatable dquin, real(dp), dimension(:,:), intent(inout), allocatable dquin, real(dp), intent(inout) scferror, integer piter, real(dp), intent(in) pulaycoef, integer, intent(in) mpulay, integer, intent(in) verbose)

Mixing the charges to acelerate scf convergence.

# **Parameters**

charges	System charges.
oldcharges	Old charges of the system.
dqin	Matrix for charges history in.
dqout	Matrix for charges history out.
scferror	SCF error.
piter	scf iteration number.
pulaycoef	Coefficient for pulay mixing (generally between 0.01 and 0.1).
mpulay	Number of matrices stored (generally 3-5).
verbose	Different levels of verbosity.

Definition at line 103 of file prg\_pulaymixer\_mod.F90.

# 9.21.3 Variable Documentation

9.21.3.1 integer, parameter prg\_pulaymixer\_mod::dp = kind(1.0d0) [private]

Definition at line 14 of file prg\_pulaymixer\_mod.F90.

# 9.22 prg\_quantumdynamics\_mod Module Reference

A module to add in common quantum dynamical operations.

### **Functions/Subroutines**

• subroutine, public prg\_kick\_density (kick\_direc, kick\_mag, dens, norbs, mdim, S, SINV, which\_atom, r, bml-type, thresh)

Provides perturbation to initial density matrix in the form of an electric field kick. This routine does:  $\rho_{\hat{kick}} = \exp{\frac{-i}{\hbar}\hat{V}\hat{\rho}\hat{S}}\exp{\frac{i}{\hbar}\hat{V}\hat{S}^{-1}}$  where  $\hat{V}$  is the field disturbance.

• subroutine, public prg\_get\_sparsity\_cplxmat (matrix\_type, element\_type, thresh, a\_dense)

This computes the sparsity of a complex matrix given a threshold value This routine does:  $f = \frac{N_0}{N_{tot}}$  where f is the sparsity,  $N_0$  is the number of values less than the threshold, and  $N_{tot}$  is the total number of values. The sparsity and threshold are printed to the screen.

subroutine, public prg\_get\_sparsity\_realmat (matrix\_type, element\_type, thresh, a\_dense)

This computes the sparsity of a real matrix given a threshold value This routine does:  $f=\frac{N_0}{N_{tot}}$  where f is the sparsity,  $N_0$  is the number of values less than the threshold, and  $N_{tot}$  is the total number of values. The sparsity and threshold are printed to the screen.

subroutine, public prg\_kick\_density\_bml (kick\_direc, kick\_mag, rho\_bml, s\_bml, sinv\_bml, mdim, which\_
 atom, r, matrix\_type, thresh)

Provides perturbation to initial density matrix in the form of an electric field kick given input matricies in BML format. This routine does:  $\rho_{k\hat{i}ck} = \exp{\frac{-i}{\hbar}\hat{V}\hat{\rho}\hat{S}} \exp{\frac{i}{\hbar}\hat{V}\hat{S}^{-1}}$  where  $\hat{V}$  is the field disturbance.

subroutine, public prg\_lvni\_bml (h1\_bml, sinv\_bml, dt, hbar, rhoold\_bml, rho\_bml, aux\_bml, matrix\_type, mdim, thresh)

Performs Liouville-von Neumann integration using leap-frog method. This routine does:  $\hat{\rho}(t+\Delta t)=\hat{\rho}(t-\Delta t)+2\Delta t \frac{\partial \hat{\rho}(t)}{\partial t}$  where the time derivative of the density matrix is defined as follows:  $\frac{\partial \hat{\rho}(t)}{\partial t}=\frac{-i}{\hbar}\left(S^{-1}\hat{H}(t)\hat{\rho}(t)-\hat{\rho}(t)\hat{H}(t)S^{-1}\right)$ .

• subroutine, public prg\_getcharge (rho\_bml, s\_bml, charges, aux\_bml, z, spindex, N, nats, thresh)

Constructs the charges from the density matrix.

• subroutine, public prg\_getdipole (charges, r, mu)

This routine computes the dipole moment of the system with units determined by the units of the coordinate matrix and charges given.

• subroutine, public prg\_excitation (fill\_mat, orbit\_orig, orbit\_exci)

Produce an excitation in the initially calculated density matrix to.

### **Variables**

• integer, parameter dp = kind(1.0d0)

# 9.22.1 Detailed Description

A module to add in common quantum dynamical operations.

This module contains routines that perform the following tasks: apply an excitation or perturbation to the initial density matrix, compute the comutator of two two matricies, calculate the sparsity of a real or complex matrix, and time evolve a density matrix using Liouville-von Neumann equation with the leap-frog method of integration.

### 9.22.2 Function/Subroutine Documentation

9.22.2.1 subroutine, public prg\_quantumdynamics\_mod::prg\_excitation ( integer, dimension(:), intent(inout) *fill\_mat*, integer, intent(in) *orbit\_orig*, integer, intent(in) *orbit\_exci* )

Produce an excitation in the initially calculated density matrix to.

Definition at line 307 of file prg\_quantumdynamics\_mod.F90.

9.22.2.2 subroutine, public prg\_quantumdynamics\_mod::prg\_get\_sparsity\_cplxmat ( character(len=\*), intent(in) matrix\_type, character(len=\*), intent(in) element\_type, real(dp), intent(in) thresh, complex(dp), dimension(:,:), intent(in) a dense )

This computes the sparsity of a complex matrix given a threshold value This routine does:  $f = \frac{N_0}{N_{tot}}$  where f is the sparsity,  $N_0$  is the number of values less than the threshold, and  $N_{tot}$  is the total number of values. The sparsity and threshold are printed to the screen.

#### **Parameters**

matrix_type	the BML matrix type
element_type	the BML element type
thresh	the threshold for sparsity evaluation
a_dense	the dense complex matrix to be evaluated for sparsity

Definition at line 98 of file prg\_quantum dynamics\_mod.F90.

9.22.2.3 subroutine, public prg\_quantumdynamics\_mod::prg\_get\_sparsity\_realmat ( character(len=\*), intent(in) *matrix\_type*, character(len=\*), intent(in) *element\_type*, real(dp), intent(in) *thresh*, real(dp), dimension(:,:), intent(in) *a\_dense* )

This computes the sparsity of a real matrix given a threshold value This routine does:  $f=\frac{N_0}{N_{tot}}$  where f is the sparsity,  $N_0$  is the number of values less than the threshold, and  $N_{tot}$  is the total number of values. The sparsity and threshold are printed to the screen.

#### **Parameters**

matrix_type	the BML matrix type
element_type	the BML element type
thresh	the threshold for sparsity evaluation
a_dense	the dense real matrix to be evaluated for sparsity

Definition at line 122 of file prg\_quantumdynamics\_mod.F90.

9.22.2.4 subroutine, public prg\_quantumdynamics\_mod::prg\_getcharge ( type(bml\_matrix\_t), intent(in) rho\_bml, type(bml\_matrix\_t), intent(in) s\_bml, real(dp), dimension(:), allocatable charges, type(bml\_matrix\_t) aux\_bml, real(dp), dimension(:), intent(in), allocatable spindex, integer, dimension(:), intent(in), allocatable N, integer nats, real(dp), intent(in) thresh)

Constructs the charges from the density matrix.

#### **Parameters**

rho_bml	Density matrix in BML format.
over_bml	Overlap matrix in BML format.
charges	the array of charges.
aux_bml	the auxiliary matrix in BML format.
spindex	Start and end index for every atom in the system.
Z	
nats	the number of atoms
N	
thresh	threshold for the BML matrix

Definition at line 247 of file prg\_quantumdynamics\_mod.F90.

9.22.2.5 subroutine, public prg\_quantumdynamics\_mod::prg\_getdipole ( real(dp), dimension(:), intent(in) charges, real(dp), dimension(:,:), intent(in) r, real(dp), dimension(3), intent(inout) mu )

This routine computes the dipole moment of the system with units determined by the units of the coordinate matrix and charges given.

#### **Parameters**

charges	Charge on each atom.
r	Coordinate matrix of the atoms.
р	Dipole moment vector.

Definition at line 282 of file prg\_quantumdynamics\_mod.F90.

9.22.2.6 subroutine, public prg\_quantumdynamics\_mod::prg\_kick\_density ( integer, intent(in) kick\_direc, real(dp) kick\_mag, complex(dp), dimension(:,:), intent(inout), allocatable dens, integer, intent(in) norbs, integer, intent(in) mdim, complex(dp), dimension(:,:), allocatable SINV, integer, dimension(:), intent(in), allocatable which\_atom, real(dp), dimension(:,:), allocatable r, character(len=\*), intent(in) bmltype, real(dp) thresh)

Provides perturbation to initial density matrix in the form of an electric field kick. This routine does:  $\rho_{\hat{kick}} = \exp{\frac{-i}{\hbar}\hat{V}\hat{\rho}\hat{S}}\exp{\frac{i}{\hbar}\hat{V}\hat{S}^{-1}}$  where  $\hat{V}$  is the field disturbance.

#### **Parameters**

kick_direc	the direction of the kick in the electric field
kick_mag	the magnitude of the kick in the electric field
dens	the initial density matrix to be kicked.
norbs	the number of orbitals in the density matrix
S	the overlap matrix
SINV	the inverse of the overlap matrix
which_atom	vector containing atom identification
r	direction vector for kick based on atom and kick_direc
bmltype	type of BML matrix desired for faster computation
thresh	threshold for BML matrix conversion

Definition at line 43 of file prg\_quantumdynamics\_mod.F90.

9.22.2.7 subroutine, public prg\_quantumdynamics\_mod::prg\_kick\_density\_bml ( integer, intent(in) kick\_direc, real(dp) kick\_mag, type(bml\_matrix\_t) rho\_bml, type(bml\_matrix\_t) s\_bml, type(bml\_matrix\_t) sinv\_bml, integer mdim, integer, dimension(:), intent(in), allocatable which\_atom, real(dp), dimension(:,:), allocatable r, character(len=\*), intent(in) matrix\_type, real(dp) thresh)

Provides perturbation to initial density matrix in the form of an electric field kick given input matricies in BML format. This routine does:  $\rho_{\hat{k}ick} = \exp{\frac{-i}{\hbar}\hat{V}\hat{\rho}\hat{S}} \exp{\frac{i}{\hbar}\hat{V}\hat{S}^{-1}}$  where  $\hat{V}$  is the field disturbance.

#### **Parameters**

kick_direc	the direction of the kick in the electric field
kick_mag	the magnitude of the kick in the electric field
rho_bml	the initial density matrix to be kicked in BML format.
s_bml	the overlap matrix
sinv_bml	the inverse of the overlap matrix
mdim	maximum number of nonzero values per row in BML matrix
which_atom	vector containing atom identification
r	position vector for kicked atom
matrix_type	the type of BML format
thresh	the threshold for the BML matrix

Definition at line 154 of file prg\_quantumdynamics\_mod.F90.

9.22.2.8 subroutine, public prg\_quantumdynamics\_mod::prg\_lvni\_bml ( type(bml\_matrix\_t) h1\_bml, type(bml\_matrix\_t) sinv\_bml, real(dp) dt, real(dp) hbar, type(bml\_matrix\_t) rhoold\_bml, type(bml\_matrix\_t) rho\_bml, type(bml\_matrix\_t) aux\_bml, character(len=\*), intent(in) matrix\_type, integer mdim, real(dp), intent(in) thresh )

Performs Liouville-von Neumann integration using leap-frog method. This routine does:  $\hat{\rho}(t+\Delta t)=\hat{\rho}(t-\Delta t)+2\Delta t \frac{\partial \hat{\rho}(t)}{\partial t}$  where the time derivative of the density matrix is defined as follows:  $\frac{\partial \hat{\rho}(t)}{\partial t}=\frac{-i}{\hbar}\left(S^{-1}\hat{H}(t)\hat{\rho}(t)-\hat{\rho}(t)\hat{H}(t)S^{-1}\right)$ .

### **Parameters**

Н	the Hamiltonian matrix at time t
sinv_bml	the inverse overlap matrix
dt	the timestep for integration
hbar	the Dirac constant (generally taken to be 1 in simulation units)
rho_old	the density matrix at previous time-step
rho_bml	the density matrix at current time-step
aux_bml	the temp matrix used for value storage during computation
matrix_type	the type of BML matrix
thresh	the threshold for the BML matrix

Definition at line 211 of file prg\_quantumdynamics\_mod.F90.

#### 9.22.3 Variable Documentation

**9.22.3.1** integer, parameter prg\_quantumdynamics\_mod::dp = kind(1.0d0) [private]

Definition at line 14 of file prg\_quantumdynamics\_mod.F90.

# 9.23 prg\_response\_mod Module Reference

Module to compute the density matrix response and related quantities.

### **Data Types**

type respdata\_type

### **Functions/Subroutines**

• subroutine, public prg\_parse\_response (RespData, filename)

The parser for the calculation of the DM response.

• subroutine, public prg\_compute\_dipole (charges, coordinate, dipoleMoment, factor, verbose)

To compute the dipole moment of the system. The units of the dipole moment are determined by the units of the coordinates and charges that are given.

subroutine, public prg\_write\_dipole\_tcl (dipoleMoment, file, factor, verbose)

To visualize a dipole moment using VMD. This will prg\_generate a .tcl script that could be run using VMD To visualize with VMD: \$ vmd -e dipole.tcl.

• subroutine, public prg\_compute\_polarizability (rsp\_bml, prt\_bml, polarizability, factor, verbose)

To compute the polarizability of the system. The units of the directional polarizability are determined by the units of the perturbation and Hamiltonian. This equation can be found in [5] equation 4a. Note that in equation 4a of the reference there is a 2 that account for the double occupancy which is not present in this case cause the density matrix construction is done by taking the occupancy into account.

• subroutine, public prg\_pert\_from\_file (prt\_bml, norb)

Read perturbation from file.

subroutine, public prg\_compute\_response\_rs (ham\_bml, prt\_bml, rsp\_bml, lambda, bndfil, threshold, ver-bose)

Computes the first order response density matrix using Rayleigh Schrodinger Perturbation theory The transformation hereby performed are:

• subroutine, public prg\_compute\_response\_fd (ham\_bml, prt\_bml, rsp\_bml, prg\_delta, bndfil, threshold, verbose)

Computes the first order response density matrix using finite differences. The transformation hereby performed are:

• subroutine, public prg\_pert\_constant\_field (field, intensity, coordinate, lambda, prt\_bml, threshold, spindex, norbi, verbose, over\_bml)

Apply a constant field perturbation through the dipole moment operator (  $\hat{\mu}=e\hat{r}$ ). In the matrix representation, this is:  $H^{(1)}=\lambda \frac{1}{2}(S\,er\cdot {\bf E}+\,er\cdot {\bf E}S)$ . The symmetrization is done in order to preserve the Hermiticity of H. In this case the whole system will be affected by the field. In a latter version we will add the possibility of applying this field to a region of the system. In this implementation e=1 and units can be transformed by using the parameter  $\lambda$ .

• subroutine, public prg\_pert\_sin\_pot (direction, lx, coordinate, lambda, prt\_bml, threshold, spindex, norbi, verbose, over bml)

Apply a sinusoidal length dependent potential  $(\sin(\tilde{\textbf{r}}_x))$  where  $\textbf{r}_x$  is the x coordinate. The Hamiltonian gets modified as follows:  $H^{(1)} = \frac{1}{2}\lambda(S\sin(\tilde{\textbf{r}}_x) + \sin(\tilde{\textbf{r}}_x)S)$ .  $\tilde{\textbf{r}}_x = 2\pi(\textbf{r}/l_x) - \pi$ . The symmetrization is done in order to preserve the Hermiticity of H. Units can be transformed by using the parameter  $\lambda$ .

• subroutine, public prg\_pert\_cos\_pot (direction, lx, coordinate, lambda, prt\_bml, threshold, spindex, norbi, verbose, over bml)

Apply a cosine length dependent potential  $(\cos(\tilde{\textbf{r}}_x))$  where  $\textbf{r}_x$  is the x coordinate. The Hamiltonian gets modified as follows:  $H^{(1)} = \frac{1}{2}\lambda(S\sin(\tilde{\textbf{r}}_x) + \sin(\tilde{\textbf{r}}_x)S)$ .  $\tilde{\textbf{r}}_x = 2\pi(\textbf{r}/l_x) - \pi$ . The symmetrization is done in order to preserve the Hermiticity of H. Units can be transformed by using the parameter  $\lambda$ .

• subroutine, public prg\_compute\_response\_sp2 (ham\_bml, prt\_bml, rsp\_bml, rho\_bml, lambda, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, threshold, verbose)

Finds the first order response matrix from a Hamiltonian matrix.

• subroutine, public prg\_project\_response (rsp\_bml, over\_bml, spindex, norbi, coordinates, rspfunc, verbose) Project the response onto atomic positions. First order response to the perturbation ( $\rho^{(1)}$ ) projected onto the atomic position. Basically:  $rsp(i) = \sum_{\alpha \in i} \rho_{\alpha\alpha}^{(1)}$ , where orbital  $\alpha$  belong to atom i.

## **Variables**

- integer, parameter dp = kind(1.0d0)
- real(dp), parameter pi = 3.14159265358979323846264338327950 dp

## 9.23.1 Detailed Description

Module to compute the density matrix response and related quantities.

## Todo Add the response scf

Change name response\_SP2 to dm\_prt\_response

Change name response\_rs to rs\_prt\_response

More information about the theory can be found at [4] and Niklasson2015

# 9.23.2 Function/Subroutine Documentation

9.23.2.1 subroutine, public prg\_response\_mod::prg\_compute\_dipole ( real(dp), dimension(:), intent(in) charges, real(dp), dimension(:,:), intent(in) coordinate, real(dp), dimension(3), intent(inout) dipoleMoment, real(dp), intent(in) factor, integer verbose )

To compute the dipole moment of the system. The units of the dipole moment are determined by the units of the coordinates and charges that are given.

#### **Parameters**

charges	Charges on each atomic position.
coordinate	Coordinates of the atoms.
nats	Number of atoms.
dipoleMoment	Dipole moment vector.
factor	Unit conversion factor (use 1.0 is no conversion is required).
verbose	To give different verbosity levels. If coordinates are in and charges are in fractions of electron, then transformation ea2debye form LATTE lib can be used to change units to Debye.

Definition at line 120 of file prg\_response\_mod.F90.

9.23.2.2 subroutine, public prg\_response\_mod::prg\_compute\_polarizability ( type(bml\_matrix\_t), intent(in) rsp\_bml, type(bml\_matrix\_t), intent(in) prt\_bml, real(dp), intent(inout) polarizability, real(dp), intent(in) factor, integer verbose )

To compute the polarizability of the system. The units of the directional polarizability are determined by the units of the perturbation and Hamiltonian. This equation can be found in [5] equation 4a. Note that in equation 4a of the reference there is a 2 that account for the double occupancy which is not present in this case cause the density matrix construction is done by taking the occupancy into account.

#### **Parameters**

charges	Charges on each atomic position.
coordinate	Coordinates of the atoms.
nats	Number of atoms.
dipoleMoment	Dipole moment vector.
factor	Unit conversion factor (use 1.0 is no conversion is required).
verbose	To give different verbosity levels. If coordinates are in and charges are in fractions of electron, then transformation ea2debye form LATTE lib can be used to change units to Debye.

Definition at line 198 of file prg\_response\_mod.F90.

9.23.2.3 subroutine, public prg\_response\_mod::prg\_compute\_response\_fd ( type(bml\_matrix\_t), intent(in) ham\_bml, type(bml\_matrix\_t), intent(in) prt\_bml, type(bml\_matrix\_t), intent(inout) rsp\_bml, real(dp) prg\_delta, real(dp), intent(in) bndfil, real(dp), intent(in) threshold, integer verbose)

Computes the first order response density matrix using finite differences. The transformation hereby performed are:

- $H^+ = H^{(0)} + \delta H^{(1)}$
- $H^- = H^{(0)} \delta H^{(1)}$
- $\rho^+ = f(H^+)$
- $\rho^- = f(H^-)$
- $\rho^{(1)} = (\rho^+ \rho^-)/(2\delta)$ . Where f denotes the Fermi function (construction of the density matrix)

# Parameters

ham_bml	Hamiltonian in bml format ( $H^{\left(0\right)}$ ).
prt_bml	Perturbation in bml format ( $H^{(1)}$ ).
rsp_bml	First order response to the perturbation ( $\rho^{(1)}$ ).
bndfil	Filing factor.
threshold	Threshold value for matrix elements.
verbose	Different levels of verbosity.

# Warning

This works only for the prg\_orthogonalized form of ham\_bml. The response must be in the prg\_orthogonalized form.

Definition at line 379 of file prg\_response\_mod.F90.

Here is the call graph for this function:



9.23.2.4 subroutine, public prg\_response\_mod::prg\_compute\_response\_rs ( type(bml\_matrix\_t), intent(in) ham\_bml, type(bml\_matrix\_t), intent(in) prt\_bml, type(bml\_matrix\_t), intent(inout) rsp\_bml, real(dp) lambda, real(dp), intent(in) bndfil, real(dp), intent(in) threshold, integer verbose )

Computes the first order response density matrix using Rayleigh Schrodinger Perturbation theory The transformation hereby performed are:

- $V = C^{\dagger}H^{(1)}C$
- $\tilde{V}_{ij} = \frac{V_{ij}}{\epsilon_i \epsilon_i}$ , with  $\tilde{V}_{ii} = 0 \, \forall i$ .
- $C^{(1)} = C\tilde{V}$
- And finally:  $ho^{(1)}=Cf(C^{(1)})^\dagger+C^{(1)}fC^\dagger$

#### **Parameters**

ham_bml	Hamiltonian in bml format ( $H^{(0)}$ ).
prt_bml	Perturbation in bml format ( $H^{(1)}$ ).
rsp_bml	First order response to the perturbation ( $ ho^{(1)}$ ).
bndfil	Filing factor.
threshold	Threshold value for matrix elements.
verbose	Different levels of verbosity.

# Warning

This works only for the prg\_orthogonalized form of ham\_bml. The response must be in the prg\_orthogonalized form.

Definition at line 249 of file prg response mod.F90.

9.23.2.5 subroutine, public prg\_response\_mod::prg\_compute\_response\_sp2 ( type(bml\_matrix\_t), intent(in) ham\_bml, type(bml\_matrix\_t), intent(in) prt\_bml, type(bml\_matrix\_t), intent(inout) rsp\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp) lambda, real(dp), intent(in) bndfil, integer, intent(in) minsp2iter, integer, intent(in) massp2iter, character(len=\*), intent(in) sp2conv, real(dp), intent(in) idemtol, real(dp), intent(in) threshold, integer verbose)

Finds the first order response matrix from a Hamiltonian matrix.

Definition at line 652 of file prg\_response\_mod.F90.

9.23.2.6 subroutine, public prg\_response\_mod::prg\_parse\_response ( type(respdata\_type) RespData, character(len=\*) filename )

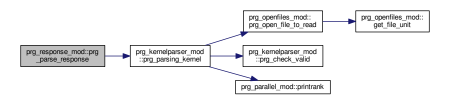
The parser for the calculation of the DM response.

#### **Parameters**

RespData	Response data type.
filename	Name of the file to parse.

Definition at line 45 of file prg\_response\_mod.F90.

Here is the call graph for this function:



9.23.2.7 subroutine, public prg\_response\_mod::prg\_pert\_constant\_field ( real(dp), dimension(3), intent(in) field, real(dp) intensity, real(dp), dimension(:,:), intent(in) coordinate, real(dp) lambda, type(bml\_matrix\_t), intent(inout) prt\_bml, real(dp) threshold, integer, dimension(:), intent(in) spindex, integer, dimension(:), intent(in) norbi, integer, intent(in) verbose, type(bml\_matrix\_t), intent(in), optional over\_bml)

Apply a constant field perturbation through the dipole moment operator (  $\hat{\mu}=e\hat{\bf r}$ ). In the matrix representation, this is:  $H^{(1)}=\lambda \frac{1}{2}(\, S\, e{\bf r}\cdot{\bf E}+\, e{\bf r}\cdot{\bf E}S)$ . The symmetrization is done in order to preserve the Hermiticity of H. In this case the whole system will be affected by the field. In a latter version we will add the possibility of applying this field to a region of the system. In this implementation e=1 and units can be transformed by using the parameter  $\lambda$ .

### Note

If the Hamiltonian is already in the prg orthogonalized form, then parameter over bml can be omitted.

#### **Parameters**

field	Direction of the applied field ( $\hat{\mathbf{E}}$ ).
intensity	Intensity of the field ( $  \mathbf{E}  $ )
coordinate	Coordinates of the system ( r).
lambda	Constant to premultiply the perturbation ( $\lambda$ ).
prt_bml	Perturbation in bml format ( $H^{(1)}$ ).
threshold	Threshold value for bml format matrices.
spindex	Species index. It gives the species index of a particular atom.
norbi	Number of orbitals for each atomic site.
verbose	Different levels of verbosity.
over_bml	It has to be present for a nonorthogonal representation ( $S$ ).

Definition at line 446 of file prg\_response\_mod.F90.

9.23.2.8 subroutine, public prg\_response\_mod::prg\_pert\_cos\_pot ( character *direction*, real(dp) *lx*, real(dp), dimension(:,:), intent(in) *coordinate*, real(dp) *lambda*, type(bml\_matrix\_t), intent(inout) *prt\_bml*, real(dp) *threshold*, integer, dimension(:), intent(in) *spindex*, integer, dimension(:), intent(in) *norbi*, integer, intent(in) *verbose*, type(bml\_matrix\_t), intent(in), optional *over\_bml*)

Apply a cosine length dependent potential ( $\cos(\tilde{\mathbf{r}}_x)$ ) where  $\mathbf{r}_x$  is the x coordinate. The Hamiltonian gets modified as follows:  $H^{(1)} = \frac{1}{2}\lambda(S\sin(\tilde{\mathbf{r}}_x) + \sin(\tilde{\mathbf{r}}_x)S)$ .  $\tilde{\mathbf{r}}_x = 2\pi(\mathbf{r}/l_x) - \pi$ . The symmetrization is done in order to preserve the Hermiticity of H. Units can be transformed by using the parameter  $\lambda$ .

#### Note

If the Hamiltonian is already in the prg orthogonalized form, then parameter over bml can be omitted.

#### **Parameters**

direction	Direction of the potential gradient (x,y or z).
lx	Lenght of the box in x direction.
coordinate	Coordinates of the system ( r).
lambda	Constant to premultiply the perturbation ( $\lambda$ ).
prt_bml	Perturbation in bml format ( $H^{(1)}$ ).
threshold	Threshold value for bml format matrices.
norbi	Number of orbitals for each atomic site.
verbose	Different levels of verbosity.
over_bml	It has to be present for a nonorthogonal representation ( $S$ ).

Definition at line 591 of file prg\_response\_mod.F90.

9.23.2.9 subroutine, public prg\_response\_mod::prg\_pert\_from\_file ( type(bml\_matrix\_t), intent(inout) prt\_bml, integer norb )

Read perturbation from file.

Todo Add read perturbation from file

Definition at line 223 of file prg\_response\_mod.F90.

9.23.2.10 subroutine, public prg\_response\_mod::prg\_pert\_sin\_pot ( character *direction*, real(dp) *lx*, real(dp), dimension(:,:), intent(in) *coordinate*, real(dp) *lambda*, type(bml\_matrix\_t), intent(inout) *prt\_bml*, real(dp) *threshold*, integer, dimension(:), intent(in) *spindex*, integer, dimension(:), intent(in) *norbi*, integer, intent(in) *verbose*, type(bml\_matrix\_t), intent(in), optional *over\_bml*)

Apply a sinusoidal length dependent potential (  $\sin(\tilde{\mathbf{r}}_x)$ ) where  $\mathbf{r}_x$  is the x coordinate. The Hamiltonian gets modified as follows:  $H^{(1)} = \frac{1}{2}\lambda(S\sin(\tilde{\mathbf{r}}_x) + \sin(\tilde{\mathbf{r}}_x)S)$ .  $\tilde{\mathbf{r}}_x = 2\pi(\mathbf{r}/l_x) - \pi$ . The symmetrization is done in order to preserve the Hermiticity of H. Units can be transformed by using the parameter  $\lambda$ .

### Note

If the Hamiltonian is already in the prg\_orthogonalized form, then parameter over\_bml can be omitted.

#### **Parameters**

direction	Direction of the potential gradient (x,y or z).
lx	Length of the box in x direction.
coordinate	Coordinates of the system ( r).
lambda	Constant to premultiply the perturbation ( $\lambda$ ).
prt_bml	Perturbation in bml format ( $H^{(1)}$ ).
threshold	Threshold value for bml format matrices.
norbi	Number of orbitals for each atomic site.
verbose	Different levels of verbosity.
over_bml	It has to be present for a nonorthogonal representation ( $S$ ).

Definition at line 523 of file prg\_response\_mod.F90.

9.23.2.11 subroutine, public prg\_response\_mod::prg\_project\_response ( type(bml\_matrix\_t), intent(inout) rsp\_bml, type(bml\_matrix\_t), intent(in) over\_bml, integer, dimension(:), intent(in) spindex, integer, dimension(:), intent(in) norbi, real(dp), dimension(:,:), intent(in) coordinates, real(dp), dimension(:), intent(inout), allocatable rspfunc, integer, intent(in) verbose)

Project the response onto atomic positions. First order response to the perturbation (  $\rho^{(1)}$ ) projected onto the atomic position. Basically:  $rsp(i) = \sum_{\alpha \in i} \rho_{\alpha\alpha}^{(1)}$ , where orbital  $\alpha$  belong to atom i.

### **Parameters**

rsp_bml	First order response density matrix.
spindex	It gives the species index of a particular atom.
norbi	Number of orbitals of species i.
coordinates	Atomic coordinates.
rspfunc	Response function at atomic positions.
verbose	Different levels of verbosity.

Definition at line 795 of file prg\_response\_mod.F90.

9.23.2.12 subroutine, public prg\_response\_mod::prg\_write\_dipole\_tcl ( real(dp), dimension(3), intent(in) dipoleMoment, character(\*), intent(in) file, real(dp), intent(in) factor, integer verbose )

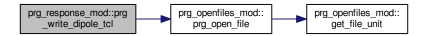
To visualize a dipole moment using VMD. This will prg\_generate a .tcl script that could be run using VMD To visualize with VMD: \$ vmd -e dipole.tcl.

### **Parameters**

dipoleMoment	Dipole moment vector.
file	PDB/XYZ file to load for visualization.
factor	Arbitrary scale for visualization.
verbose	To give different verbosity levels.

Definition at line 158 of file prg\_response\_mod.F90.

Here is the call graph for this function:



#### 9.23.3 Variable Documentation

9.23.3.1 integer, parameter prg\_response\_mod::dp = kind(1.0d0) [private]

Definition at line 18 of file prg\_response\_mod.F90.

Definition at line 19 of file prg\_response\_mod.F90.

# 9.24 prg\_sp2\_fermi\_mod Module Reference

The SP2 Fermi module.

## **Functions/Subroutines**

subroutine, public prg\_sp2\_fermi\_init (h\_bml, nsteps, nocc, tscale, threshold, occErrLimit, traceLimit, x\_bml, mu, beta, h1, hN, sgnlist)

Truncated SP2 prg\_initialization.

• subroutine, public prg\_sp2\_fermi\_init\_norecs (h\_bml, nsteps, nocc, tscale, threshold, occErrLimit, traceLimit, x bml, mu, beta, h1, hN, sgnlist, verbose)

Truncated SP2 prg\_initialization. This routine also gives back the Number of SP2 recursive steps that gets a Pseudo-Fermi distribution with a temperature close to the target temperature which is entered using parameter beta = (1/KbT).

• subroutine, public prg\_sp2\_fermi (h\_bml, osteps, nsteps, nocc, mu, beta, h1, hN, sgnlist, threshold, eps, traceLimit, x\_bml)

Calculate Truncated SP2.

• subroutine, public prg\_sp2\_entropy\_function (mu, h1, hN, nsteps, sgnlist, GG, ee)

Calculate SP2 entropy function using gaussian quadrature. Note that GG and ee are allocated and returned from this routine.

• real(dp) function, public sp2 entropy ts (D0 bml, GG, ee)

Test SP2 entropy. Get the entropy contribution TS to the total free energy.

real(dp) function, public sp2\_inverse (f, mu, h1, hN, nsteps, sgnlist)

Calculate the SP2 inverse.

• real(dp) function absmaxderivative (func, de)

Gets the absolute maximum of the derivative of a function.

## Variables

• integer, parameter dp = kind(1.0d0)

# 9.24.1 Detailed Description

The SP2 Fermi module.

This subroutine implements Niklasson's truncated SP2 density matrix purification algorithm.

## 9.24.2 Function/Subroutine Documentation

9.24.2.1 real(dp) function prg\_sp2\_fermi\_mod::absmaxderivative ( real(dp), dimension(:), intent(in) func, real(dp), intent(in) de
) [private]

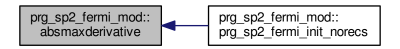
Gets the absolute maximum of the derivative of a function.

#### **Parameters**

func.	
de	Energy step.

Definition at line 614 of file prg\_sp2\_fermi\_mod.F90.

Here is the caller graph for this function:



9.24.2.2 subroutine, public prg\_sp2\_fermi\_mod::prg\_sp2\_entropy\_function ( real(dp), intent(in) *mu*, real(dp), intent(in) *h1*, real(dp), intent(in) *hN*, integer, intent(in) *nsteps*, integer, dimension(:), intent(in) *sgnlist*, real(dp), dimension(:), intent(inout), allocatable *ee* )

Calculate SP2 entropy function using gaussian quadrature. Note that GG and ee are allocated and returned from this routine.

#### **Parameters**

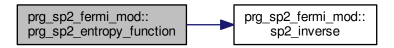
ти	Shifted chemical potential
h1	Minimum scaled Gershgorin bound
hN	Maximum scaled Gershgorin bound

### **Parameters**

nsteps	Number of SP2 steps
sgnlist	SP2 sequence
GG	Entropy function
ee	1D mesh

Definition at line 479 of file prg\_sp2\_fermi\_mod.F90.

Here is the call graph for this function:



9.24.2.3 subroutine, public prg\_sp2\_fermi\_mod::prg\_sp2\_fermi ( type(bml\_matrix\_t), intent(in) h\_bml, integer, intent(in) osteps, integer, intent(in) nsteps, real(dp), intent(in) nocc, real(dp), intent(inout) mu, real(dp), intent(inout) beta, real(dp), intent(inout) h1, real(dp), intent(inout) hN, integer, dimension(:), intent(in) sgnlist, real(dp), intent(in) threshold, real(dp), intent(in) eps, real(dp), intent(in) traceLimit, type(bml\_matrix\_t), intent(inout) x\_bml)

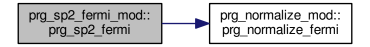
Calculate Truncated SP2.

# Parameters

h_bml	Hamiltonian matrix
osteps	Outer loop steps
nsteps	Number of sequence branches
посс	Number of occupation states
ти	Shifted chemical potential
beta	Inverse temperature
h1	Minimum scaled Gershgorin bound.
hN	Maximum scaled Gershgorin bound.
sgnlist	SP2 sequence
threshold	Threshold for multiplies
eps	Occupation error limit
traceLimit	Trace limit
x_bml	Output density matrix

Definition at line 386 of file prg\_sp2\_fermi\_mod.F90.

Here is the call graph for this function:



9.24.2.4 subroutine, public prg\_sp2\_fermi\_mod::prg\_sp2\_fermi\_init ( type(bml\_matrix\_t), intent(in) h\_bml, integer, intent(in) nsteps, real(dp), intent(in) nocc, real(dp), intent(in) tscale, real(dp), intent(in) threshold, real(dp), intent(in) occErrLimit, real(dp), intent(in) traceLimit, type(bml\_matrix\_t), intent(inout) x\_bml, real(dp), intent(inout) mu, real(dp), intent(inout) beta, real(dp), intent(inout) h1, real(dp), intent(inout) hN, integer, dimension(:), intent(inout) sgnlist)

Truncated SP2 prg\_initialization.

#### **Parameters**

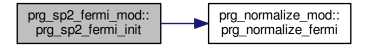
h_bml	Input Hamiltonian matrix.
nsteps	Number of sp2 iterations.
nocc	Number of occupied states.
tscale	Temperature rescaling factor.
threshold	Threshold for multiplication.
occErrLimit	Occupation error limit.
traceLimit	Trace limit.
x_bml	Output prg_initial matrix.
mu	Shifted chemical potential
beta	Output inverse temperature.
h1	Output temperature-scaled minimum gershgorin bound.
hN	Output temperature-scaled maximum gershgorin bound.
sgnlist	SP2 sequence

Calculate Gershgorin bounds and rescale

Determine sequence branching first time through

Definition at line 45 of file prg\_sp2\_fermi\_mod.F90.

Here is the call graph for this function:



9.24.2.5 subroutine, public prg\_sp2\_fermi\_mod::prg\_sp2\_fermi\_init\_norecs ( type(bml\_matrix\_t), intent(in) h\_bml, integer, intent(inout) nsteps, real(dp), intent(in) nocc, real(dp), intent(in) tscale, real(dp), intent(in) threshold, real(dp), intent(in) occErrLimit, real(dp), intent(in) traceLimit, type(bml\_matrix\_t), intent(inout) x\_bml, real(dp), intent(inout) mu, real(dp), intent(inout) beta, real(dp), intent(inout) h1, real(dp), intent(inout) hN, integer, dimension(:), intent(inout) sgnlist, integer, optional verbose)

Truncated SP2 prg\_initialization. This routine also gives back the Number of SP2 recursive steps that gets a Pseudo-Fermi distribution with a temperature close to the target temperature which is entered using parameter beta = (1/KbT).

### **Parameters**

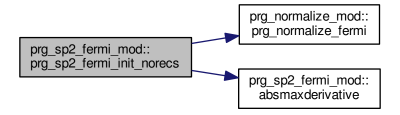
h_bml	Input Hamiltonian matrix.
nsteps	Output number of sp2 iterations.
nocc	Number of occupied states.
tscale	Temperature rescaling factor.
threshold	Threshold for multiplication.
occErrLimit	Occupation error limit.
traceLimit	Trace limit.
x_bml	Output prg_initial matrix.
mu	Shifted chemical potential
beta	Input guess and output inverse temperature.
h1	Output temperature-scaled minimum gershgorin bound.
hN	Output temperature-scaled maximum gershgorin bound.
sgnlist	SP2 sequence
verbose	Optional parameter for verbosity.

Calculate Gershgorin bounds and rescale

Determine sequence branching first time through

Definition at line 200 of file prg\_sp2\_fermi\_mod.F90.

Here is the call graph for this function:



9.24.2.6 real(dp) function, public prg\_sp2\_fermi\_mod::sp2\_entropy\_ts ( type(bml\_matrix\_t), intent(in) D0\_bml, real(dp), dimension(\*), intent(in) GG, real(dp), dimension(\*), intent(in) ee )

Test SP2 entropy. Get the entropy contribution TS to the total free energy.

#### **Parameters**

D0_bml	BML matrix
GG	Entropy function
ee	1D mesh
TS	Energy contribution

Definition at line 537 of file prg\_sp2\_fermi\_mod.F90.

9.24.2.7 real(dp) function, public prg\_sp2\_fermi\_mod::sp2\_inverse ( real(dp), intent(in) f, real(dp), intent(in) mu, real(dp), intent(in) h1, real(dp), intent(in) hN, integer, intent(in) nsteps, integer, dimension(:), intent(in) sgnlist)

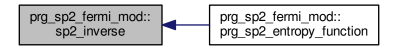
Calculate the SP2 inverse.

# **Parameters**

f	Occupation factor	
ти	Shifted chemical potential	
h1	Minimum scaled Gershgorin bound	
hN	Maximum scaled Gershgorin bound	
nsteps	Numbers of SP2 iterations	
sgnlist	SP2 sequence	
ee	Energy value	

Definition at line 589 of file prg\_sp2\_fermi\_mod.F90.

Here is the caller graph for this function:



### 9.24.3 Variable Documentation

**9.24.3.1** integer, parameter prg\_sp2\_fermi\_mod::dp = kind(1.0d0) [private]

Definition at line 18 of file prg\_sp2\_fermi\_mod.F90.

# 9.25 prg\_sp2\_mod Module Reference

The SP2 module.

# **Functions/Subroutines**

• subroutine, public prg\_sp2\_basic (h\_bml, rho\_bml, threshold, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, verbose)

Calculates the density matrix from a Hamiltonian matrix by purification. The method implemented here is the very first verion of the SP2 method.

- subroutine, public prg\_sp2\_alg2 (h\_bml, rho\_bml, threshold, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, verbose)
- subroutine, public prg\_sp2\_alg2\_genseq (h\_bml, rho\_bml, threshold, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, pp, icount, vv, verbose)
- subroutine, public prg\_sp2\_alg2\_seq (h\_bml, rho\_bml, threshold, pp, icount, vv, verbose)
- subroutine, public prg\_prg\_sp2\_alg2\_seq\_inplace (rho\_bml, threshold, pp, icount, vv, mineval, maxeval, verbose)
- subroutine, public prg\_sp2\_alg1 (h\_bml, rho\_bml, threshold, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, verbose)
- subroutine, public prg\_sp2\_alg1\_genseq (h\_bml, rho\_bml, threshold, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, pp, icount, vv)
- subroutine, public prg\_sp2\_alg1\_seq (h\_bml, rho\_bml, threshold, pp, icount, vv)
- subroutine, public prg\_prg\_sp2\_alg1\_seq\_inplace (rho\_bml, threshold, pp, icount, vv, mineval, maxeval)
- subroutine, public prg\_sp2\_submatrix (ham\_bml, rho\_bml, threshold, pp, icount, vv, mineval, maxeval, core
   —size)

Perform SP2 algorithm using sequence and calculate norm for a submatrix.

subroutine, public prg\_sp2\_submatrix\_inplace (rho\_bml, threshold, pp, icount, vv, mineval, maxeval, core\_
 size)

**Variables** 

• integer, parameter dp = kind(1.0d0)

## 9.25.1 Detailed Description

The SP2 module.

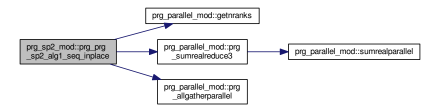
This subroutine implements Niklasson's SP2 density matrix purification algorithm.

#### 9.25.2 Function/Subroutine Documentation

9.25.2.1 subroutine, public prg\_sp2\_mod::prg\_prg\_sp2\_alg1\_seq\_inplace (type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp), intent(in) threshold, integer, dimension(:), intent(inout) pp, integer, intent(inout) icount, real(dp), dimension(:), intent(inout) vv, real(dp), intent(in) mineval, real(dp), intent(in) maxeval)

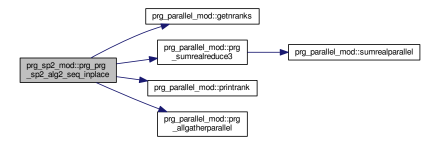
Definition at line 968 of file prg sp2 mod.F90.

Here is the call graph for this function:



9.25.2.2 subroutine, public prg\_sp2\_mod::prg\_prg\_sp2\_alg2\_seq\_inplace (type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp), intent(in) threshold, integer, dimension(:), intent(inout) pp, integer, intent(inout) icount, real(dp), dimension(:), intent(inout) vv, real(dp), intent(in), optional mineval, real(dp), intent(in), optional maxeval, integer, intent(in), optional verbose)

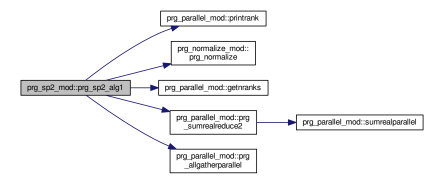
Definition at line 525 of file prg\_sp2\_mod.F90.



9.25.2.3 subroutine, public prg\_sp2\_mod::prg\_sp2\_alg1 ( type(bml\_matrix\_t), intent(in) h\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp), intent(in) threshold, real(dp), intent(in) bndfil, integer, intent(in) minsp2iter, integer, intent(in) maxsp2iter, character(len=\*), intent(in) sp2conv, real(dp), intent(in) idemtol, integer, intent(in), optional verbose )

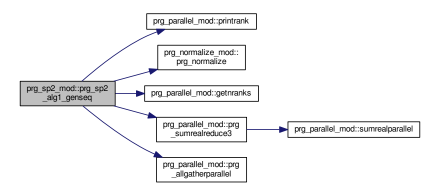
Definition at line 621 of file prg\_sp2\_mod.F90.

Here is the call graph for this function:



9.25.2.4 subroutine, public prg\_sp2\_mod::prg\_sp2\_alg1\_genseq ( type(bml\_matrix\_t), intent(in) h\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp), intent(in) threshold, real(dp), intent(in) bndfil, integer, intent(in) minsp2iter, integer, intent(in) maxsp2iter, character(len=\*), intent(in) sp2conv, real(dp), intent(in) idemtol, integer, dimension(:), intent(inout) pp, integer, intent(inout) icount, real(dp), dimension(:), intent(inout) vv)

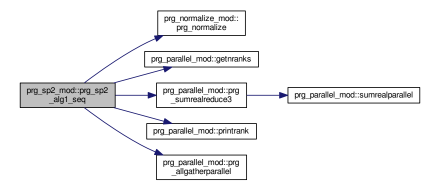
Definition at line 748 of file prg\_sp2\_mod.F90.



9.25.2.5 subroutine, public prg\_sp2\_mod::prg\_sp2\_alg1\_seq ( type(bml\_matrix\_t), intent(in) h\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp), intent(in) threshold, integer, dimension(:), intent(inout) pp, integer, intent(inout) icount, real(dp), dimension(:), intent(inout) vv )

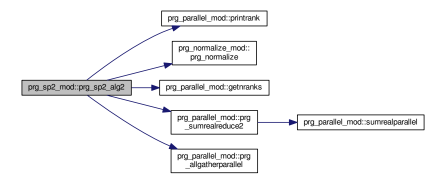
Definition at line 877 of file prg\_sp2\_mod.F90.

Here is the call graph for this function:



9.25.2.6 subroutine, public prg\_sp2\_mod::prg\_sp2\_alg2 ( type(bml\_matrix\_t), intent(in) h\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp), intent(in) threshold, real(dp), intent(in) bndfil, integer, intent(in) minsp2iter, integer, intent(in) maxsp2iter, character(len=\*), intent(in) sp2conv, real(dp), intent(in) idemtol, integer, intent(in), optional verbose )

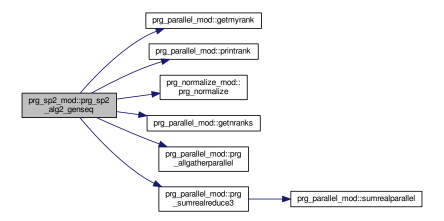
Definition at line 150 of file prg\_sp2\_mod.F90.



9.25.2.7 subroutine, public prg\_sp2\_mod::prg\_sp2\_alg2\_genseq ( type(bml\_matrix\_t), intent(in) h\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp), intent(in) threshold, real(dp), intent(in) bndfil, integer, intent(in) minsp2iter, integer, intent(in) maxsp2iter, character(len=\*), intent(in) sp2conv, real(dp), intent(in) idemtol, integer, dimension(:), intent(inout) pp, integer, intent(inout) icount, real(dp), dimension(:), intent(inout) vv, integer, intent(in), optional verbose)

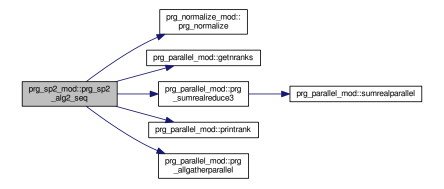
Definition at line 282 of file prg sp2 mod.F90.

Here is the call graph for this function:



9.25.2.8 subroutine, public prg\_sp2\_mod::prg\_sp2\_alg2\_seq ( type(bml\_matrix\_t), intent(in) h\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp), intent(in) threshold, integer, dimension(:), intent(inout) pp, integer, intent(inout) icount, real(dp), dimension(:), intent(inout) vv, integer, intent(in), optional verbose)

Definition at line 429 of file prg\_sp2\_mod.F90.



9.25.2.9 subroutine, public prg\_sp2\_mod::prg\_sp2\_basic ( type(bml\_matrix\_t), intent(in) h\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp) threshold, real(dp) bndfil, integer minsp2iter, integer maxsp2iter, character(len=\*), intent(in) sp2conv, real(dp) idemtol, integer verbose )

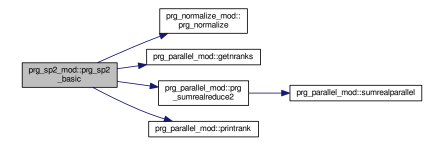
Calculates the density matrix from a Hamiltonian matrix by purification. The method implemented here is the very first verion of the SP2 method.

#### **Parameters**

h_bml	Input Hamiltonian matrix
rho_bml	Output density matrix
threshold	Threshold for sparse matrix algebra
bndfil	Bond
minsp2iter	Minimum sp2 iterations
maxsp2iter	Maximum SP2 iterations
sp2conv	Convergence type
idemtol	Idempotency tolerance
verbose	A verbosity level

Definition at line 49 of file prg sp2 mod.F90.

Here is the call graph for this function:



9.25.2.10 subroutine, public prg\_sp2\_mod::prg\_sp2\_submatrix ( type(bml\_matrix\_t), intent(in) ham\_bml, type(bml\_matrix\_t), intent(inout) rho\_bml, real(dp), intent(in) threshold, integer, dimension(:), intent(in) pp, integer, intent(in) icount, real(dp), dimension(:), intent(inout) vv, real(dp), intent(in) mineval, real(dp), intent(in) maxeval, integer, intent(in) core\_size)

Perform SP2 algorithm using sequence and calculate norm for a submatrix.

### **Parameters**

rho_bml	Input Hamiltonian/Output density matrix
threshold	Threshold for sparse matrix algebra
рр	Vector containing sequence of 0s and 1s
icount	Sequence count
VV	Vector of sum of squares per iteration
mineval	Min value used for normalization (optional)
Gemerated by Do	ky ម្នាំក្នុងx value used for normalization (optional)
core_size	Number of core rows

Definition at line 1058 of file prg\_sp2\_mod.F90.

9.25.2.11 subroutine, public prg\_sp2\_mod::prg\_sp2\_submatrix\_inplace ( type(bml\_matrix\_t), intent(inout) *rho\_bml*, real(dp), intent(in) *threshold*, integer, dimension(:), intent(inout) *pp*, integer, intent(inout) *icount*, real(dp), dimension(:), intent(inout) *vv*, real(dp), intent(in) *mineval*, real(dp), intent(in) *maxeval*, integer, intent(in) *core\_size* )

Definition at line 1127 of file prg\_sp2\_mod.F90.

Here is the caller graph for this function:



### 9.25.3 Variable Documentation

9.25.3.1 integer, parameter prg\_sp2\_mod::dp = kind(1.0d0) [private]

Definition at line 18 of file prg\_sp2\_mod.F90.

# 9.26 prg\_sp2parser\_mod Module Reference

SP2 parser.

# **Data Types**

• type sp2data\_type

General SP2 solver type.

# **Functions/Subroutines**

subroutine, public prg\_parse\_sp2 (sp2data, filename)
 The parser for SP2 solver.

### **Variables**

integer, parameter dp = kind(1.0d0)

## 9.26.1 Detailed Description

### SP2 parser.

This module is used to parse all the input variables for the SP2 method electronic structure solver. Adding a new input keyword to the parser:

- If the variable is real, we have to increase nkey re.
- Add the keyword (character type) in the keyvector\_re vector.
- Add a default value (real type) in the valvector\_re.
- Define a new variable and pass the value through valvector\_re(num) where num is the position of the new keyword in the vector.

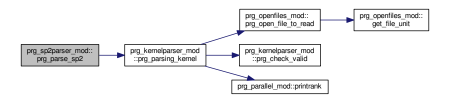
### 9.26.2 Function/Subroutine Documentation

9.26.2.1 subroutine, public prg\_sp2parser\_mod::prg\_parse\_sp2 ( type(sp2data\_type), intent(inout) sp2data, character(len=\*) filename )

The parser for SP2 solver.

Definition at line 50 of file prg\_sp2parser\_mod.F90.

Here is the call graph for this function:



### 9.26.3 Variable Documentation

**9.26.3.1** integer, parameter prg\_sp2parser\_mod::dp = kind(1.0d0) [private]

Definition at line 22 of file prg\_sp2parser\_mod.F90.

# 9.27 prg\_subgraphloop\_mod Module Reference

The subgraphloop module.

## **Functions/Subroutines**

- subroutine, public prg\_subgraphsp2loop (h\_bml, g\_bml, rho\_bml, gp, threshold)
- subroutine, public prg\_collectmatrixfromparts (gp, rho\_bml)

Collect distributed parts into same matrix.

- subroutine, public prg\_balanceparts (gp)
- subroutine, public prg\_partordering (gp)

Set row ordering bases on parts.

• subroutine, public prg\_getgrouppartitionhalosfromgraph (gp, g\_bml, hnode, djflag)

Get core+halo indeces for all partitions only using the graph.

• subroutine, public prg\_getpartitionhalosfromgraph (gp, g\_bml, djflag)

Get core+halo indeces for all partitions only using the graph.

## **Variables**

• integer, parameter dp = kind(1.0d0)

## 9.27.1 Detailed Description

The subgraphloop module.

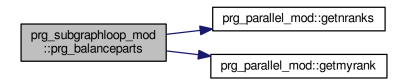
## 9.27.2 Function/Subroutine Documentation

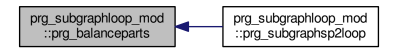
9.27.2.1 subroutine, public prg\_subgraphloop\_mod::prg\_balanceparts (type (graph\_partitioning\_t), intent(inout) gp)

Renumber parts Handle unbalanced numbers of parts.

Definition at line 165 of file prg\_subgraphloop\_mod.F90.

Here is the call graph for this function:





9.27.2.2 subroutine, public prg\_subgraphloop\_mod::prg\_collectmatrixfromparts ( type (graph\_partitioning\_t), intent(inout) gp, type (bml\_matrix\_t), intent(inout) rho\_bml )

Collect distributed parts into same matrix.

### **Parameters**

gp	Graph partitioning
rho_bml	Matrix to be collected into

Definition at line 133 of file prg\_subgraphloop\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



9.27.2.3 subroutine, public prg\_subgraphloop\_mod::prg\_getgrouppartitionhalosfromgraph ( type (graph\_partitioning\_t), intent(inout) *gp,* type (bml\_matrix\_t), intent(in) *g\_bml,* integer, dimension(\*), intent(in) *hnode,* logical, intent(in) *djflag* )

Get core+halo indeces for all partitions only using the graph.

### **Parameters**

gp	Graph partitioning
g_bml	Graph
hnode	Group start indeces
djflg	Double jump flag (true/false)

Determine halo elements for each subgraph

Definition at line 292 of file prg\_subgraphloop\_mod.F90.

9.27.2.4 subroutine, public prg\_subgraphloop\_mod::prg\_getpartitionhalosfromgraph ( type (graph\_partitioning\_t), intent(inout) *gp,* type (bml\_matrix\_t), intent(in) *g\_bml,* logical, intent(in) *djflag* )

Get core+halo indeces for all partitions only using the graph.

#### **Parameters**

gp	Graph partitioning
g_bml	Graph
djflg	Double jump flag (true/false)

Determine halo elements for each subgraph

Definition at line 337 of file prg\_subgraphloop\_mod.F90.

9.27.2.5 subroutine, public prg\_subgraphloop\_mod::prg\_partordering (type (graph\_partitioning\_t), intent(inout) gp)

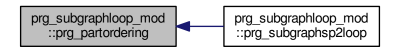
Set row ordering bases on parts.

### **Parameters**

gp	Graph partitioning
----	--------------------

Definition at line 263 of file prg\_subgraphloop\_mod.F90.

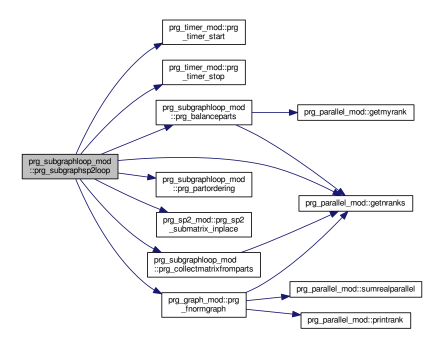
Here is the caller graph for this function:



9.27.2.6 subroutine, public prg\_subgraphloop\_mod::prg\_subgraphsp2loop ( type (bml\_matrix\_t), intent(in) h\_bml, type (bml\_matrix\_t), intent(in) g\_bml, type (bml\_matrix\_t), intent(inout) rho\_bml, type (graph\_partitioning\_t), intent(inout) gp, real(dp), intent(in) threshold )

Definition at line 37 of file prg\_subgraphloop\_mod.F90.

Here is the call graph for this function:



## 9.27.3 Variable Documentation

**9.27.3.1** integer, parameter prg\_subgraphloop\_mod::dp = kind(1.0d0) [private]

Definition at line 18 of file prg\_subgraphloop\_mod.F90.

# 9.28 prg\_syrotation\_mod Module Reference

A module to rotate the coordinates of a sybsystem in chemical systems.

# **Data Types**

type rotation\_type
 Rotation type.

## **Functions/Subroutines**

- subroutine, public prg\_parse\_rotation (rot, filename)

  The parser for rotation.
- subroutine, public prg\_rotate (rot, r, verbose)
   Rotation routine.

#### **Variables**

• integer, parameter dp = kind(1.0d0)

## 9.28.1 Detailed Description

A module to rotate the coordinates of a sybsystem in chemical systems.

It works by specifying two orientations and a rotation point.

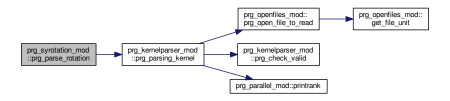
## 9.28.2 Function/Subroutine Documentation

9.28.2.1 subroutine, public prg\_syrotation\_mod::prg\_parse\_rotation ( type(rotation\_type), intent(inout) rot, character(len=\*) filename )

The parser for rotation.

Definition at line 46 of file prg\_syrotation\_mod.F90.

Here is the call graph for this function:



9.28.2.2 subroutine, public prg\_syrotation\_mod::prg\_rotate ( type(rotation\_type), intent(in) *rot*, real(dp), dimension(:,:), intent(inout) *r*, integer, intent(in) *verbose* )

Rotation routine.

It works by indicating the orientations (v1 and v1) and a rotation center. The orientation can be passed either directly by setting v1 and v2 or by indicating two points pQ1 and pQ2. Orientation can also be specified with an atom position if patom1 and patom2 indices are not zero this atoms are used to determine the initial and final orientation.

#### **Parameters**

rot	Rotation type
r	Coordinates to be rotated
verbose	Verbosity level

## Example:

```
rot%patom1 = 4
rot%patom2 = 0
rot%catom2 = 6
rot%v2 = 0.0; rot%v2(1) = 1
call prg_rotate(rot,r)
```

The latter will orient the system such that atom 4 points to the (1,0,0) direction.

Definition at line 139 of file prg\_syrotation\_mod.F90.

#### 9.28.3 Variable Documentation

```
9.28.3.1 integer, parameter prg_syrotation_mod::dp = kind(1.0d0) [private]
```

Definition at line 11 of file prg syrotation mod.F90.

# 9.29 prg\_system\_mod Module Reference

A module to read and handle chemical systems.

# **Data Types**

· type estruct\_type

Electronic structure type.

type system\_type

System type.

### **Functions/Subroutines**

• subroutine, public prg get nameandext (fullfilename, filename, ext)

Get the name and extension of a file.

• subroutine, public prg\_parse\_system (system, filename, extin)

The parser for the chemical system.

• subroutine, public prg\_write\_system (system, filename, extension)

Write system in .xyz, .dat or pdb file.

• subroutine, public prg\_write\_trajectory (system, iter, each, prg\_deltat, filename, extension)

Write trajectory in .xyz, .dat or pdb file.

subroutine, public prg\_write\_trajectoryandproperty (system, iter, each, prg\_deltat, scalarprop, filename, extension)

Write trajectory and atomic properties. Only pdb file.

subroutine, public prg\_make\_random\_system (system, nats, seed, lx, ly, lz)

Make random Xx system.

• subroutine prg\_parameters\_to\_vectors (abc\_angles, lattice\_vector)

Transforms the lattice parameters into lattice vectors.

• subroutine <a href="mailto:precamparameters">prg\_vectors\_to\_parameters</a> (lattice\_vector, abc\_angles)

Transforms the lattice vectors into lattice parameters.

subroutine, public prg\_get\_origin (coords, origin)

Get the origin of the coordinates.

subroutine, public prg\_get\_distancematrix (coords, dmat)

Get the distance matrix.

• subroutine, public prg\_translateandfoldtobox (coords, lattice\_vectors, origin, verbose)

Translate and fold to box.

• subroutine, public prg\_centeratbox (coords, lattice\_vectors, verbose)

Translate geometric center to the center of the box.

• subroutine, public prg wraparound (coords, lattice vectors, index, verbose)

Wrap around atom i using pbc.

• subroutine, public prg translatetogeomcandfoldtobox (coords, lattice vectors, origin)

Translate to geometric center.

subroutine, public prg\_replicate (coords, symbols, lattice\_vectors, nx, ny, nz)

Extend/replicate system along lattice vectors.

• subroutine, public prg\_get\_recip\_vects (lattice\_vectors, recip\_vectors, volr, volk)

Get the volume of the cell and the reciprocal vectors: This soubroutine computes:

• subroutine, public prg get dihedral (coords, id1, id2, id3, id4, dihedral)

Get the dihedral angle given four atomic positions.

• subroutine, public <a href="mailto:prg\_get\_covgraph">prg\_get\_covgraph</a> (sy, nnStructMindist, nnStruct, nrnnstruct, bml\_type, factor, gcov\_bml, mdimin, verbose)

Get the covalency graph in bml format.

- subroutine prg\_get\_covgraph\_int (sy, nnStructMindist, nnStruct, nrnnstruct, bml\_type, factor, gcov\_bml, mdimin, verbose)
- subroutine, public prg\_get\_covgraph\_h (sy, nnStructMindist, nnStruct, nrnnstruct, rcut, graph\_h, mdimin, verbose)

Get the covanlency graph.

subroutine, public prg\_get\_subsystem (sy, Isize, indices, sbsy, verbose)

Get a subsystem out of the total system.

subroutine, public prg\_destroy\_subsystems (sbsy, verbose)

Destroy allocated subsystem.

• subroutine, public prg\_molpartition (sy, npart, nnStructMindist, nnStruct, nrnnstruct, hetatm, gp, verbose)

\*Partition by molecule.\*

• subroutine, public prg\_get\_partial\_atomgraph (rho\_bml, hindex, gch\_bml, threshold, verbose)

Get partial subgraph based on the Density matrix.

subroutine, public prg\_collect\_graph\_p (rho\_bml, nc, nats, hindex, chindex, graph\_p, threshold, mdimin, ver-bose)

Collect the small graph to build the full graph.

subroutine, public prg\_merge\_graph (graph\_p, graph\_h)

Get partial subgraph based on the Density matrix.

• subroutine, public prg\_merge\_graph\_adj (graph\_p, graph\_h, xadj, adjncy)

Get partial subgraph based on the Density matrix.

• subroutine, public prg\_adj2bml (xadj, adjncy, bml\_type, g\_bml)

prg\_adj2bml

• subroutine, public prg\_graph2bml (graph, bml\_type, g\_bml)

Graph2bml.

• subroutine, public prg\_graph2vector (graph, vector, maxnz)

Vectorize graph.

• subroutine, public prg\_vector2graph (vector, graph, maxnz)

Back to graph.

subroutine, public prg\_sortadj (xadj, adjncy)

Sort adj NOTE: this might not be needed anymre since the bml\_get\_adj routine is sorting the values.

#### **Variables**

• integer, parameter dp = kind(1.0d0)

## 9.29.1 Detailed Description

A module to read and handle chemical systems.

This module will be used to build and handle a molecular system.

#### 9.29.2 Function/Subroutine Documentation

9.29.2.1 subroutine, public prg\_system\_mod::prg\_adj2bml ( integer, dimension(:), intent(in) xadj, integer, dimension(:), intent(in) adjncy, character(20), intent(in) bml\_type, type(bml\_matrix\_t), intent(inout) g\_bml )

prg\_adj2bml

#### **Parameters**

xadj	CSR start values for the adjacency matrix.
adjncy	CSR positions of adjacency matrix.
bml_type	bml format.
g_bml	graph in bml format.

Definition at line 2296 of file prg\_system\_mod.F90.

9.29.2.2 subroutine, public prg\_system\_mod::prg\_centeratbox ( real(dp), dimension(:,:), intent(inout), allocatable *coords*, real(dp), dimension(:,:), intent(in) *lattice\_vectors*, integer, intent(in), optional *verbose* )

Translate geometric center to the center of the box.

## **Parameters**

coords	Coordinates of the system (see system_type).
lattice_vectors	System lattice vectors.
verbose	Verbosity level.

Definition at line 1290 of file prg\_system\_mod.F90.

9.29.2.3 subroutine, public prg\_system\_mod::prg\_collect\_graph\_p ( type(bml\_matrix\_t), intent(in) rho\_bml, integer, intent(in) nc, integer, intent(in) nats, integer, dimension(:,:), intent(in) hindex, integer, dimension(:), intent(in) chindex, integer, dimension(:), intent(in) mdimin, integer, intent(in) mdimin, integer, intent(in), optional verbose )

Collect the small graph to build the full graph.

#### **Parameters**

rho_bml	Density matix in bml format.
nc	Number of core atoms.
nats	Number of atoms.
hindex	Hindex for the small part (see haindex)
chindex	Core-hallo index for the small part.
graph_p	Graph in an "ellpack" format.
threshold	Threshold to buil the density based atom projected graph.
verbose	Verbosity level.

Definition at line 2079 of file prg\_system\_mod.F90.

9.29.2.4 subroutine, public prg\_system\_mod::prg\_destroy\_subsystems ( type(system\_type), intent(inout) sbsy, integer, intent(in), optional verbose )

Destroy allocated subsystem.

This routine will deallocate all the arrays of the structures.

#### **Parameters**

sy	System to de deallocated (see system_type).
----	---

Definition at line 1864 of file prg\_system\_mod.F90.

9.29.2.5 subroutine, public prg\_system\_mod::prg\_get\_covgraph ( type(system\_type), intent(in) sy, real(dp), dimension(:,:), intent(in) nnStructMindist, integer, dimension(:,:), intent(in) nnStruct, integer, dimension(:), intent(in) nrnnstruct, character(20), intent(in) bml\_type, real(dp) factor, type(bml\_matrix\_t), intent(inout) gcov\_bml, integer, intent(in) mdimin, integer, intent(in), optional verbose )

Get the covalency graph in bml format.

This is the graph composed by the covalent bonds (edges) that are determined with the VDW radius.

## **Parameters**

sy	System structure (see system_type).
nnStructMindist	Minimun distance between atoms.
nnStruct	The neigbors J to I within Rcut that are all within the box.
nrnnstruct	Number of neigbors to I within Rcut that are all within the box.
bml_type	The bml type for constructing the graph.
gconv_bml	Covanlency graph in bml format.
verbose	Verbosity level.

Definition at line 1579 of file prg\_system\_mod.F90.

9.29.2.6 subroutine, public prg\_system\_mod::prg\_get\_covgraph\_h ( type(system\_type), intent(in) sy, real(dp), dimension(:,:), intent(in) nnStructMindist, integer, dimension(:,:), intent(in) nnStruct, integer, dimension(:), intent(in) nnstruct, real(dp), intent(in) rcut, integer, dimension(:,:), intent(inout), allocatable graph\_h, integer, intent(in) mdimin, integer, intent(in), optional verbose)

Get the covanlency graph.

This is the graph composed by the covalent bonds (edges) that are determined with the VDW radius.

#### **Parameters**

sy	System structure (see system_type).
nnStructMindist	Minimun distance between atoms.
nnStruct	The neigbors J to I within Rcut that are all within the box.
nrnnstruct	Number of neigbors to I within Rcut that are all within the box.
bml_type	The bml type for constructing the graph.
gconv_bml	Covanlency graph in bml format.
verbose	Verbosity level.

Definition at line 1712 of file prg\_system\_mod.F90.

9.29.2.7 subroutine prg\_system\_mod::prg\_get\_covgraph\_int ( type(system\_type), intent(in) sy, real(dp), dimension(:,:), intent(in) nnStructMindist, integer, dimension(:,:), intent(in) nnStruct, integer, dimension(:), intent(in) nrnnstruct, character(20), intent(in) bml\_type, real(dp) factor, type(bml\_matrix\_t), intent(inout) gcov\_bml, integer, intent(in) mdimin, integer, intent(in), optional verbose ) [private]

Definition at line 1653 of file prg\_system\_mod.F90.

9.29.2.8 subroutine, public prg\_system\_mod::prg\_get\_dihedral ( real(dp), dimension(:,:), intent(in) coords, integer, intent(in) id1, integer, intent(in) id3, integer, intent(in) id4, real(dp), intent(out) dihedral )

Get the dihedral angle given four atomic positions.

#### **Parameters**

sy	System structure
id1	Atom index 1
id2	Atom index 1
id3	Atom index 1
id4	Atom index 1
dihedral	Output dihedral angle

Definition at line 1523 of file prg\_system\_mod.F90.

9.29.2.9 subroutine, public prg\_system\_mod::prg\_get\_distancematrix ( real(dp), dimension(:,:), intent(in) coords, real(dp), dimension(:,:), intent(out), allocatable dmat )

Get the distance matrix.

#### **Parameters**

coords	Coordinates of the system (see system_type).
dmat	Distance matrix (nats x nats).

Definition at line 1216 of file prg\_system\_mod.F90.

9.29.2.10 subroutine, public prg\_system\_mod::prg\_get\_nameandext ( character(30), intent(in) *fullfilename*, character(30), intent(inout) *filename*, character(3), intent(inout) *ext* )

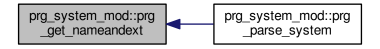
Get the name and extension of a file.

#### **Parameters**

fullfilename	Full filename.
filename	Filename of the system.
extension	Extension of the file.

Definition at line 203 of file prg\_system\_mod.F90.

Here is the caller graph for this function:



9.29.2.11 subroutine, public prg\_system\_mod::prg\_get\_origin ( real(dp), dimension(:,:), intent(in) coords, real(dp), dimension(:), intent(inout), allocatable origin )

Get the origin of the coordinates.

## **Parameters**

coords	Coordinates of teh system (see system_type).
origin	$(\min(x), \min(y), \min(z))$ set as the origin of the system.

Definition at line 1180 of file prg\_system\_mod.F90.

9.29.2.12 subroutine, public prg\_system\_mod::prg\_get\_partial\_atomgraph ( type(bml\_matrix\_t), intent(in) *rho\_bml*, integer, dimension(:,:), intent(in) *hindex*, type(bml\_matrix\_t), intent(inout) *gch\_bml*, real(dp), intent(in) *threshold*, integer, intent(in), optional *verbose* )

Get partial subgraph based on the Density matrix.

#### **Parameters**

rho_bml	Density matix in bml format.
hindex	Start and end index for every atom in the system.
gch_bml	Atom based graph in bml format.
threshold	Threshold value for constructing the graph.
verbose	Verbosity levels.

Definition at line 2013 of file prg system mod.F90.

9.29.2.13 subroutine, public prg\_system\_mod::prg\_get\_recip\_vects ( real(dp), dimension(:,:), intent(in) lattice\_vectors, real(dp), dimension(:,:), intent(inout), allocatable recip\_vectors, real(dp), intent(inout) volr, real(dp), intent(inout) volk )

Get the volume of the cell and the reciprocal vectors: This soubroutine computes:

- $b_1 = \frac{1}{V_2} a_1 \times a_2$
- $b_2 = \frac{1}{V_c} a_2 \times a_3$
- $b_3 = \frac{1}{V_2} a_3 \times a_1$
- $V_c = ||a_1 \cdot (a_2 \times a_3)||$
- $V_{BZ} = ||b_1 \cdot (b_2 \times b_3)||$

## **Parameters**

lattice_vectors	Lattice vectors for the system.
recip_vectors	Reciprocal vectors of the system.
volr	Volume of the cell.
volk	Volume of the reciprocal cell.

Definition at line 1474 of file prg\_system\_mod.F90.

9.29.2.14 subroutine, public prg\_system\_mod::prg\_get\_subsystem ( type(system\_type), intent(in) sy, integer, intent(in) lsize, integer, dimension(:), intent(in) indices, type(system\_type), intent(inout) sbsy, integer, intent(in), optional verbose )

Get a subsystem out of the total system.

This will get a subsystem from the total system guided by a partition.

#### **Parameters**

sy	System structure (see system_type).
Isize	Core+Hallo subsystem size.
indices	Partition indices.
sbsy	Subsystem to be extracted.

Definition at line 1777 of file prg\_system\_mod.F90.

9.29.2.15 subroutine, public prg\_system\_mod::prg\_graph2bml ( integer, dimension(:,:), intent(inout), allocatable *graph*, character(20), intent(in) *bml\_type*, type(bml\_matrix\_t), intent(inout) *g\_bml* )

Graph2bml.

#### **Parameters**

graph	Atom based graph in "ellpack" like format.
bml_type	Bml type (usually ellpack for graph starage)
g_bml	Graph in bml format.

Definition at line 2330 of file prg\_system\_mod.F90.

9.29.2.16 subroutine, public prg\_system\_mod::prg\_graph2vector ( integer, dimension(:,:), intent(inout) *graph*, integer, dimension(:), allocatable *vector*, integer *maxnz* )

Vectorize graph.

#### **Parameters**

graph	Ellpack graph.
vector	Vector to store the graph.

Definition at line 2373 of file prg\_system\_mod.F90.

9.29.2.17 subroutine, public prg\_system\_mod::prg\_make\_random\_system ( type(system\_type), intent(out) system, integer nats, integer seed, real(dp) lx, real(dp) lz )

Make random Xx system.

#### **Parameters**

system	System to be construucted.
nats	Number of atoms.
lx	length of the box for the x coordinate.
ly	length of the box for the y coordinate.
lz	length of the box for the z coordinate.

Definition at line 1056 of file prg\_system\_mod.F90.

9.29.2.18 subroutine, public prg\_system\_mod::prg\_merge\_graph ( integer, dimension(:,:), intent(inout) graph\_p, integer, dimension(:,:), intent(inout) graph\_h)

Get partial subgraph based on the Density matrix.

#### **Parameters**

graph⊷	Density matix based graph in bml format.
_p	
graph←	Hamiltonian matix based graph in bml format.
_h	

Definition at line 2171 of file prg\_system\_mod.F90.

9.29.2.19 subroutine, public prg\_system\_mod::prg\_merge\_graph\_adj ( integer, dimension(:,:), intent(inout), allocatable graph\_p, integer, dimension(:,:), intent(inout), allocatable graph\_h, integer, dimension(:), intent(inout), allocatable adjncy )

Get partial subgraph based on the Density matrix.

#### **Parameters**

graph⇔	Density matix based graph in "ellpack type format".
_p	
graph⇔	Hamiltonian matix based graph in "ellpack type format".
_h	
xadj	CSR start values for the adjacency matrix.
adjncy	CSR positions of adjacency matrix.

Definition at line 2222 of file prg\_system\_mod.F90.

9.29.2.20 subroutine, public prg\_system\_mod::prg\_molpartition ( type(system\_type), intent(in) sy, integer, intent(inout) npart, real(dp), dimension(:,:), intent(in) nnStructMindist, integer, dimension(:,:), intent(in) nrnstruct, character(2), intent(in) hetatm, type(graph\_partitioning\_t), intent(inout) gp, integer, intent(inout), optional verbose)

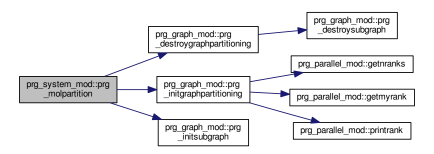
Partition by molecule.

#### **Parameters**

sy	System structure.
npart	Number of parts.
nnStructMindist	Minimum distance between neighbors.
nnStruct	The neighbors J to I within Rcut that are all within the box.
nrnnstruct	Number of neighbors to I within Rcut that are all within the box.
hetatm	Atom to be taken as the "center" of the by molecule partition.
gp	Graph partition structure.
Gerferales by Doxygen	Verbosity level.

Definition at line 1926 of file prg\_system\_mod.F90.

Here is the call graph for this function:



9.29.2.21 subroutine prg\_system\_mod::prg\_parameters\_to\_vectors ( real(dp), dimension(2,3), intent(in) abc\_angles, real(dp), dimension(3,3), intent(out) lattice\_vector ) [private]

Transforms the lattice parameters into lattice vectors.

#### **Parameters**

abc_angles	2x3 array containing the lattice parameters. abc_angles(1,1) = a, abc_angles(1,2) = b, and abc_angles(1,3) = c abc_angles(2,1) = $\alpha$ , abc_angles(2,2) = $\beta$ and abc_angles(2,3) = $\gamma$
lattice_vector	3x3 array containing the lattice vectors. lattice_vector(1,:) = $\overrightarrow{a}$

Definition at line 1102 of file prg\_system\_mod.F90.

Here is the caller graph for this function:



9.29.2.22 subroutine, public prg\_system\_mod::prg\_parse\_system ( type(system\_type), intent(out) system, character(len=\*) filename, character(3), intent(in), optional extin )

The parser for the chemical system.

#### **Parameters**

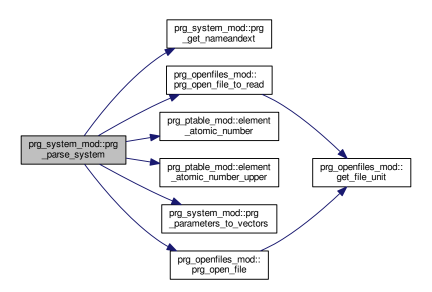
system	System to be constructed.
filename	Filename of the system.
extin	Extension of the file.

Assignment of species index for every atom.

**Todo** Integrate this loop in the loop for building the splist.

Definition at line 226 of file prg\_system\_mod.F90.

Here is the call graph for this function:



9.29.2.23 subroutine, public prg\_system\_mod::prg\_replicate ( real(dp), dimension(:,:), intent(inout), allocatable *coords*, character(2), dimension(:), intent(inout), allocatable *symbols*, real(dp), dimension(:,:), intent(inout) *lattice\_vectors*, integer, intent(in) *nx*, integer, intent(in) *ny*, integer, intent(in) *nz* )

Extend/replicate system along lattice vectors.

#### **Parameters**

coords	Coordinates of the system (see system_type).
symbols	Symbols for elements.
lattice_vectors	System lattice vectors.
nx	Number of lattice points in the v1 direction.
ny	Number of lattice points in the v2 direction.
nz	Number of lattice points in the v2 direction.

Definition at line 1414 of file prg\_system\_mod.F90.

9.29.2.24 subroutine, public prg\_system\_mod::prg\_sortadj ( integer, dimension(:), intent(inout) xadj, integer, dimension(:), intent(inout), allocatable adjncy )

Sort adj NOTE: this might not be needed anymre since the bml\_get\_adj routine is sorting the values.

Definition at line 2430 of file prg\_system\_mod.F90.

9.29.2.25 subroutine, public prg\_system\_mod::prg\_translateandfoldtobox ( real(dp), dimension(:,:), intent(inout), allocatable coords, real(dp), dimension(:,:), intent(in) lattice\_vectors, real(dp), dimension(:), intent(inout), allocatable origin, integer, intent(in), optional verbose )

Translate and fold to box.

#### **Parameters**

coords	Coordinates of the system (see system_type).
lattice_vectors	System lattice vectors.
origin	(min(x),min(y),min(z)) set as the origin of the system.

Definition at line 1240 of file prg\_system\_mod.F90.

9.29.2.26 subroutine, public prg\_system\_mod::prg\_translatetogeomcandfoldtobox ( real(dp), dimension(:,:), intent(inout), allocatable *coords*, real(dp), dimension(:,:), intent(in) *lattice\_vectors*, real(dp), dimension(:), intent(inout), allocatable *origin* )

Translate to geometric center.

#### **Parameters**

coords	Coordinates of the system (see system_type).
lattice_vectors	System lattice vectors.
origin	(min(x),min(y),min(z)) set as the origin of the system.

Definition at line 1373 of file prg\_system\_mod.F90.

9.29.2.27 subroutine, public prg\_system\_mod::prg\_vector2graph ( integer, dimension(:), intent(inout), allocatable *vector*, integer, dimension(:,:), intent(inout) *graph*, integer *maxnz* )

Back to graph.

## **Parameters**

vector	Vector to store the graph.
graph	Ellpack graph.

Definition at line 2402 of file prg\_system\_mod.F90.

9.29.2.28 subroutine prg\_system\_mod::prg\_vectors\_to\_parameters ( real(dp), dimension(3,3), intent(in) *lattice\_vector*, real(dp), dimension(2,3), intent(out) *abc\_angles* ) [private]

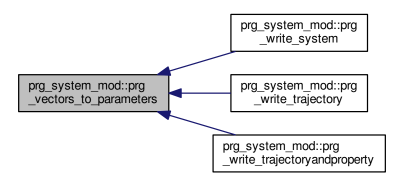
Transforms the lattice vectors into lattice parameters.

#### **Parameters**

lattice_vector	3x3 array containing the lattice vectors. lattice_vector(1,:) = $\overrightarrow{a}$
abc_angles	2x3 array containing the lattice parameters. abc_angles(1,1) = a, abc_angles(1,2) = b and abc_angles(1,3) = c abc_angles(2,1) = $\alpha$ , abc_angles(2,2) = $\beta$ , and abc_angles(2,3) = $\gamma$ .

Definition at line 1144 of file prg\_system\_mod.F90.

Here is the caller graph for this function:



9.29.2.29 subroutine, public prg\_system\_mod::prg\_wraparound ( real(dp), dimension(:,:), intent(inout), allocatable *coords*, real(dp), dimension(:,:), intent(in) *lattice\_vectors*, integer, intent(in) *index*, integer, intent(in), optional *verbose* )

Wrap around atom i using pbc.

#### **Parameters**

coords	Coordinates of the system (see system_type).
lattice_vectors	System lattice vectors.
index	Index atom to wrap around

Definition at line 1330 of file prg\_system\_mod.F90.

9.29.2.30 subroutine, public prg\_system\_mod::prg\_write\_system ( type(system\_type), intent(in) system, character(\*) filename, character(3) extension )

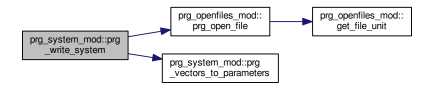
Write system in .xyz, .dat or pdb file.

#### **Parameters**

system	System to be constructed.
filename	File name.
extension	Extension of the file.

Definition at line 635 of file prg\_system\_mod.F90.

Here is the call graph for this function:



9.29.2.31 subroutine, public prg\_system\_mod::prg\_write\_trajectory ( type(system\_type), intent(in) system, integer, intent(in) system, integer, intent(in) each, real(dp), intent(in) prg\_deltat, character(\*) filename, character(3) extension )

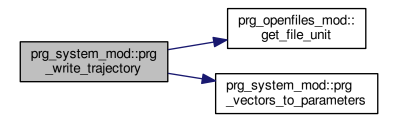
Write trajectory in .xyz, .dat or pdb file.

#### **Parameters**

system	System to be appended to the trajectory file.
iter	Simulation step.
each	Writing frequency.
filename	File name for the trajectory.
extension	Extension of the file.

Definition at line 823 of file prg\_system\_mod.F90.

Here is the call graph for this function:



9.29.2.32 subroutine, public prg\_system\_mod::prg\_write\_trajectoryandproperty ( type(system\_type), intent(in) system, integer, intent(in) iter, integer, intent(in) each, real(dp), intent(in) prg\_deltat, real(dp), dimension(:), intent(in) scalarprop, character(\*) filename, character(3) extension )

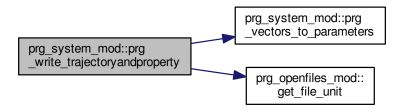
Write trajectory and atomic properties. Only pdb file.

#### **Parameters**

system	System to be appended to the trajectory file.
iter	Simulation step.
each	Writing frequency.
prg_deltat	Integration step.
scalarprop	Scalar property to plot on atoms.
filename	File name for the trajectory.
extension	Extension of the file.

Definition at line 946 of file prg\_system\_mod.F90.

Here is the call graph for this function:



#### 9.29.3 Variable Documentation

```
9.29.3.1 integer, parameter prg_system_mod::dp = kind(1.0d0) [private]
```

Definition at line 16 of file prg\_system\_mod.F90.

# 9.30 prg\_timer\_mod Module Reference

The timer module.

## **Data Types**

· type timer\_status\_t

Timer status type.

#### **Functions/Subroutines**

• subroutine, public timer\_prg\_init ()

Initialize timers.

• subroutine prg\_timer\_getid ()

Get timer id.

• subroutine, public prg\_timer\_shutdown ()

Done with timers.

• subroutine, public <a href="mailto:prg\_timer\_start">prg\_timer\_start</a> (itimer, tag)

Start Timing.

• subroutine, public prg\_timer\_stop (itimer, verbose)

Stop timing.

- subroutine, public prg\_timer\_collect ()
- subroutine, public prg\_timer\_results ()
- real(8) function, public time2milliseconds ()
- subroutine, public prg\_print\_date\_and\_time (tag)
- character(2) function, private int2char (ival)

#### **Variables**

- integer, parameter dp = kind(1.0d0)
- integer, public loop\_timer
- integer, public sp2\_timer
- integer, public genx\_timer
- integer, public part\_timer
- integer, public subgraph\_timer
- integer, public deortho\_timer
- integer, public ortho\_timer
- integer, public zdiag\_timer
- integer, public graphsp2\_timer
- integer, public subind\_timer
- integer, public subext timer
- integer, public subsp2\_timer

- · integer, public suball\_timer
- integer, public bmult\_timer
- integer, public badd\_timer
- integer, public dyn\_timer
- integer, public mdloop\_timer
- integer, public buildz timer
- integer, public realcoul\_timer
- integer, public recipcoul\_timer
- integer, public pairpot\_timer
- integer, public halfverlet\_timer
- integer, public pos\_timer
- · integer, public nlist timer
- integer tstart\_clock
- integer tstop\_clock
- integer tclock\_rate
- integer tclock\_max
- integer num\_timers
- type(timer\_status\_t), dimension(:), allocatable ptimer

## 9.30.1 Detailed Description

The timer module.

Sets up timers that can be used to time other routines.

Example use of dynamic timing:

```
call timer_prg_init()
```

call prg\_timer\_start(dyn\_timer,"timer\_tag")

.... code lines ...

call prg\_timer\_stop(dyn\_timer,1)

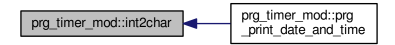
This will write the time it takes to execute "code lines" and it will name it "timer\_tag"

## 9.30.2 Function/Subroutine Documentation

9.30.2.1 character(2) function, private prg\_timer\_mod::int2char( integer, intent(in) ival ) [private]

Definition at line 394 of file prg\_timer\_mod.F90.

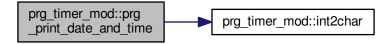
Here is the caller graph for this function:



9.30.2.2 subroutine, public prg\_timer\_mod::prg\_print\_date\_and\_time ( character(len=\*), intent(in) tag )

Definition at line 371 of file prg\_timer\_mod.F90.

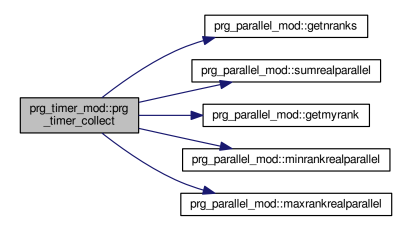
Here is the call graph for this function:



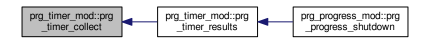
9.30.2.3 subroutine, public prg\_timer\_mod::prg\_timer\_collect( )

Definition at line 253 of file prg\_timer\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:



9.30.2.4 subroutine prg\_timer\_mod::prg\_timer\_getid( ) [private]

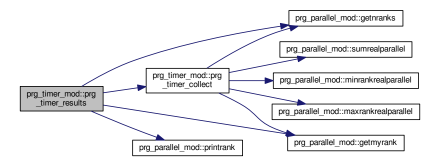
Get timer id.

Definition at line 200 of file prg\_timer\_mod.F90.

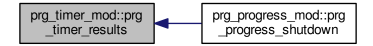
9.30.2.5 subroutine, public prg\_timer\_mod::prg\_timer\_results ( )

Definition at line 317 of file prg\_timer\_mod.F90.

Here is the call graph for this function:



Here is the caller graph for this function:

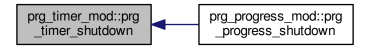


9.30.2.6 subroutine, public prg\_timer\_mod::prg\_timer\_shutdown()

Done with timers.

Definition at line 205 of file prg\_timer\_mod.F90.

Here is the caller graph for this function:



9.30.2.7 subroutine, public prg\_timer\_mod::prg\_timer\_start ( integer, intent(in) itimer, character(len=\*), intent(in), optional tag )

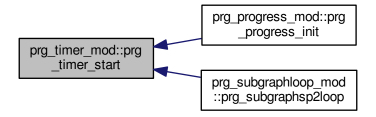
Start Timing.

#### **Parameters**

itimer	The index of the timer to start.
tag	Optional parameter to retag the timer on the fly.

Definition at line 215 of file prg\_timer\_mod.F90.

Here is the caller graph for this function:



9.30.2.8 subroutine, public prg\_timer\_mod::prg\_timer\_stop ( integer, intent(in) itimer, integer, intent(in), optional verbose )

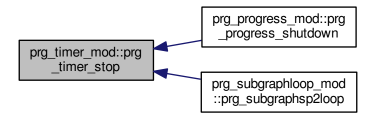
Stop timing.

## **Parameters**

itimer	The index of the timer to stop.
verbose	Optional parameters to print partial times.

Definition at line 233 of file prg\_timer\_mod.F90.

Here is the caller graph for this function:



9.30.2.9 real(8) function, public prg\_timer\_mod::time2milliseconds ( )

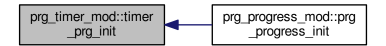
Definition at line 360 of file prg\_timer\_mod.F90.

9.30.2.10 subroutine, public prg\_timer\_mod::timer\_prg\_init ( )

Initialize timers.

Definition at line 132 of file prg\_timer\_mod.F90.

Here is the caller graph for this function:



## 9.30.3 Variable Documentation

9.30.3.1 integer, public prg\_timer\_mod::badd\_timer

Definition at line 48 of file prg\_timer\_mod.F90.

9.30.3.2 integer, public prg\_timer\_mod::bmult\_timer Definition at line 48 of file prg\_timer\_mod.F90. 9.30.3.3 integer, public prg\_timer\_mod::buildz\_timer Definition at line 49 of file prg\_timer\_mod.F90. 9.30.3.4 integer, public prg\_timer\_mod::deortho\_timer Definition at line 45 of file prg\_timer\_mod.F90. 9.30.3.5 integer, parameter prg\_timer\_mod::dp = kind(1.0d0) [private] Definition at line 32 of file prg\_timer\_mod.F90. 9.30.3.6 integer, public prg\_timer\_mod::dyn\_timer Definition at line 49 of file prg\_timer\_mod.F90. 9.30.3.7 integer, public prg\_timer\_mod::genx\_timer Definition at line 44 of file prg\_timer\_mod.F90. 9.30.3.8 integer, public prg\_timer\_mod::graphsp2\_timer Definition at line 46 of file prg\_timer\_mod.F90. 9.30.3.9 integer, public prg\_timer\_mod::halfverlet\_timer Definition at line 51 of file prg\_timer\_mod.F90. 9.30.3.10 integer, public prg\_timer\_mod::loop\_timer Definition at line 44 of file prg\_timer\_mod.F90. 9.30.3.11 integer, public prg\_timer\_mod::mdloop\_timer Definition at line 49 of file prg\_timer\_mod.F90.

```
9.30.3.12 integer, public prg_timer_mod::nlist_timer
Definition at line 51 of file prg_timer_mod.F90.
9.30.3.13 integer prg_timer_mod::num_timers [private]
Definition at line 122 of file prg_timer_mod.F90.
9.30.3.14 integer, public prg_timer_mod::ortho_timer
Definition at line 46 of file prg_timer_mod.F90.
9.30.3.15 integer, public prg_timer_mod::pairpot_timer
Definition at line 50 of file prg_timer_mod.F90.
9.30.3.16 integer, public prg_timer_mod::part_timer
Definition at line 45 of file prg_timer_mod.F90.
9.30.3.17 integer, public prg_timer_mod::pos_timer
Definition at line 51 of file prg_timer_mod.F90.
9.30.3.18 type (timer_status_t), dimension(:), allocatable prg_timer_mod::ptimer [private]
Definition at line 124 of file prg_timer_mod.F90.
9.30.3.19 integer, public prg_timer_mod::realcoul_timer
Definition at line 50 of file prg_timer_mod.F90.
9.30.3.20
          integer, public prg_timer_mod::recipcoul_timer
Definition at line 50 of file prg_timer_mod.F90.
9.30.3.21 integer, public prg_timer_mod::sp2_timer
Definition at line 44 of file prg_timer_mod.F90.
```

```
9.30.3.22 integer, public prg_timer_mod::suball_timer
Definition at line 48 of file prg_timer_mod.F90.
9.30.3.23 integer, public prg_timer_mod::subext_timer
Definition at line 47 of file prg_timer_mod.F90.
9.30.3.24 integer, public prg_timer_mod::subgraph_timer
Definition at line 45 of file prg_timer_mod.F90.
9.30.3.25 integer, public prg_timer_mod::subind_timer
Definition at line 47 of file prg_timer_mod.F90.
9.30.3.26 integer, public prg_timer_mod::subsp2_timer
Definition at line 47 of file prg_timer_mod.F90.
9.30.3.27 integer prg_timer_mod::tclock_max [private]
Definition at line 121 of file prg_timer_mod.F90.
9.30.3.28 integer prg_timer_mod::tclock_rate [private]
Definition at line 121 of file prg_timer_mod.F90.
9.30.3.29 integer prg_timer_mod::tstart_clock [private]
Definition at line 121 of file prg_timer_mod.F90.
9.30.3.30 integer prg_timer_mod::tstop_clock [private]
Definition at line 121 of file prg_timer_mod.F90.
9.30.3.31 integer, public prg_timer_mod::zdiag_timer
```

Definition at line 46 of file prg\_timer\_mod.F90.

# 9.31 prg\_xlbo\_mod Module Reference

A module to perform XLBO integration.

## **Data Types**

type xlbo\_type

General xlbo solver type.

#### **Functions/Subroutines**

```
• subroutine, public prg_parse_xlbo (xlbo, filename)
```

The parser for XLBO parser.

• subroutine, public prg\_xlbo\_nint (charges, n, n\_0, n\_1, n\_2, n\_3, n\_4, n\_5, mdstep, xl)

This routine integrates the dynamical variable "n".

• subroutine, public prg\_xlbo\_fcoulupdate (fcoul, charges, n)

Adjust forces for the linearized XLBOMD functional.

## **Variables**

```
• integer, parameter dp = kind(1.0d0)
```

```
• real(dp), parameter c0 = -6.0_dp
```

Coefficients for modified Verlet integration.

- real(dp), parameter c1 = 14.0\_dp
- real(dp), parameter c2 = -8.0\_dp
- real(dp), parameter c3 = -3.0\_dp
- real(dp), parameter c4 = 4.0 dp
- real(dp), parameter c5 = -1.0\_dp
- real(dp), parameter kappa = 1.82\_dp

Coefficients for modified Verlet integration.

- real(dp), parameter alpha = 0.018\_dp
- real(dp), parameter cc = 0.9\_dp

## 9.31.1 Detailed Description

A module to perform XLBO integration.

This module will be used to compute integrate the dynamical variable "n" in xlbo.

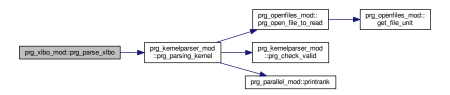
## 9.31.2 Function/Subroutine Documentation

9.31.2.1 subroutine, public prg\_xlbo\_mod::prg\_parse\_xlbo ( type(xlbo\_type), intent(inout) xlbo, character(len=\*) filename )

The parser for XLBO parser.

Definition at line 62 of file prg\_xlbo\_mod.F90.

Here is the call graph for this function:



9.31.2.2 subroutine, public prg\_xlbo\_mod::prg\_xlbo\_fcoulupdate ( real(dp), dimension(:,:), intent(inout) *fcoul*, real(dp), dimension(:), intent(inout) *charges*, real(dp), dimension(:), intent(inout) *n* )

Adjust forces for the linearized XLBOMD functional.

**Parameters** 

charges

Definition at line 158 of file prg\_xlbo\_mod.F90.

9.31.2.3 subroutine, public prg\_xlbo\_mod::prg\_xlbo\_nint ( real(dp), dimension(:), intent(in), allocatable *charges*, real(dp), dimension(:), intent(inout), allocatable *n\_0*, real(dp), dimension(:), intent(inout), allocatable *n\_0*, real(dp), dimension(:), intent(inout), allocatable *n\_2*, real(dp), dimension(:), intent(inout), allocatable *n\_3*, real(dp), dimension(:), intent(inout), allocatable *n\_4*, real(dp), dimension(:), intent(inout), allocatable *n\_5*, integer, intent(in) *mdstep*, type(xlbo\_type), intent(in) *xl*)

This routine integrates the dynamical variable "n".

**Parameters** 

charges

Definition at line 118 of file prg\_xlbo\_mod.F90.

## 9.31.3 Variable Documentation

```
9.31.3.1 real(dp), parameter prg_xlbo_mod::alpha = 0.018_dp [private]
Definition at line 28 of file prg_xlbo_mod.F90.
9.31.3.2 real(dp), parameter prg_xlbo_mod::c0 = -6.0_dp [private]
Coefficients for modified Verlet integration.
Definition at line 19 of file prg_xlbo_mod.F90.
9.31.3.3 real(dp), parameter prg_xlbo_mod::c1 = 14.0_dp [private]
Definition at line 20 of file prg_xlbo_mod.F90.
9.31.3.4 real(dp), parameter prg_xlbo_mod::c2 = -8.0_dp [private]
Definition at line 21 of file prg_xlbo_mod.F90.
9.31.3.5 real(dp), parameter prg_xlbo_mod::c3 = -3.0_dp [private]
Definition at line 22 of file prg xlbo mod.F90.
9.31.3.6 real(dp), parameter prg_xlbo_mod::c4 = 4.0_dp [private]
Definition at line 23 of file prg_xlbo_mod.F90.
9.31.3.7 real(dp), parameter prg_xlbo_mod::c5 = -1.0_dp [private]
Definition at line 24 of file prg_xlbo_mod.F90.
9.31.3.8 real(dp), parameter prg_xlbo_mod::cc = 0.9_dp [private]
Definition at line 29 of file prg xlbo mod.F90.
9.31.3.9 integer, parameter prg_xlbo_mod::dp = kind(1.0d0) [private]
Definition at line 16 of file prg_xlbo_mod.F90.
9.31.3.10 real(dp), parameter prg_xlbo_mod::kappa = 1.82_dp [private]
Coefficients for modified Verlet integration.
```

Definition at line 27 of file prg\_xlbo\_mod.F90.

# 9.32 prg\_xlkernel\_mod Module Reference

Add name.

### **Data Types**

type xlk\_type

#### **Functions/Subroutines**

- subroutine, public prg\_parse\_xlkernel (input, filename)
  - The parser for the mixer routines.
- subroutine, public prg\_fermi (D0, QQ, ee, gap, Fe\_vec, mu0, H, Z, Nocc, T, OccErrLim, MaxIt, HDIM)
- subroutine, public prg\_kernel\_fermi\_full (KK, JJ, D0, mu0, mu1, T, RX, RY, RZ, LBox, Hubbard\_U, Element
   — Type, Nr\_atoms, MaxIt, eps, m, HDIM, Max\_Nr\_Neigh, Coulomb\_acc, TIMERATIO, nnRx, nnRy, nnRz,
   nrnnlist, nnType, H\_INDEX\_START, H\_INDEX\_END, H, S, Z, Nocc, Znuc, QQ, ee, Fe\_vec)
- subroutine, public prg\_v\_kernel\_fermi (D0, dq\_dv, v, mu0, mu1, T, RX, RY, RZ, LBox, Hubbard\_U, Element
   \_Type, Nr\_atoms, MaxIt, eps, m, HDIM, Max\_Nr\_Neigh, Coulomb\_acc, TIMERATIO, nnRx, nnRy, nnRz,
   nrnnlist, nnType, H\_INDEX\_START, H\_INDEX\_END, H, S, Z, Nocc, Znuc, QQ, ee, Fe\_vec)
- subroutine, private prg get deriv finite temp (P1, H0, H1, Nocc, T, Q, ev, fe, mu0, eps, HDIM)
- subroutine, private prg\_mmult (alpha, A, B, beta, C, TA, TB, HDIM)
- subroutine, private prg\_eig (A, Q, ee, type, HDIM)
- subroutine, private prg inv (X, XI, HDIM)
- subroutine, public prg rank1 (verbose)

Rank1 kernel ....

## **Variables**

integer, parameter dp = kind(1.0d0)

#### 9.32.1 Detailed Description

Add name.

XL kernel (To be integrated)

Note

This module is still not functional

## 9.32.2 Function/Subroutine Documentation

9.32.2.1 subroutine, private prg\_xlkernel\_mod::prg\_eig ( real(prec), dimension(hdim,hdim), intent(in) A, real(prec), dimension(hdim,hdim), intent(out) Q, real(prec), dimension(hdim), intent(out) ee, character(1), intent(in) type, integer(prec), intent(in) HDIM ) [private]

Definition at line 384 of file prg\_xlkernel\_mod.F90.

9.32.2.2 subroutine, public prg\_xlkernel\_mod::prg\_fermi ( real(prec), dimension(hdim,hdim), intent(out) *D0*, real(prec), dimension(hdim,hdim), intent(out) *QQ*, real(prec), dimension(hdim), intent(out) *ee*, real(prec), intent(out) *gap*, real(prec), dimension(hdim), intent(out) *Fe\_vec*, real(prec), intent(inout) *mu0*, real(prec), dimension(hdim,hdim), intent(in) *H*, real(prec), dimension(hdim,hdim), intent(in) *Z*, integer(prec), intent(in) *Nocc*, real(prec), intent(in) *T*, real(prec), intent(in) *OccErrLim*, integer(prec), intent(in) *MaxIt*, integer(prec), intent(in) *HDIM* )

Definition at line 88 of file prg xlkernel mod.F90.

9.32.2.3 subroutine, private prg\_xlkernel\_mod::prg\_get\_deriv\_finite\_temp ( real(prec), dimension(hdim,hdim), intent(out) P1, real(prec), dimension(hdim,hdim), intent(in) H0, real(prec), dimension(hdim,hdim), intent(in) H1, integer(prec), intent(in) Nocc, real(prec), intent(in) T, real(prec), dimension(hdim,hdim), intent(in) Q, real(prec), dimension(hdim), intent(in) ev, real(prec), dimension(hdim), intent(in) fe, real(prec), intent(inout) mu0, real(prec), intent(in) eps, integer(prec), intent(in) HDIM) [private]

Definition at line 306 of file prg\_xlkernel\_mod.F90.

9.32.2.4 subroutine, private prg\_xlkernel\_mod::prg\_inv ( real(prec), dimension(hdim,hdim), intent(in) X, real(prec), dimension(hdim,hdim), intent(out) XI, integer(prec), intent(in) HDIM ) [private]

Definition at line 411 of file prg\_xlkernel\_mod.F90.

9.32.2.5 subroutine, public prg\_xlkernel\_mod::prg\_kernel\_fermi\_full ( real(prec), dimension(nr\_atoms,nr\_atoms), intent(out) KK, real(prec), dimension(nr\_atoms,nr\_atoms), intent(inout) DO, real(prec), intent(inout) mu0, real(prec), intent(inout) mu1, real(prec), intent(in) T, real(prec), dimension(nr\_atoms), intent(in) RX, real(prec), dimension(nr\_atoms), intent(in) RY, real(prec), dimension(nr\_atoms), intent(in) RZ, real(prec), dimension(3), intent(in) LBox, real(prec), dimension(nr\_atoms), intent(in) Hubbard\_U, character(10), dimension(nr\_atoms), intent(in) Element\_Type, integer(prec), intent(in) Nr\_atoms, integer(prec), intent(in) Maxlt, real(prec), intent(in) eps, integer(prec), intent(in) m, integer(prec), intent(in) HDIM, integer(prec), intent(in) Max\_Nr\_Neigh, real(prec), intent(in) Coulomb\_acc, real(prec), intent(in) TIMERATIO, real(prec), dimension(nr\_atoms,max\_nr\_neigh), intent(in) nnRx, real(prec), dimension(nr\_atoms,max\_nr\_neigh), intent(in) nnRz, integer(prec), dimension(nr\_atoms), intent(in) nnnlist, integer(prec), dimension(nr\_atoms,max\_nr\_neigh), intent(in) nnType, integer(prec), dimension(nr\_atoms), intent(in) H\_INDEX\_START, integer(prec), dimension(nr\_atoms), intent(in) S, real(prec), dimension(hdim,hdim), intent(in) Z, integer(prec), intent(in) Nocc, real(prec), dimension(nr\_atoms), intent(in) Znuc, real(prec), dimension(hdim,hdim), intent(in) intent(in) QQ, real(prec), dimension(hdim), intent(in) ee, real(prec), dimension(hdim), intent(in) Fe\_vec)

Definition at line 144 of file prg\_xlkernel\_mod.F90.

9.32.2.6 subroutine, private prg\_xlkernel\_mod::prg\_mmult ( real(prec), intent(in) alpha, real(prec), dimension(hdim,hdim), intent(in) A, real(prec), dimension(hdim,hdim), intent(in) B, real(prec), intent(in) beta, real(prec), dimension(hdim,hdim), intent(inout) C, character(1), intent(in) TA, character(1), intent(in) TB, integer(prec), intent(in) HDIM ) [private]

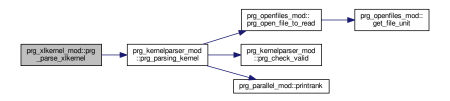
Definition at line 366 of file prg\_xlkernel\_mod.F90.

9.32.2.7 subroutine, public prg\_xlkernel\_mod::prg\_parse\_xlkernel ( type(xlk\_type), intent(inout) input, character(len=\*) filename )

The parser for the mixer routines.

Definition at line 39 of file prg xlkernel mod.F90.

Here is the call graph for this function:



9.32.2.8 subroutine, public prg\_xlkernel\_mod::prg\_rank1 ( integer, intent(in) verbose )

Rank1 kernel ....

#### **Parameters**

param1	
verbose	Different levels of verbosity.

Definition at line 439 of file prg\_xlkernel\_mod.F90.

9.32.2.9 subroutine, public prg\_xlkernel\_mod::prg\_v\_kernel\_fermi ( real(prec), dimension(hdim,hdim), intent(inout) D0, real(prec), dimension(nr\_atoms), intent(out) dq\_dv, real(prec), dimension(nr\_atoms), intent(in) v, real(prec), intent(inout) mu0, real(prec), intent(inout) mu1, real(prec), intent(in) T, real(prec), dimension(nr\_atoms), intent(in) RZ, real(prec), dimension(3), intent(in) LBox, real(prec), dimension(nr\_atoms), intent(in) Hubbard\_U, character(10), dimension(nr\_atoms), intent(in) Element\_Type, integer(prec), intent(in) Nr\_atoms, integer(prec), intent(in) MaxIt, real(prec), intent(in) eps, integer(prec), intent(in) m, integer(prec), intent(in) HDIM, integer(prec), intent(in) Max\_Nr\_Neigh, real(prec), intent(in) Coulomb\_acc, real(prec), intent(in) TIMERATIO, real(prec), dimension(nr\_atoms,max\_nr\_neigh), intent(in) nnRx, real(prec), dimension(nr\_atoms,max\_nr\_neigh), intent(in) nnRx, real(prec), dimension(nr\_atoms), intent(in) nnnIst, integer(prec), dimension(nr\_atoms,max\_nr\_neigh), intent(in) nnType, integer(prec), dimension(nr\_atoms), intent(in) H\_INDEX\_START, integer(prec), dimension(nr\_atoms), intent(in) Nnntitent(in) Nnntitent(in) Nnntitent(in) Nnntitent(in) Nntitent(in) Nntitent(

Definition at line 235 of file prg xlkernel mod.F90.

# 9.32.3 Variable Documentation

**9.32.3.1** integer, parameter prg\_xlkernel\_mod::dp = kind(1.0d0) [private]

Definition at line 15 of file prg\_xlkernel\_mod.F90.

# **Chapter 10**

# **Data Type Documentation**

10.1 prg\_chebyshev\_mod::chebdata\_type Type Reference

General Cheb solver type.

## **Public Attributes**

- character(100) flavor
- character(100) bml\_type
- character(100) jobname
- integer mdim
- integer ncoeffs
- integer ndim
- integer verbose
- integer npts
- real(dp) atr
- real(dp) bndfil
- real(dp) ef
- real(dp) estep
- real(dp) fermitol
- real(dp) kbt
- real(dp) threshold
- logical getef
- logical jon
- logical trkfunc

# 10.1.1 Detailed Description

General Cheb solver type.

Definition at line 28 of file prg\_chebyshev\_mod.F90.

## 10.1.2 Member Data Documentation

10.1.2.1 real(dp) prg\_chebyshev\_mod::chebdata\_type::atr

Definition at line 33 of file prg\_chebyshev\_mod.F90.

10.1.2.2 character(100) prg\_chebyshev\_mod::chebdata\_type::bml\_type

Definition at line 30 of file prg\_chebyshev\_mod.F90.

10.1.2.3 real(dp) prg\_chebyshev\_mod::chebdata\_type::bndfil

Definition at line 33 of file prg\_chebyshev\_mod.F90.

10.1.2.4 real(dp) prg\_chebyshev\_mod::chebdata\_type::ef

Definition at line 33 of file prg\_chebyshev\_mod.F90.

10.1.2.5 real(dp) prg\_chebyshev\_mod::chebdata\_type::estep

Definition at line 33 of file prg\_chebyshev\_mod.F90.

10.1.2.6 real(dp) prg\_chebyshev\_mod::chebdata\_type::fermitol

Definition at line 34 of file prg\_chebyshev\_mod.F90.

10.1.2.7 character(100) prg\_chebyshev\_mod::chebdata\_type::flavor

Definition at line 29 of file prg\_chebyshev\_mod.F90.

10.1.2.8 logical prg\_chebyshev\_mod::chebdata\_type::getef

Definition at line 35 of file prg\_chebyshev\_mod.F90.

10.1.2.9 character(100) prg\_chebyshev\_mod::chebdata\_type::jobname

Definition at line 30 of file prg chebyshev mod.F90.

10.1.2.10 logical prg\_chebyshev\_mod::chebdata\_type::jon

Definition at line 35 of file prg\_chebyshev\_mod.F90.

10.1.2.11 real(dp) prg\_chebyshev\_mod::chebdata\_type::kbt

Definition at line 34 of file prg\_chebyshev\_mod.F90.

10.1.2.12 integer prg\_chebyshev\_mod::chebdata\_type::mdim

Definition at line 31 of file prg\_chebyshev\_mod.F90.

10.1.2.13 integer prg\_chebyshev\_mod::chebdata\_type::ncoeffs

Definition at line 31 of file prg\_chebyshev\_mod.F90.

10.1.2.14 integer prg\_chebyshev\_mod::chebdata\_type::ndim

Definition at line 31 of file prg\_chebyshev\_mod.F90.

10.1.2.15 integer prg\_chebyshev\_mod::chebdata\_type::npts

Definition at line 32 of file prg chebyshev mod.F90.

10.1.2.16 real(dp) prg\_chebyshev\_mod::chebdata\_type::threshold

Definition at line 34 of file prg chebyshev mod.F90.

10.1.2.17 logical prg\_chebyshev\_mod::chebdata\_type::trkfunc

Definition at line 35 of file prg\_chebyshev\_mod.F90.

10.1.2.18 integer prg\_chebyshev\_mod::chebdata\_type::verbose

Definition at line 31 of file prg\_chebyshev\_mod.F90.

The documentation for this type was generated from the following file:

• /home/christian/qmd-progress/src/prg\_chebyshev\_mod.F90

## 10.2 prg\_system\_mod::estruct\_type Type Reference

Electronic structure type.

### **Public Attributes**

integer norbs

Number of orbitals of the system.

integer nel

Number of electrons.

• integer, dimension(:,:), allocatable hindex

Hindex.

• type(bml\_matrix\_t) ham

SCC-Hamiltonian of the system.

• type(bml\_matrix\_t) ham0

Hamiltonian of the system.

• type(bml\_matrix\_t) oham

Orthogonalized Hamiltonian.

type(bml\_matrix\_t) over

Overlap matrix of the system.

• type(bml\_matrix\_t) rho

Density matrix of the system.

type(bml\_matrix\_t) orho

Orthogonalized density matrix.

• type(bml\_matrix\_t) zmat

Congruence transformation.

• real(dp), dimension(:), allocatable coul\_pot\_r

Real Coulombic contribution.

real(dp), dimension(:), allocatable coul\_pot\_k

Reciprocal Coulombic contribution.

• real(dp), dimension(:,:), allocatable skforce

Slater Koster force.

• real(dp), dimension(:,:), allocatable fpul

Pulay force.

• real(dp), dimension(:,:), allocatable fscoul

Nonorthogonal Coulombic force.

real(dp) eband

Band energy.

## 10.2.1 Detailed Description

Electronic structure type.

The electronic structure type.

Definition at line 19 of file prg\_system\_mod.F90.

## 10.2.2 Member Data Documentation

10.2.2.1 real(dp), dimension(:), allocatable prg\_system\_mod::estruct\_type::coul\_pot\_k

Reciprocal Coulombic contribution.

Definition at line 55 of file prg\_system\_mod.F90.

10.2.2.2 real(dp), dimension(:), allocatable prg\_system\_mod::estruct\_type::coul\_pot\_r

Real Coulombic contribution.

Definition at line 52 of file prg\_system\_mod.F90.

10.2.2.3 real(dp) prg\_system\_mod::estruct\_type::eband

Band energy.

Definition at line 67 of file prg\_system\_mod.F90.

10.2.2.4 real(dp), dimension(:,:), allocatable prg\_system\_mod::estruct\_type::fpul

Pulay force.

Definition at line 61 of file prg\_system\_mod.F90.

10.2.2.5 real(dp), dimension(:,:), allocatable prg\_system\_mod::estruct\_type::fscoul

Nonorthogonal Coulombic force.

Definition at line 64 of file prg\_system\_mod.F90.

10.2.2.6 type(bml\_matrix\_t) prg\_system\_mod::estruct\_type::ham

SCC-Hamiltonian of the system.

Definition at line 31 of file prg\_system\_mod.F90.

10.2.2.7 type(bml\_matrix\_t) prg\_system\_mod::estruct\_type::ham0

Hamiltonian of the system.

Definition at line 34 of file prg system mod.F90.

10.2.2.8 integer, dimension(:,:), allocatable prg\_system\_mod::estruct\_type::hindex

Hindex.

Definition at line 28 of file prg\_system\_mod.F90.

10.2.2.9 integer prg\_system\_mod::estruct\_type::nel

Number of electrons.

Definition at line 25 of file prg\_system\_mod.F90.

10.2.2.10 integer prg\_system\_mod::estruct\_type::norbs

Number of orbitals of the system.

Definition at line 22 of file prg\_system\_mod.F90.

10.2.2.11 type(bml\_matrix\_t) prg\_system\_mod::estruct\_type::oham

Orthogonalized Hamiltonian.

Definition at line 37 of file prg\_system\_mod.F90.

10.2.2.12 type(bml\_matrix\_t) prg\_system\_mod::estruct\_type::orho

Orthogonalized density matrix.

Definition at line 46 of file prg\_system\_mod.F90.

10.2.2.13 type(bml\_matrix\_t) prg\_system\_mod::estruct\_type::over

Overlap matrix of the system.

Definition at line 40 of file prg\_system\_mod.F90.

10.2.2.14 type(bml\_matrix\_t) prg\_system\_mod::estruct\_type::rho

Density matrix of the system.

Definition at line 43 of file prg\_system\_mod.F90.

10.2.2.15 real(dp), dimension(:,:), allocatable prg\_system\_mod::estruct\_type::skforce

Slater Koster force.

Definition at line 58 of file prg\_system\_mod.F90.

10.2.2.16 type(bml\_matrix\_t) prg\_system\_mod::estruct\_type::zmat

Congruence transformation.

Definition at line 49 of file prg\_system\_mod.F90.

The documentation for this type was generated from the following file:

/home/christian/qmd-progress/src/prg\_system\_mod.F90

## 10.3 prg\_genz\_mod::genzspdata Type Reference

contains the data for the genZ driver.

### **Public Member Functions**

- procedure prg\_init
- procedure prg\_generate
- procedure prg\_allocatezspmat

### **Public Attributes**

- integer verbose
- integer nfirst
- integer nrefi
- integer nreff
- · real(dp) numthresi
- · real(dp) numthresf
- · logical integration

### 10.3.1 Detailed Description

contains the data for the genZ driver.

Definition at line 65 of file prg\_genz\_mod.F90.

#### 10.3.2 Member Function/Subroutine Documentation

```
10.3.2.1 procedure prg_genz_mod::genzspdata::prg_allocatezspmat ( )
```

Definition at line 77 of file prg\_genz\_mod.F90.

10.3.2.2 procedure prg\_genz\_mod::genzspdata::prg\_generate ( )

Definition at line 76 of file prg\_genz\_mod.F90.

10.3.2.3 procedure prg\_genz\_mod::genzspdata::prg\_init ( )

Definition at line 75 of file prg\_genz\_mod.F90.

### 10.3.3 Member Data Documentation

10.3.3.1 logical prg\_genz\_mod::genzspdata::integration

Definition at line 73 of file prg\_genz\_mod.F90.

10.3.3.2 integer prg\_genz\_mod::genzspdata::nfirst Definition at line 68 of file prg\_genz\_mod.F90. 10.3.3.3 integer prg\_genz\_mod::genzspdata::nreff Definition at line 70 of file prg\_genz\_mod.F90. 10.3.3.4 integer prg\_genz\_mod::genzspdata::nrefi Definition at line 69 of file prg\_genz\_mod.F90. 10.3.3.5 real(dp) prg\_genz\_mod::genzspdata::numthresf Definition at line 72 of file prg\_genz\_mod.F90. 10.3.3.6 real(dp) prg\_genz\_mod::genzspdata::numthresi Definition at line 71 of file prg\_genz\_mod.F90. 10.3.3.7 integer prg\_genz\_mod::genzspdata::verbose Definition at line 67 of file prg\_genz\_mod.F90. The documentation for this type was generated from the following file: /home/christian/qmd-progress/src/prg\_genz\_mod.F90

# 10.4 prg\_genz\_mod::genzspinp Type Reference

Input for the genz driver. This type controlls all the variables that are needed by genz.

### **Public Attributes**

· integer verbose

To have different levels of verbose.

· integer nfirst

!Lentgth of the "firsts iteration period".

integer nrefi

!Initial number of recursive refinements.

· integer nreff

!Initial number of recursive refinements.

• real(dp) numthresi

Initial threshold value.

• real(dp) numthresf

Final threshold value.

· logical integration

If we want to do XL integration scheme for Z.

· integer igenz

To keep track of the genz iterations.

logical zsp

Logical variable to compute in sparse or dense mode.

· integer mdim

Max nonzero elements per row for every row see [1].

character(20) bml\_type

Matrix format (Dense or Ellpack).

### 10.4.1 Detailed Description

Input for the genz driver. This type controlls all the variables that are needed by genz.

Definition at line 26 of file prg\_genz\_mod.F90.

### 10.4.2 Member Data Documentation

10.4.2.1 character(20) prg\_genz\_mod::genzspinp::bml\_type

Matrix format (Dense or Ellpack).

Definition at line 59 of file prg\_genz\_mod.F90.

10.4.2.2 integer prg\_genz\_mod::genzspinp::igenz

To keep track of the genz iterations.

Definition at line 50 of file prg\_genz\_mod.F90.

Generated by Doxygen

10.4.2.3 logical prg\_genz\_mod::genzspinp::integration If we want to do XL integration scheme for Z. Definition at line 47 of file prg\_genz\_mod.F90. 10.4.2.4 integer prg\_genz\_mod::genzspinp::mdim Max nonzero elements per row for every row see [1]. Definition at line 56 of file prg\_genz\_mod.F90. 10.4.2.5 integer prg\_genz\_mod::genzspinp::nfirst !Lentgth of the "firsts iteration period". Definition at line 32 of file prg\_genz\_mod.F90. 10.4.2.6 integer prg\_genz\_mod::genzspinp::nreff !Initial number of recursive refinements. Definition at line 38 of file prg\_genz\_mod.F90. 10.4.2.7 integer prg\_genz\_mod::genzspinp::nrefi !Initial number of recursive refinements. Definition at line 35 of file prg\_genz\_mod.F90. 10.4.2.8 real(dp) prg\_genz\_mod::genzspinp::numthresf Final threshold value. Definition at line 44 of file prg genz mod.F90. 10.4.2.9 real(dp) prg\_genz\_mod::genzspinp::numthresi Initial threshold value. Definition at line 41 of file prg\_genz\_mod.F90.

10.4.2.10 integer prg\_genz\_mod::genzspinp::verbose

To have different levels of verbose.

Definition at line 29 of file prg\_genz\_mod.F90.

10.4.2.11 logical prg\_genz\_mod::genzspinp::zsp

Logical variable to compute in sparse or dense mode.

Definition at line 53 of file prg\_genz\_mod.F90.

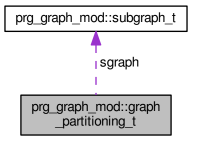
The documentation for this type was generated from the following file:

/home/christian/qmd-progress/src/prg\_genz\_mod.F90

## 10.5 prg\_graph\_mod::graph\_partitioning\_t Type Reference

Trace per iteration.

Collaboration diagram for prg\_graph\_mod::graph\_partitioning\_t:



### **Private Attributes**

• character(len=100) pname

Partition name.

· integer myrank

Local processor.

• integer totalprocs

Number of processors.

integer totalparts

Total number of global partitions.

• integer totalnodes

Total number of global groups, nodes (or matrix rows)

• integer totalnodes2

Total number of global nodes (or matrix rows)

integer globalpartmin

Minimum global part number.

integer globalpartmax

Maximum global part number.

· integer globalpartextent

Total global parts.

• integer, dimension(:), allocatable localpartmin

Minimum part per processor.

• integer, dimension(:), allocatable localpartmax

Maximum part per processor.

• integer, dimension(:), allocatable localpartextent

Number of parts per processor.

integer, dimension(:), allocatable order

Original ordering if required.

• integer, dimension(:), allocatable reorder

Reordering if required.

· integer nparts

Total number of local partitions.

• integer, dimension(:), allocatable nnodesinpart

Number of nodes in each local partition.

• integer, dimension(:), allocatable nnodesinpartall

Number of nodes in each partition.

• integer, dimension(100) pp

Sequence for SP2.

integer maxiter

Number of SP2 iterations.

• real(dp) ehomo

Homo value.

• real(dp) elumo

Lumo value.

· real(dp) mineval

Min eval for prg\_normalize.

· real(dp) maxeval

Max eval for prg\_normalize.

real(dp), dimension(100) vv

Trace per iteration.

• type(subgraph\_t), dimension(:), allocatable sgraph

Subgraph details.

## 10.5.1 Detailed Description

Trace per iteration.

Graph partitioning type

Definition at line 57 of file prg\_graph\_mod.F90.

### 10.5.2 Member Data Documentation

10.5.2.1 real(dp) prg\_graph\_mod::graph\_partitioning\_t::ehomo [private]

Homo value.

Definition at line 117 of file prg\_graph\_mod.F90.

10.5.2.2 real(dp) prg\_graph\_mod::graph\_partitioning\_t::elumo [private] Lumo value. Definition at line 120 of file prg\_graph\_mod.F90. **10.5.2.3** integer prg\_graph\_mod::graph\_partitioning\_t::globalpartextent [private] Total global parts. Definition at line 84 of file prg\_graph\_mod.F90. **10.5.2.4** integer prg\_graph\_mod::graph\_partitioning\_t::globalpartmax [private] Maximum global part number. Definition at line 81 of file prg\_graph\_mod.F90. **10.5.2.5** integer prg\_graph\_mod::graph\_partitioning\_t::globalpartmin [private] Minimum global part number. Definition at line 78 of file prg\_graph\_mod.F90. **10.5.2.6** integer, dimension(:), allocatable prg\_graph\_mod::graph\_partitioning\_t::localpartextent [private] Number of parts per processor. Definition at line 93 of file prg\_graph\_mod.F90. 10.5.2.7 integer, dimension(:), allocatable prg\_graph\_mod::graph\_partitioning\_t::localpartmax [private] Maximum part per processor. Definition at line 90 of file prg graph mod.F90. 10.5.2.8 integer, dimension(:), allocatable prg\_graph\_mod::graph\_partitioning\_t::localpartmin [private] Minimum part per processor. Definition at line 87 of file prg\_graph\_mod.F90. 10.5.2.9 real(dp) prg\_graph\_mod::graph\_partitioning\_t::maxeval [private] Max eval for prg\_normalize.

Generated by Doxygen

Definition at line 126 of file prg\_graph\_mod.F90.

```
10.5.2.10 integer prg_graph_mod::graph_partitioning_t::maxiter [private]
Number of SP2 iterations.
Definition at line 114 of file prg_graph_mod.F90.
10.5.2.11 real(dp) prg_graph_mod::graph_partitioning_t::mineval [private]
Min eval for prg_normalize.
Definition at line 123 of file prg_graph_mod.F90.
10.5.2.12 integer prg_graph_mod::graph_partitioning_t::myrank [private]
Local processor.
Definition at line 63 of file prg graph mod.F90.
10.5.2.13 integer, dimension(:), allocatable prg_graph_mod::graph_partitioning_t::nnodesinpart [private]
Number of nodes in each local partition.
Definition at line 105 of file prg_graph_mod.F90.
10.5.2.14 integer, dimension(:), allocatable prg_graph_mod::graph_partitioning_t::nnodesinpartall [private]
Number of nodes in each partition.
Definition at line 108 of file prg_graph_mod.F90.
10.5.2.15 integer prg_graph_mod::graph_partitioning_t::nparts [private]
Total number of local partitions.
Definition at line 102 of file prg graph mod.F90.
10.5.2.16 integer, dimension(:), allocatable prg_graph_mod::graph_partitioning_t::order [private]
Original ordering if required.
Definition at line 96 of file prg_graph_mod.F90.
10.5.2.17 character(len=100) prg_graph_mod::graph_partitioning_t::pname [private]
Partition name.
```

Definition at line 60 of file prg\_graph\_mod.F90.

```
10.5.2.18 integer, dimension(100) prg_graph_mod::graph_partitioning_t::pp [private]
Sequence for SP2.
Definition at line 111 of file prg_graph_mod.F90.
10.5.2.19 integer, dimension(:), allocatable prg_graph_mod::graph_partitioning_t::reorder [private]
Reordering if required.
Definition at line 99 of file prg_graph_mod.F90.
10.5.2.20 type (subgraph_t), dimension(:), allocatable prg_graph_mod::graph_partitioning_t::sgraph [private]
Subgraph details.
Definition at line 132 of file prg_graph_mod.F90.
10.5.2.21 integer prg_graph_mod::graph_partitioning_t::totalnodes [private]
Total number of global groups, nodes (or matrix rows)
Definition at line 72 of file prg_graph_mod.F90.
10.5.2.22 integer prg_graph_mod::graph_partitioning_t::totalnodes2 [private]
Total number of global nodes (or matrix rows)
Definition at line 75 of file prg_graph_mod.F90.
10.5.2.23 integer prg_graph_mod::graph_partitioning_t::totalparts [private]
Total number of global partitions.
Definition at line 69 of file prg_graph_mod.F90.
10.5.2.24 integer prg_graph_mod::graph_partitioning_t::totalprocs [private]
Number of processors.
```

Definition at line 66 of file prg\_graph\_mod.F90.

10.5.2.25 real(dp), dimension(100) prg\_graph\_mod::graph\_partitioning\_t::vv [private]

Trace per iteration.

Definition at line 129 of file prg\_graph\_mod.F90.

The documentation for this type was generated from the following file:

/home/christian/qmd-progress/src/prg\_graph\_mod.F90

## 10.6 prg\_graphsp2parser\_mod::gsp2data\_type Type Reference

General SP2 solver type.

### **Public Attributes**

- character(20) jobname
- character(50) hamfile
- · integer verbose
- · integer minsp2iter
- · integer maxsp2iter
- integer nodesperpart
- integer natoms
- integer partition\_count
- real(dp) sp2tol
- real(dp) threshold
- real(dp) bndfil
- real(dp) gthreshold
- real(dp) errlimit
- · integer mdim
- integer ndim
- character, dimension(3) sdim
- real(dp), dimension(3) pdim
- character(20) bml\_type
- character(10) sp2conv
- character(10) graph\_element
- character(10) partition\_type
- character(10) partition\_refinement
- logical double\_jump
- real(dp) covgfact
- real(dp) nlgcut
- integer parteach

## 10.6.1 Detailed Description

General SP2 solver type.

Definition at line 26 of file prg\_graphsp2parser\_mod.F90.

### 10.6.2 Member Data Documentation

10.6.2.1 character(20) prg\_graphsp2parser\_mod::gsp2data\_type::bml\_type

Definition at line 44 of file prg\_graphsp2parser\_mod.F90.

10.6.2.2 real(dp) prg\_graphsp2parser\_mod::gsp2data\_type::bndfil

Definition at line 37 of file prg\_graphsp2parser\_mod.F90.

10.6.2.3 real(dp) prg\_graphsp2parser\_mod::gsp2data\_type::covgfact

Definition at line 50 of file prg\_graphsp2parser\_mod.F90.

10.6.2.4 logical prg\_graphsp2parser\_mod::gsp2data\_type::double\_jump

Definition at line 49 of file prg\_graphsp2parser\_mod.F90.

10.6.2.5 real(dp) prg\_graphsp2parser\_mod::gsp2data\_type::errlimit

Definition at line 39 of file prg\_graphsp2parser\_mod.F90.

10.6.2.6 character(10) prg\_graphsp2parser\_mod::gsp2data\_type::graph\_element

Definition at line 46 of file prg\_graphsp2parser\_mod.F90.

 $10.6.2.7 \quad real (dp) \ prg\_graphsp2 parser\_mod::gsp2 data\_type::gthreshold$ 

Definition at line 38 of file prg\_graphsp2parser\_mod.F90.

10.6.2.8 character(50) prg\_graphsp2parser\_mod::gsp2data\_type::hamfile

Definition at line 28 of file prg\_graphsp2parser\_mod.F90.

10.6.2.9 character(20) prg\_graphsp2parser\_mod::gsp2data\_type::jobname

Definition at line 27 of file prg\_graphsp2parser\_mod.F90.

10.6.2.10 integer prg\_graphsp2parser\_mod::gsp2data\_type::maxsp2iter

Definition at line 31 of file prg\_graphsp2parser\_mod.F90.

10.6.2.11 integer prg\_graphsp2parser\_mod::gsp2data\_type::mdim Definition at line 40 of file prg\_graphsp2parser\_mod.F90. 10.6.2.12 integer prg\_graphsp2parser\_mod::gsp2data\_type::minsp2iter Definition at line 30 of file prg\_graphsp2parser\_mod.F90. 10.6.2.13 integer prg\_graphsp2parser\_mod::gsp2data\_type::natoms Definition at line 33 of file prg\_graphsp2parser\_mod.F90. 10.6.2.14 integer prg\_graphsp2parser\_mod::gsp2data\_type::ndim Definition at line 41 of file prg\_graphsp2parser\_mod.F90. 10.6.2.15 real(dp) prg\_graphsp2parser\_mod::gsp2data\_type::nlgcut Definition at line 51 of file prg\_graphsp2parser\_mod.F90. 10.6.2.16 integer prg\_graphsp2parser\_mod::gsp2data\_type::nodesperpart Definition at line 32 of file prg\_graphsp2parser\_mod.F90. 10.6.2.17 integer prg\_graphsp2parser\_mod::gsp2data\_type::parteach Definition at line 52 of file prg\_graphsp2parser\_mod.F90. 10.6.2.18 integer prg\_graphsp2parser\_mod::gsp2data\_type::partition\_count Definition at line 34 of file prg\_graphsp2parser\_mod.F90. 10.6.2.19 character(10) prg\_graphsp2parser\_mod::gsp2data\_type::partition\_refinement Definition at line 48 of file prg\_graphsp2parser\_mod.F90. 10.6.2.20 character(10) prg\_graphsp2parser\_mod::gsp2data\_type::partition\_type

Definition at line 47 of file prg\_graphsp2parser\_mod.F90.

10.6.2.21 real(dp), dimension(3) prg\_graphsp2parser\_mod::gsp2data\_type::pdim

Definition at line 43 of file prg\_graphsp2parser\_mod.F90.

10.6.2.22 character, dimension(3) prg\_graphsp2parser\_mod::gsp2data\_type::sdim

Definition at line 42 of file prg\_graphsp2parser\_mod.F90.

10.6.2.23 character(10) prg\_graphsp2parser\_mod::gsp2data\_type::sp2conv

Definition at line 45 of file prg\_graphsp2parser\_mod.F90.

10.6.2.24 real(dp) prg\_graphsp2parser\_mod::gsp2data\_type::sp2tol

Definition at line 35 of file prg\_graphsp2parser\_mod.F90.

10.6.2.25 real(dp) prg\_graphsp2parser\_mod::gsp2data\_type::threshold

Definition at line 36 of file prg\_graphsp2parser\_mod.F90.

10.6.2.26 integer prg\_graphsp2parser\_mod::gsp2data\_type::verbose

Definition at line 29 of file prg\_graphsp2parser\_mod.F90.

The documentation for this type was generated from the following file:

/home/christian/qmd-progress/src/prg\_graphsp2parser\_mod.F90

## 10.7 prg\_pulaymixer\_mod::mx\_type Type Reference

### **Public Attributes**

• character(20) mixertype

Type or mixing scheme to be used (Linear or Pulay)

integer verbose

Verbosity level.

· integer mpulay

Pulay dimension for matrix.

real(dp) mixcoeff

Coefficient for mixing.

logical mixeron

Mixer on or off (Not implemented)

## 10.7.1 Detailed Description

Definition at line 16 of file prg\_pulaymixer\_mod.F90.

### 10.7.2 Member Data Documentation

10.7.2.1 real(dp) prg\_pulaymixer\_mod::mx\_type::mixcoeff

Coefficient for mixing.

Definition at line 28 of file prg\_pulaymixer\_mod.F90.

10.7.2.2 logical prg\_pulaymixer\_mod::mx\_type::mixeron

Mixer on or off (Not implemented)

Definition at line 31 of file prg\_pulaymixer\_mod.F90.

10.7.2.3 character(20) prg\_pulaymixer\_mod::mx\_type::mixertype

Type or mixing scheme to be used (Linear or Pulay)

Definition at line 19 of file prg\_pulaymixer\_mod.F90.

10.7.2.4 integer prg\_pulaymixer\_mod::mx\_type::mpulay

Pulay dimension for matrix.

Definition at line 25 of file prg\_pulaymixer\_mod.F90.

10.7.2.5 integer prg\_pulaymixer\_mod::mx\_type::verbose

Verbosity level.

Definition at line 22 of file prg\_pulaymixer\_mod.F90.

The documentation for this type was generated from the following file:

• /home/christian/qmd-progress/src/prg\_pulaymixer\_mod.F90

## 10.8 prg\_extras\_mod::prg\_memory\_consumption Interface Reference

### **Private Member Functions**

• subroutine prg\_memory\_consumption (vm\_peak, vm\_size, pid, ppid)

## 10.8.1 Detailed Description

Definition at line 15 of file prg\_extras\_mod.F90.

### 10.8.2 Constructor & Destructor Documentation

10.8.2.1 subroutine prg\_extras\_mod::prg\_memory\_consumption::prg\_memory\_consumption ( integer(c\_long\_long), intent(inout) vm\_peak, integer(c\_long\_long), intent(inout) vm\_size, integer(c\_long\_long), intent(inout) ppid, integer(c\_long\_long), intent(inout) ppid ) [private]

Definition at line 17 of file prg\_extras\_mod.F90.

The documentation for this interface was generated from the following file:

/home/christian/qmd-progress/src/prg\_extras\_mod.F90

## 10.9 prg\_parallel\_mod::rankreducedata\_t Type Reference

Data structure for rection over MPI ranks.

### **Private Attributes**

real(dp) val

Data value.

MPI rank.

integer rank

## 10.9.1 Detailed Description

Data structure for rection over MPI ranks.

Definition at line 72 of file prg\_parallel\_mod.F90.

### 10.9.2 Member Data Documentation

10.9.2.1 integer prg\_parallel\_mod::rankreducedata\_t::rank [private]

MPI rank.

Definition at line 78 of file prg\_parallel\_mod.F90.

10.9.2.2 real(dp) prg\_parallel\_mod::rankreducedata\_t::val [private]

Data value.

Definition at line 75 of file prg\_parallel\_mod.F90.

The documentation for this type was generated from the following file:

/home/christian/qmd-progress/src/prg parallel mod.F90

## 10.10 prg\_response\_mod::respdata\_type Type Reference

### **Public Attributes**

- character(20) respmode
- character(20) typeofpert
- character(20) bmltype
- integer mdim
- real(dp) numthresh
- · logical computedipole
- logical getresponse
- real(dp) fieldintensity
- real(dp), dimension(3) field

## 10.10.1 Detailed Description

Definition at line 21 of file prg\_response\_mod.F90.

### 10.10.2 Member Data Documentation

10.10.2.1 character(20) prg\_response\_mod::respdata\_type::bmltype

Definition at line 24 of file prg\_response\_mod.F90.

10.10.2.2 logical prg\_response\_mod::respdata\_type::computedipole

Definition at line 27 of file prg\_response\_mod.F90.

10.10.2.3 real(dp), dimension(3) prg\_response\_mod::respdata\_type::field

Definition at line 30 of file prg\_response\_mod.F90.

10.10.2.4 real(dp) prg\_response\_mod::respdata\_type::fieldintensity

Definition at line 29 of file prg\_response\_mod.F90.

10.10.2.5 logical prg\_response\_mod::respdata\_type::getresponse

Definition at line 28 of file prg\_response\_mod.F90.

10.10.2.6 integer prg\_response\_mod::respdata\_type::mdim

Definition at line 25 of file prg response mod.F90.

10.10.2.7 real(dp) prg\_response\_mod::respdata\_type::numthresh

Definition at line 26 of file prg\_response\_mod.F90.

10.10.2.8 character(20) prg\_response\_mod::respdata\_type::respmode

Definition at line 22 of file prg response mod.F90.

10.10.2.9 character(20) prg\_response\_mod::respdata\_type::typeofpert

Definition at line 23 of file prg\_response\_mod.F90.

The documentation for this type was generated from the following file:

/home/christian/qmd-progress/src/prg\_response\_mod.F90

## 10.11 prg\_syrotation\_mod::rotation\_type Type Reference

Rotation type.

#### **Public Attributes**

- character(20) jobname
- character(50) typeofrot
- integer patom1

Atomic point to determine the initial orientation.

integer patom2

Atomic point to determine initial orientation.

· integer catom

Atomic point to determine the rotation center.

integer catom2

Atomic point to determine a second rotation center.

• real(dp), dimension(3) pq1

Point to determine initial orientation.

• real(dp), dimension(3) pq2

Point to determine final orientation.

• real(dp), dimension(3) v1

Initial orientation.

• real(dp), dimension(3) v2

Final orientation.

• real(dp), dimension(3) vq

Center of rotation.

• integer, dimension(2) rotate\_atoms

First and last rotated atom in the list.

## 10.11.1 Detailed Description

Rotation type.

Definition at line 14 of file prg\_syrotation\_mod.F90.

#### 10.11.2 Member Data Documentation

10.11.2.1 integer prg\_syrotation\_mod::rotation\_type::catom

Atomic point to determine the rotation center.

Definition at line 22 of file prg\_syrotation\_mod.F90.

10.11.2.2 integer prg\_syrotation\_mod::rotation\_type::catom2

Atomic point to determine a second rotation center.

Definition at line 24 of file prg syrotation mod.F90.

10.11.2.3 character(20) prg\_syrotation\_mod::rotation\_type::jobname

Definition at line 15 of file prg\_syrotation\_mod.F90.

10.11.2.4 integer prg\_syrotation\_mod::rotation\_type::patom1

Atomic point to determine the initial orientation.

Definition at line 18 of file prg\_syrotation\_mod.F90.

10.11.2.5 integer prg\_syrotation\_mod::rotation\_type::patom2

Atomic point to determine initial orientation.

Definition at line 20 of file prg\_syrotation\_mod.F90.

10.11.2.6 real(dp), dimension(3) prg\_syrotation\_mod::rotation\_type::pq1

Point to determine initial orientation.

Definition at line 26 of file prg\_syrotation\_mod.F90.

10.11.2.7 real(dp), dimension(3) prg\_syrotation\_mod::rotation\_type::pq2

Point to determine final orientation.

Definition at line 28 of file prg syrotation mod.F90.

10.11.2.8 integer, dimension(2) prg\_syrotation\_mod::rotation\_type::rotate\_atoms

First and last rotated atom in the list.

Definition at line 36 of file prg\_syrotation\_mod.F90.

10.11.2.9 character(50) prg\_syrotation\_mod::rotation\_type::typeofrot

Definition at line 16 of file prg\_syrotation\_mod.F90.

10.11.2.10 real(dp), dimension(3) prg\_syrotation\_mod::rotation\_type::v1

Initial orientation.

Definition at line 30 of file prg\_syrotation\_mod.F90.

10.11.2.11 real(dp), dimension(3) prg\_syrotation\_mod::rotation\_type::v2

Final orientation.

Definition at line 32 of file prg\_syrotation\_mod.F90.

10.11.2.12 real(dp), dimension(3) prg\_syrotation\_mod::rotation\_type::vq

Center of rotation.

Definition at line 34 of file prg\_syrotation\_mod.F90.

The documentation for this type was generated from the following file:

/home/christian/qmd-progress/src/prg\_syrotation\_mod.F90

## 10.12 prg\_sp2parser\_mod::sp2data\_type Type Reference

General SP2 solver type.

### **Public Attributes**

- character(20) jobname
- integer verbose
- integer minsp2iter
- integer maxsp2iter
- real(dp) sp2tol
- real(dp) threshold
- real(dp) bndfil
- integer mdim
- integer ndim
- character, dimension(3) sdim
- real(dp), dimension(3) pdim
- character(20) bml\_type
- character(10) sp2conv
- character(10) flavor

### 10.12.1 Detailed Description

General SP2 solver type.

Definition at line 26 of file prg\_sp2parser\_mod.F90.

### 10.12.2 Member Data Documentation

10.12.2.1 character(20) prg\_sp2parser\_mod::sp2data\_type::bml\_type

Definition at line 38 of file prg\_sp2parser\_mod.F90.

10.12.2.2 real(dp) prg\_sp2parser\_mod::sp2data\_type::bndfil

Definition at line 33 of file prg\_sp2parser\_mod.F90.

10.12.2.3 character(10) prg\_sp2parser\_mod::sp2data\_type::flavor

Definition at line 40 of file prg\_sp2parser\_mod.F90.

10.12.2.4 character(20) prg\_sp2parser\_mod::sp2data\_type::jobname

Definition at line 27 of file prg\_sp2parser\_mod.F90.

10.12.2.5 integer prg\_sp2parser\_mod::sp2data\_type::maxsp2iter

Definition at line 30 of file prg\_sp2parser\_mod.F90.

10.12.2.6 integer prg\_sp2parser\_mod::sp2data\_type::mdim

Definition at line 34 of file prg\_sp2parser\_mod.F90.

10.12.2.7 integer prg\_sp2parser\_mod::sp2data\_type::minsp2iter

Definition at line 29 of file prg sp2parser mod.F90.

10.12.2.8 integer prg\_sp2parser\_mod::sp2data\_type::ndim

Definition at line 35 of file prg\_sp2parser\_mod.F90.

10.12.2.9 real(dp), dimension(3) prg\_sp2parser\_mod::sp2data\_type::pdim

Definition at line 37 of file prg\_sp2parser\_mod.F90.

10.12.2.10 character, dimension(3) prg\_sp2parser\_mod::sp2data\_type::sdim

Definition at line 36 of file prg\_sp2parser\_mod.F90.

10.12.2.11 character(10) prg\_sp2parser\_mod::sp2data\_type::sp2conv

Definition at line 39 of file prg\_sp2parser\_mod.F90.

10.12.2.12 real(dp) prg\_sp2parser\_mod::sp2data\_type::sp2tol

Definition at line 31 of file prg\_sp2parser\_mod.F90.

10.12.2.13 real(dp) prg\_sp2parser\_mod::sp2data\_type::threshold

Definition at line 32 of file prg\_sp2parser\_mod.F90.

10.12.2.14 integer prg\_sp2parser\_mod::sp2data\_type::verbose

Definition at line 28 of file prg sp2parser mod.F90.

The documentation for this type was generated from the following file:

/home/christian/qmd-progress/src/prg\_sp2parser\_mod.F90

# 10.13 prg\_graph\_mod::subgraph\_t Type Reference

Subgraph type.

### **Private Attributes**

integer part

Partition number.

· integer hsize

Size of original matrix (h x h)

• integer Isize

Size of full subgraph (I x I)

integer Ilsize

Size of core subgraph.

• integer, dimension(:), allocatable core\_halo\_index

Indeces from original matrix for subgraph core+halo extraction.

• integer, dimension(:), allocatable nodeinpart Nodes in this partition.

### 10.13.1 Detailed Description

Subgraph type.

Definition at line 31 of file prg\_graph\_mod.F90.

### 10.13.2 Member Data Documentation

10.13.2.1 integer, dimension(:), allocatable prg\_graph\_mod::subgraph\_t::core\_halo\_index [private]

Indeces from original matrix for subgraph core+halo extraction.

Definition at line 46 of file prg\_graph\_mod.F90.

10.13.2.2 integer prg\_graph\_mod::subgraph\_t::hsize [private]

Size of original matrix (h x h)

Definition at line 37 of file prg\_graph\_mod.F90.

10.13.2.3 integer prg\_graph\_mod::subgraph\_t::llsize [private]

Size of core subgraph.

Definition at line 43 of file prg\_graph\_mod.F90.

10.13.2.4 integer prg\_graph\_mod::subgraph\_t::lsize [private]

Size of full subgraph (I x I)

Definition at line 40 of file prg\_graph\_mod.F90.

10.13.2.5 integer, dimension(:), allocatable prg\_graph\_mod::subgraph\_t::nodeinpart [private]

Nodes in this partition.

Definition at line 49 of file prg\_graph\_mod.F90.

10.13.2.6 integer prg\_graph\_mod::subgraph\_t::part [private]

Partition number.

Definition at line 34 of file prg\_graph\_mod.F90.

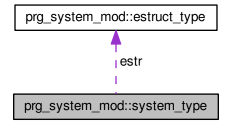
The documentation for this type was generated from the following file:

• /home/christian/qmd-progress/src/prg\_graph\_mod.F90

## 10.14 prg\_system\_mod::system\_type Type Reference

System type.

Collaboration diagram for prg\_system\_mod::system\_type:



### **Public Attributes**

· integer nats

Number of atoms of the system.

character(2), dimension(:), allocatable symbol

Chemical Symbols for every atom of the system. Symbol can be recovered using ptable module and calling the following routine:

• integer, dimension(:), allocatable atomic number

Atomic number for every atom in the system.

real(dp), dimension(:,:), allocatable coordinate

Coordinates of every atom in the system. Allocation:

real(dp), dimension(:,:), allocatable velocity

Velocities for every atom in the system. Allocation:

real(dp), dimension(:,:), allocatable force

Forces acting on every atom in the system. Allocation:

real(dp), dimension(:), allocatable net\_charge

Charges of every atom in the system. Allocation:

• real(dp), dimension(:), allocatable mass

Mass of every atom in the system. These can be automatically loaded by using the structures of the ptable mod:

real(dp), dimension(:,:), allocatable lattice vector

Lattice vectors of the system. Use the prg\_vectors\_to\_parameters and parameters\_to\_vector to transform from lattice vector to lattice parameters. Allocation:

• real(dp), dimension(:,:), allocatable recip vector

Reciprocal vectors of the system. Allocation:

real(dp) volr

Volume of the system (direct space).

real(dp) volk

Volume of the system (direct space).

integer nsp

Number of different species. Number of species or number of different antom types (symbols) in the system. This integer is alwas less or equal than the total number of atoms ( $nsp \le nats$ ). This information can also be found in tbparams structure and the following equality holds:

• integer, dimension(:), allocatable spindex

Species index. It gives the species index of a particulat atom. Allocation:

character(2), dimension(:), allocatable splist

Species symbol list. A list with the different species e.g. H, C, N, etc with the order corresponding to the appearence in systemsymbol. Allocation:

integer, dimension(:), allocatable spatnum

Species atomic number list. A list with the atomic numbers for every species Allocation:

real(dp), dimension(:), allocatable spmass

Species mass list. A list with the atomic mass for every species Allocation:

real(dp), dimension(:), allocatable userdef

User define field.

• integer, dimension(:), allocatable resindex

Residue index.

• type(estruct\_type) estr

Electronic structure.

### 10.14.1 Detailed Description

System type.

The molecular system type.

Definition at line 72 of file prg\_system\_mod.F90.

### 10.14.2 Member Data Documentation

10.14.2.1 integer, dimension(:), allocatable prg\_system\_mod::system\_type::atomic\_number

Atomic number for every atom in the system.

Definition at line 86 of file prg\_system\_mod.F90.

10.14.2.2 real(dp), dimension(:,:), allocatable prg\_system\_mod::system\_type::coordinate

Coordinates of every atom in the system. Allocation:

```
coordinate(3, nats)
```

Definition at line 91 of file prg\_system\_mod.F90.

10.14.2.3 type(estruct\_type) prg\_system\_mod::system\_type::estr

Electronic structure.

Definition at line 184 of file prg system mod.F90.

10.14.2.4 real(dp), dimension(:,:), allocatable prg\_system\_mod::system\_type::force

Forces acting on every atom in the system. Allocation:

```
force(3,nats)
```

Definition at line 101 of file prg\_system\_mod.F90.

10.14.2.5 real(dp), dimension(:,:), allocatable prg\_system\_mod::system\_type::lattice\_vector

Lattice vectors of the system. Use the prg\_vectors\_to\_parameters and parameters\_to\_vector to transform from lattice vector to lattice parameters. Allocation:

```
lattice_vector(3,3)
v1 = lattice_vector(1,:)
v2 = lattice_vector(2,:)
v3 = lattice_vector(3,:)
```

Definition at line 123 of file prg\_system\_mod.F90.

10.14.2.6 real(dp), dimension(:), allocatable prg\_system\_mod::system\_type::mass

Mass of every atom in the system. These can be automatically loaded by using the structures of the ptable mod:

```
system%mass(i) = mass(mystem%atomic_number(i))
```

Allocation:

```
mass (nats)
```

Definition at line 113 of file prg\_system\_mod.F90.

```
10.14.2.7 integer prg_system_mod::system_type::nats
```

Number of atoms of the system.

Definition at line 75 of file prg\_system\_mod.F90.

```
10.14.2.8 real(dp), dimension(:), allocatable prg_system_mod::system_type::net_charge
```

Charges of every atom in the system. Allocation:

```
net_charge(nats)
```

Definition at line 106 of file prg\_system\_mod.F90.

```
10.14.2.9 integer prg_system_mod::system_type::nsp
```

Number of different species. Number of species or number of different antom types (symbols) in the system. This integer is alwas less or equal than the total number of atoms ( $nsp \le nats$ ). This information can also be found in tbparams structure and the following equality holds:

```
system%nsp = tbparams%nsp
```

Definition at line 146 of file prg\_system\_mod.F90.

10.14.2.10 real(dp), dimension(:,:), allocatable prg\_system\_mod::system\_type::recip\_vector

Reciprocal vectors of the system. Allocation:

```
recip_vector(3,3)
v1 = recip_vector(1,:)
v2 = recip_vector(2,:)
v3 = recip_vector(3,:)
```

Definition at line 131 of file prg\_system\_mod.F90.

10.14.2.11 integer, dimension(:), allocatable prg\_system\_mod::system\_type::resindex Residue index. Definition at line 181 of file prg\_system\_mod.F90. 10.14.2.12 integer, dimension(:), allocatable prg\_system\_mod::system\_type::spatnum Species atomic number list. A list with the atomic numbers for every species Allocation: spatnum(nsp) Definition at line 169 of file prg system mod.F90. 10.14.2.13 integer, dimension(:), allocatable prg\_system\_mod::system\_type::spindex Species index. It gives the species index of a particulat atom. Allocation: spindex(nats) If we need the index of atom 30 then: system%spindex(30) Definition at line 154 of file prg\_system\_mod.F90. 10.14.2.14 character(2), dimension(:), allocatable prg\_system\_mod::system\_type::splist Species symbol list. A list with the different species e.g. H, C, N, etc with the order corresponding to the appearence in systemsymbol. Allocation: splist(nsp) Definition at line 162 of file prg\_system\_mod.F90. 10.14.2.15 real(dp), dimension(:), allocatable prg\_system\_mod::system\_type::spmass Species mass list. A list with the atomic mass for every species Allocation: spmass(nsp)

Definition at line 175 of file prg\_system\_mod.F90.

10.14.2.16 character(2), dimension(:), allocatable prg\_system\_mod::system\_type::symbol

Chemical Symbols for every atom of the system. Symbol can be recovered using ptable module and calling the following routine:

```
system%symbol(i) = element_symbol(system%atomic_number(i))
```

### Allocation:

```
symbol(nats)
```

Definition at line 83 of file prg system mod.F90.

10.14.2.17 real(dp), dimension(:), allocatable prg\_system\_mod::system\_type::userdef

User define field.

Definition at line 178 of file prg system mod.F90.

10.14.2.18 real(dp), dimension(:,:), allocatable prg\_system\_mod::system\_type::velocity

Velocities for every atom in the system. Allocation:

```
velocity(3, nats)
```

Definition at line 96 of file prg\_system\_mod.F90.

10.14.2.19 real(dp) prg\_system\_mod::system\_type::volk

Volume of the system (direct space).

Note

use prg\_get\_recip\_vects in coulomb\_latte\_mod to compute this.

Definition at line 139 of file prg\_system\_mod.F90.

10.14.2.20 real(dp) prg\_system\_mod::system\_type::volr

Volume of the system (direct space).

Note

use prg\_get\_recip\_vects in coulomb\_latte\_mod to compute this.

Definition at line 135 of file prg\_system\_mod.F90.

The documentation for this type was generated from the following file:

/home/christian/qmd-progress/src/prg\_system\_mod.F90

## 10.15 prg\_timer\_mod::timer\_status\_t Type Reference

Timer status type.

#### **Private Attributes**

• character(len=20) tname

Timer name.

integer tstart

Start time.

• integer ttotal

Current total time.

· integer tcount

Current call count.

• integer minrank

Rank with min value.

· integer maxrank

Rank with max value.

· real(dp) tsum

Sum time - total time in secs.

• real(dp) minvalue

Minimum value over all ranks.

• real(dp) maxvalue

Maximum value over all ranks.

real(dp) tavg

Average value over all ranks.

· real(dp) tstdev

Stdev across all ranks.

real(dp) tpercent

Percent of time across all timers.

### 10.15.1 Detailed Description

Timer status type.

Definition at line 54 of file prg\_timer\_mod.F90.

## 10.15.2 Member Data Documentation

10.15.2.1 integer prg\_timer\_mod::timer\_status\_t::maxrank [private]

Rank with max value.

Definition at line 72 of file prg\_timer\_mod.F90.

```
10.15.2.2 real(dp) prg_timer_mod::timer_status_t::maxvalue [private]
Maximum value over all ranks.
Definition at line 81 of file prg_timer_mod.F90.
10.15.2.3 integer prg_timer_mod::timer_status_t::minrank [private]
Rank with min value.
Definition at line 69 of file prg_timer_mod.F90.
10.15.2.4 real(dp) prg_timer_mod::timer_status_t::minvalue [private]
Minimum value over all ranks.
Definition at line 78 of file prg_timer_mod.F90.
10.15.2.5 real(dp) prg_timer_mod::timer_status_t::tavg [private]
Average value over all ranks.
Definition at line 84 of file prg_timer_mod.F90.
10.15.2.6 integer prg_timer_mod::timer_status_t::tcount [private]
Current call count.
Definition at line 66 of file prg_timer_mod.F90.
10.15.2.7 character(len=20) prg_timer_mod::timer_status_t::tname [private]
Timer name.
Definition at line 57 of file prg timer mod.F90.
10.15.2.8 real(dp) prg_timer_mod::timer_status_t::tpercent [private]
Percent of time across all timers.
Definition at line 90 of file prg_timer_mod.F90.
10.15.2.9 integer prg_timer_mod::timer_status_t::tstart [private]
Start time.
```

Definition at line 60 of file prg\_timer\_mod.F90.

10.15.2.10 real(dp) prg\_timer\_mod::timer\_status\_t::tstdev [private]

Stdev across all ranks.

Definition at line 87 of file prg\_timer\_mod.F90.

10.15.2.11 real(dp) prg\_timer\_mod::timer\_status\_t::tsum [private]

Sum time - total time in secs.

Definition at line 75 of file prg timer mod.F90.

10.15.2.12 integer prg\_timer\_mod::timer\_status\_t::ttotal [private]

Current total time.

Definition at line 63 of file prg\_timer\_mod.F90.

The documentation for this type was generated from the following file:

/home/christian/qmd-progress/src/prg\_timer\_mod.F90

## 10.16 prg\_extras\_mod::to\_string Interface Reference

### **Private Member Functions**

- character(len=:) function, allocatable to\_string\_integer (i)
   Convert integer to string.
- character(len=:) function, allocatable to\_string\_long\_long (i)
   Convert integer to string.
- character(len=:) function, allocatable to\_string\_double (x)
   Convert double to string.

### 10.16.1 Detailed Description

Definition at line 25 of file prg\_extras\_mod.F90.

### 10.16.2 Member Function/Subroutine Documentation

10.16.2.1 character(len=:) function, allocatable prg\_extras\_mod::to\_string::to\_string\_double ( double precision, intent(in) x ) [private]

Convert double to string.

#### **Parameters**

x The double

### Returns

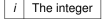
The string

Definition at line 80 of file prg\_extras\_mod.F90.

10.16.2.2 character(len=:) function, allocatable prg\_extras\_mod::to\_string::to\_string\_integer ( integer, intent(in) i ) [private]

Convert integer to string.

### **Parameters**



### Returns

The string

Definition at line 46 of file prg\_extras\_mod.F90.

10.16.2.3 character(len=:) function, allocatable prg\_extras\_mod::to\_string::to\_string\_long\_long ( integer(kind=c\_long\_long), intent(in) i ) [private]

Convert integer to string.

## **Parameters**

*i* The integer

### Returns

The string

Definition at line 62 of file prg\_extras\_mod.F90.

The documentation for this interface was generated from the following file:

• /home/christian/qmd-progress/src/prg\_extras\_mod.F90

# 10.17 prg\_xlbo\_mod::xlbo\_type Type Reference

General xlbo solver type.

# **Public Attributes**

- character(20) jobname
- · integer verbose
- · integer maxscfiter

Max SCF iterations at every XLBO MD step.

· integer maxscfinititer

Max SCF iterations for the first minit steps.

- real(dp) threshold
- · integer minit

Use SCF the first M\_prg\_init MD steps.

• real(dp) cc

Scaled prg\_delta Kernel.

# 10.17.1 Detailed Description

General xlbo solver type.

Definition at line 33 of file prg\_xlbo\_mod.F90.

## 10.17.2 Member Data Documentation

10.17.2.1 real(dp) prg\_xlbo\_mod::xlbo\_type::cc

Scaled prg\_delta Kernel.

Definition at line 51 of file prg\_xlbo\_mod.F90.

10.17.2.2 character(20) prg\_xlbo\_mod::xlbo\_type::jobname

Definition at line 35 of file prg\_xlbo\_mod.F90.

10.17.2.3 integer prg\_xlbo\_mod::xlbo\_type::maxscfinititer

Max SCF iterations for the first minit steps.

Definition at line 43 of file prg\_xlbo\_mod.F90.

10.17.2.4 integer prg\_xlbo\_mod::xlbo\_type::maxscfiter

Max SCF iterations at every XLBO MD step.

Definition at line 40 of file prg\_xlbo\_mod.F90.

10.17.2.5 integer prg\_xlbo\_mod::xlbo\_type::minit

Use SCF the first M\_prg\_init MD steps.

Definition at line 48 of file prg\_xlbo\_mod.F90.

10.17.2.6 real(dp) prg\_xlbo\_mod::xlbo\_type::threshold

Definition at line 45 of file prg\_xlbo\_mod.F90.

10.17.2.7 integer prg\_xlbo\_mod::xlbo\_type::verbose

Definition at line 37 of file prg\_xlbo\_mod.F90.

The documentation for this type was generated from the following file:

• /home/christian/qmd-progress/src/prg\_xlbo\_mod.F90

# 10.18 prg\_xlkernel\_mod::xlk\_type Type Reference

# **Public Attributes**

- character(20) kerneltype
  - Kernel type.
- integer verbose

Verbosity level.

- integer nrank
- real(dp) scalecoeff

Coefficient for mixing.

# 10.18.1 Detailed Description

Definition at line 17 of file prg\_xlkernel\_mod.F90.

# 10.18.2 Member Data Documentation

10.18.2.1 character(20) prg\_xlkernel\_mod::xlk\_type::kerneltype

Kernel type.

Definition at line 20 of file prg\_xlkernel\_mod.F90.

10.18.2.2 integer prg\_xlkernel\_mod::xlk\_type::nrank

Definition at line 23 of file prg\_xlkernel\_mod.F90.

10.18.2.3 real(dp) prg\_xlkernel\_mod::xlk\_type::scalecoeff

Coefficient for mixing.

Definition at line 26 of file prg\_xlkernel\_mod.F90.

10.18.2.4 integer prg\_xlkernel\_mod::xlk\_type::verbose

Verbosity level.

Definition at line 23 of file prg\_xlkernel\_mod.F90.

The documentation for this type was generated from the following file:

• /home/christian/qmd-progress/src/prg\_xlkernel\_mod.F90

# **Chapter 11**

# **File Documentation**

- 11.1 /home/christian/qmd-progress/README.md File Reference
- 11.2 /home/christian/qmd-progress/tests/README.md File Reference
- 11.3 /home/christian/qmd-progress/src/prg\_charges\_mod.F90 File Reference

## **Modules**

• module prg\_charges\_mod

A module to compute the Mulliken charges of a chemical system.

# **Functions/Subroutines**

• subroutine, public prg\_charges\_mod::prg\_get\_charges (rho\_bml, over\_bml, hindex, charges, numel, spindex, mdimin, threshold)

Constructs the charges from the density matrix.

• subroutine, public prg\_charges\_mod::prg\_get\_hscf (ham0\_bml, over\_bml, ham\_bml, spindex, hindex, hub-bardu, charges, coulomb\_pot\_r, coulomb\_pot\_k, mdimin, threshold)

Constructs the SCF Hamiltonian given H0, HubbardU and charges. This routine does:  $H = \sum_i U_i q_i + V_i$ ;, where U is the Hubbard parameter for every atom i. V is the coulombic potential for every atom i.

## **Variables**

- integer, parameter prg\_charges\_mod::dp = kind(1.0d0)
- 11.4 /home/christian/qmd-progress/src/prg\_chebyshev\_mod.F90 File Reference

# **Data Types**

type prg\_chebyshev\_mod::chebdata\_type
 General Cheb solver type.

## **Modules**

· module prg chebyshev mod

Module to obtain the density matrix by applying a Chebyshev polynomial expansion.

## **Functions/Subroutines**

• subroutine, public prg\_chebyshev\_mod::prg\_parse\_cheb (chebdata, filename)

Chebyshev parser. This module is used to parse all the input variables for the cheb electronic structure solver. Adding a new input keyword to the parser:

• subroutine, public prg\_chebyshev\_mod::prg\_build\_density\_cheb (ham\_bml, rho\_bml, athr, threshold, nco-effs, kbt, ef, bndfil, jon, verbose)

Builds the density matrix from  $H_0$  for a Fermi function approximated with a Chebyshev polynomial expansion.

subroutine, public prg\_chebyshev\_mod::prg\_build\_density\_cheb\_fermi (ham\_bml, rho\_bml, athr, threshold, ncoeffs, kbt, ef, bndfil, getef, fermitol, jon, npts, trkfunc, verbose)

Builds the density matrix from  $H_0$  for a Fermi function approximated with a Chebyshev polynomial expansion. In this case the self-consistent recursion is applied to converge to the correct number of electrons and obtain the Fermi level.

• real(dp) function prg\_chebyshev\_mod::jackson (ncoeffs, i, jon)

Evaluates the Jackson Kernel Coefficients.

• subroutine prg\_chebyshev\_mod::prg\_get\_chebcoeffs (npts, kbt, ef, ncoeffs, coeffs, emin, emax)

Gets the coefficients of the Chebyshev expansion.

• subroutine <a href="mailto:prg\_chebyshev\_mod::prg\_get\_chebcoeffs\_fermi\_bs">prg\_chebyshev\_mod::prg\_get\_chebcoeffs\_fermi\_bs</a> (npts, kbt, ef, tracesT, ncoeffs, coeffs, emin, emax, bndfil, norb, tol, jon, verbose)

Gets the coefficients of the Chebyshev expansion with Ef computation.

• subroutine <a href="mailto:prg\_chebyshev\_mod::prg\_get\_chebcoeffs\_fermi\_nt">prg\_chebyshev\_mod::prg\_get\_chebcoeffs\_fermi\_nt</a> (npts, kbt, ef, tracesT, ncoeffs, coeffs, emin, emax, bndfil, norb, tol, jon, verbose)

Gets the coefficients of the Chebyshev expansion with Ef computation.

real(dp) function prg\_chebyshev\_mod::tr (r, x)

Chebyshev polynomial obtained by recursion.

• real(dp) function prg\_chebyshev\_mod::fermi (e, ef, kbt)

Gives the Fermi distribution value for energy e.

real(dp) function prg\_chebyshev\_mod::absmaxderivative (func, de)

Gets the absolute maximum of the derivative of a function.

#### **Variables**

- integer, parameter prg\_chebyshev\_mod::dp = kind(1.0d0)
- real(dp), parameter prg\_chebyshev\_mod::pi = 3.14159265358979323846264338327950\_dp

# 11.5 /home/christian/qmd-progress/src/prg\_densitymatrix\_mod.F90 File Reference

## **Modules**

module prg\_densitymatrix\_mod

Module to obtain the density matrix by diagonalizing an orthogonalized Hamiltonian.

#### **Functions/Subroutines**

- subroutine, public prg\_densitymatrix\_mod::prg\_build\_density\_t0 (ham\_bml, rho\_bml, threshold, bndfil) Builds the density matrix from  $H_0$  for zero electronic temperature.  $\rho = C\Theta(\mu I - \epsilon)C^{\dagger}$  Where, C is the matrix eigenvector and  $\epsilon$  is the matrix eigenvalue.  $\Theta()$  is the Heaviside function.
- subroutine, public prg\_densitymatrix\_mod::prg\_build\_density\_t (ham\_bml, rho\_bml, threshold, bndfil, kbt, ef) Builds the density matrix from  $H_0$  for electronic temperature T.  $\rho = C f(\mu I - \epsilon) C^{\dagger}$  Where, C is the matrix eigenvector and  $\epsilon$  is the matrix eigenvalue. f is the Fermi function.
- subroutine, public prg\_densitymatrix\_mod::prg\_build\_density\_t\_fermi (ham\_bml, rho\_bml, threshold, kbt, ef, verbose)

Builds the density matrix from  $H_0$  for electronic temperature T.  $\rho = C f(\mu I - \epsilon) C^{\dagger}$  Where, C is the matrix eigenvector and  $\epsilon$  is the matrix eigenvalue. f is the Fermi function. In this routine the Fermi level is passed as an argument.

• subroutine, public prg\_densitymatrix\_mod::prg\_build\_atomic\_density (rhoat\_bml, numel, hindex, spindex, norb, bml\_type)

Builds the atomic density matrix.  $\rho_{ii} = mathcal Z_{ii}$  Where,  $mathcal Z_{ii}$  is the number of electrons for orbital i.

• subroutine, public prg\_densitymatrix\_mod::prg\_get\_flevel (eigenvalues, kbt, bndfil, tol, Ef)

Routine to compute the Fermi level given a set of eigenvalues and a temperature. It applies the Bisection method over the function:  $g(\mu) = \sum_k 2f(\epsilon_k - \mu) - N = 0$  Where  $f(\epsilon_k - \mu) = \frac{1}{1 + \exp{(\epsilon_k - \mu)/(k_b T)}}$ .

- subroutine, public prg\_densitymatrix\_mod::prg\_get\_eigenvalues (ham\_bml, eigenvalues, verbose)
  - Gets the eigenvalues of the Orthogonalized Hamiltonian.
- subroutine, public prg\_densitymatrix\_mod::prg\_check\_idempotency (mat\_bml, threshold, idempotency)
  - To check the idempotency error of a matrix. This is calculated as the Frobenius norm of  $(A A^2)$ .
- real(dp) function prg\_densitymatrix\_mod::fermi (e, ef, kbt)

Gives the Fermi distribution value for energy e.

#### **Variables**

• integer, parameter prg\_densitymatrix\_mod::dp = kind(1.0d0)

# 11.6 /home/christian/qmd-progress/src/prg\_dos\_mod.F90 File Reference

## **Modules**

· module prg dos mod

A module to compute the Density of state (DOS) and IDOS.

# **Functions/Subroutines**

- subroutine, public prg\_dos\_mod::prg\_write\_tdos (eigenvals, gamma, npts, emin, emax, filename) Writes the total DOS into a file.  $DOS(\epsilon) = \sum_k L(\epsilon \epsilon_k)$  Where  $\int_{-\infty}^{\infty} DOS(\epsilon) = Nstates$ .
- real(dp) function prg\_dos\_mod::lorentz (energy, eigenvals, loads, Gamma)

Lorentzian Function.

#### **Variables**

• integer, parameter prg\_dos\_mod::dp = kind(1.0d0)

# 11.7 /home/christian/qmd-progress/src/prg\_doxy\_mod.F90 File Reference

# 11.8 /home/christian/qmd-progress/src/prg\_extras\_mod.F90 File Reference

# **Data Types**

- interface prg\_extras\_mod::prg\_memory\_consumption
- · interface prg extras mod::to string

## **Modules**

 module prg\_extras\_mod Extra routines.

#### **Functions/Subroutines**

- character(len=:) function, allocatable prg\_extras\_mod::to\_string\_integer (i) Convert integer to string.
- character(len=:) function, allocatable prg\_extras\_mod::to\_string\_long\_long (i) Convert integer to string.
- character(len=:) function, allocatable prg\_extras\_mod::to\_string\_double (x)
   Convert double to string.
- subroutine, public prg\_extras\_mod::prg\_print\_matrix (matname, amat, i1, i2, j1, j2)

  To write a dense matrix to screen.
- real(dp) function, public prg\_extras\_mod::mls ()

To get the actual time in milliseconds.

• subroutine, public prg\_extras\_mod::prg\_delta (x, s, nn, dta)

Delta function  $||X^{\wedge}tSX - I||$ .

• subroutine, public prg\_extras\_mod::prg\_get\_mem (procname, tag)

Get proc memory.

subroutine prg\_extras\_mod::prg\_twonorm (a, nn, norm2)

Gets the norm2 of a square matrix.

# **Variables**

• integer, parameter prg\_extras\_mod::dp = kind(1.0d0)

# 11.9 /home/christian/qmd-progress/src/prg\_genz\_mod.F90 File Reference

# **Data Types**

type prg\_genz\_mod::genzspinp

Input for the genz driver. This type controlls all the variables that are needed by genz.

• type prg\_genz\_mod::genzspdata

contains the data for the genZ driver.

## **Modules**

module prg\_genz\_mod

To produce a matrix Z which is needed to orthogonalize H.

#### **Functions/Subroutines**

• subroutine, public prg\_genz\_mod::prg\_parse\_zsp (input, filename)

The parser for genz solver.

• subroutine prg\_genz\_mod::prg\_init (self, input)

Initializes the genz input variables.

subroutine prg\_genz\_mod::prg\_allocatezspmat (self, zk1\_bml, zk2\_bml, zk3\_bml, zk4\_bml, zk5\_bml, zk6←bml, norb, bml type)

Allocates the matrices for the XI integration of Z.

• subroutine, public prg\_genz\_mod::prg\_init\_zspmat (igenz, zk1\_bml, zk2\_bml, zk3\_bml, zk4\_bml, zk5\_bml, zk6\_bml, norb, bml\_type)

Initiates the matrices for the XI integration of Z.

subroutine prg\_genz\_mod::prg\_generate (self, over\_bml, zmat\_bml, igenz, mdim, bml\_type, zk1\_bml, zk2
 \_bml, zk3\_bml, zk4\_bml, zk5\_bml, zk6\_bml)

Generates the Z matrix.

subroutine, public prg\_genz\_mod::prg\_buildzdiag (smat\_bml, zmat\_bml, threshold, mdimin, bml\_type, verbose)

Usual subroutine involving diagonalization.  $Z=U\sqrt{s}U^\dagger$ , where U = eigenvectors and s = eigenvalues. The purpose of this subroutine is to have an exact way of computing z for comparing with the sparse approach.

- subroutine, public prg\_genz\_mod::prg\_buildzsparse (smat\_bml, zmat\_bml, igenz, mdim, bml\_type, zk1\_bml, zk2\_bml, zk3\_bml, zk4\_bml, zk5\_bml, zk6\_bml, nfirst, nrefi, nreff, thresholdi, thresholdi, integration, verbose)
   Inverse factorization using Niklasson's algorithm.
- subroutine, public prg\_genz\_mod::prg\_genz\_sp\_initialz0 (smat\_bml, zmat\_bml, norb, mdim, bml\_type\_
   f, threshold)

Initial estimation of Z.

subroutine, public prg\_genz\_mod::prg\_genz\_sp\_initial\_zmat (smat\_bml, zmat\_bml, norb, mdim, bml\_type\_f, threshold)

Initial estimation of Z.

• subroutine <a href="mailto:prg\_genz\_mod::prg\_genz\_sp\_int">prg\_genz\_sp\_int</a> (zmat\_bml, zk1\_bml, zk2\_bml, zk3\_bml, zk4\_bml, zk5\_bml, zk6\_bml, igenz, norb, bml\_type, threshold)

Inverse factorization using Niklasson's algorithm.

• subroutine, public prg\_genz\_mod::prg\_genz\_sp\_ref (smat\_bml, zmat\_bml, nref, norb, bml\_type, threshold)

\*\*Iterative refinement.\*\*

## **Variables**

• integer, parameter prg\_genz\_mod::dp = kind(1.0d0)

# 11.10 /home/christian/qmd-progress/src/prg\_graph\_mod.F90 File Reference

## **Data Types**

· type prg\_graph\_mod::subgraph\_t

Subgraph type.

type prg\_graph\_mod::graph\_partitioning\_t

Trace per iteration.

## **Modules**

· module prg\_graph\_mod

The graph module.

# **Functions/Subroutines**

• subroutine, public prg\_graph\_mod::prg\_initsubgraph (sg, pnum, hsize)

Initialize subgraph.

• subroutine, public prg\_graph\_mod::prg\_destroysubgraph (sg)

Destroy subgraph.

• subroutine, public prg\_graph\_mod::prg\_initgraphpartitioning (gp, pname, np, nnodes, nnodes2)

Initialize graph partitioning.

• subroutine, public prg\_graph\_mod::prg\_destroygraphpartitioning (gp)

Destroy graph partitioning.

• subroutine, public prg\_graph\_mod::prg\_printgraphpartitioning (gp)

Print graph partitioning structure data.

• subroutine, public prg\_graph\_mod::prg\_equalpartition (gp, nodesPerPart, nnodes)

Create equal graph partitions, based on number of rows/orbitals.

• subroutine, public prg\_graph\_mod::prg\_equalgrouppartition (gp, hindex, ngroup, nodesPerPart, nnodes)

Create equal group graph partitions, based on number of atoms/groups.

• subroutine, public prg\_graph\_mod::prg\_filepartition (gp, partFile)

Read graph partitions from a file, based on number of rows/orbitals.

subroutine prg\_graph\_mod::prg\_readpart (gp, partFile)

Read parts (core) from part file.

• subroutine, public prg\_graph\_mod::prg\_fnormgraph (gp)

Accumulate trace norm across all subgraphs.

## **Variables**

• integer, parameter prg\_graph\_mod::dp = kind(1.0d0)

# 11.11 /home/christian/qmd-progress/src/prg\_graphsp2parser\_mod.F90 File Reference

# **Data Types**

• type prg\_graphsp2parser\_mod::gsp2data\_type

General SP2 solver type.

# Modules

· module prg\_graphsp2parser\_mod

Graph partitioning SP2 parser.

# **Functions/Subroutines**

• subroutine, public prg\_graphsp2parser\_mod::prg\_parse\_gsp2 (gsp2data, filename)

The parser for SP2 solver.

## **Variables**

integer, parameter prg\_graphsp2parser\_mod::dp = kind(1.0d0)

# 11.12 /home/christian/qmd-progress/src/prg\_homolumo\_mod.F90 File Reference

#### **Modules**

· module prg\_homolumo\_mod

The homolumo module.

## **Functions/Subroutines**

- subroutine, public prg\_homolumo\_mod::prg\_homolumogap (vv, imax, pp, mineval, maxeval, ehomo, elumo, egap, verbose)
- subroutine, public prg\_homolumo\_mod::prg\_sp2sequence (pp, imax, mineval, maxeval, ehomo, elumo, errlimit, verbose)

#### **Variables**

integer, parameter prg\_homolumo\_mod::dp = kind(1.0d0)

# 11.13 /home/christian/qmd-progress/src/prg\_implicit\_fermi\_mod.F90 File Reference

#### Modules

module prg\_implicit\_fermi\_mod

# **Functions/Subroutines**

subroutine, public prg\_implicit\_fermi\_mod::prg\_implicit\_fermi (h\_bml, xi0\_bml, p\_bml, nsteps, nocc, mu, beta, osteps, occErrLimit, threshold)

Recursive Implicit Fermi Dirac.

## **Variables**

integer, parameter prg\_implicit\_fermi\_mod::dp = kind(1.0d0)

# 11.14 /home/christian/qmd-progress/src/prg\_initmatrices\_mod.F90 File Reference

# Modules

· module prg\_initmatrices\_mod

Initialization module.

## **Functions/Subroutines**

- subroutine, public prg\_initmatrices\_mod::prg\_init\_hsmat (ham\_bml, over\_bml, bml\_type, mdim, norb)

  Initialize Hamiltonian and Overlap Matrix.
- subroutine, public prg\_initmatrices\_mod::prg\_init\_pzmat (rho\_bml, zmat\_bml, bml\_type, mdim, norb)

  Initialize Density matrix and Inverse square root Overlap.
- subroutine, public prg\_initmatrices\_mod::prg\_init\_ortho (orthoh\_bml, orthop\_bml, bml\_type, mdim, norb)

  Initialize The orthogonal versions of Hamiltonian and Density Matrix.

#### **Variables**

• integer, parameter prg\_initmatrices\_mod::dp = kind(1.0d0)

# 11.15 /home/christian/qmd-progress/src/prg\_kernelparser\_mod.F90 File Reference

#### **Modules**

module prg\_kernelparser\_mod
 Some general parsing functions.

## **Functions/Subroutines**

- subroutine, public prg\_kernelparser\_mod::prg\_parsing\_kernel (keyvector\_char, valvector\_char, keyvector\_
  int, valvector\_int, keyvector\_re, valvector\_re, keyvector\_log, valvector\_log, filename, startstop)
  - The general parsing function. It is used to vectorize a set of "keywords" "value" pairs as included in a general input file.
- subroutine prg\_kernelparser\_mod::prg\_check\_valid (invalidc)

Check for valid keywords (checks for an = sign)

# **Variables**

• integer, parameter prg\_kernelparser\_mod::dp = kind(1.0d0)

# 11.16 /home/christian/qmd-progress/src/prg\_nonortho\_mod.F90 File Reference

## **Modules**

• module prg\_nonortho\_mod

Module to prg\_orthogonalize and prg\_deorthogonalize any operator.

# **Functions/Subroutines**

This routine performs:  $A_{ortho} = Z^{\dagger}AZ$ .

subroutine, public prg\_nonortho\_mod::prg\_deorthogonalize (orthoA\_bml, zmat\_bml, a\_bml, threshold, bml
 \_type, verbose)

This routine performs:  $A = ZA_{ortho}Z^{\dagger}$ .

## **Variables**

• integer, parameter prg\_nonortho\_mod::dp = kind(1.0d0)

# 11.17 /home/christian/gmd-progress/src/prg normalize mod.F90 File Reference

## **Modules**

· module prg\_normalize\_mod

The prg\_normalize module.

# **Functions/Subroutines**

• subroutine, public prg\_normalize\_mod::prg\_normalize (h\_bml)

Normalize a Hamiltonian matrix prior to running the SP2 algorithm.

• subroutine, public prg\_normalize\_mod::prg\_normalize\_fermi (h\_bml, h1, hN, mu)

Normalize a Hamiltonian matrix prior to running the truncated SP2 algorithm.

• subroutine, public prg\_normalize\_mod::prg\_normalize\_implicit\_fermi (h\_bml, cnst, mu)

Normalize a Hamiltonian matrix prior to running the implicit fermi dirac algorithm.

subroutine, public prg\_normalize\_mod::prg\_gershgorinreduction (gp)

Determine gershgorin bounds across all parts, local and distributed.

subroutine, public prg\_normalize\_mod::prg\_normalize\_cheb (h\_bml, mu, emin, emax, alpha, scaledmu)

Normalize a Hamiltonian matrix prior to running the Chebyshev algorithm.

#### **Variables**

• integer, parameter prg\_normalize mod::dp = kind(1.0d0)

# 11.18 /home/christian/qmd-progress/src/prg\_openfiles\_mod.F90 File Reference

# **Modules**

· module prg\_openfiles\_mod

Module to handle input output files for the PROGRESS lib.

# **Functions/Subroutines**

• integer function, public prg\_openfiles mod::get\_file\_unit (io\_max)

Returns a unit number that is not in use.

• subroutine, public prg\_openfiles\_mod::prg\_open\_file (io, name)

Opens a file to write.

• subroutine, public prg\_openfiles\_mod::prg\_open\_file\_to\_read (io, name)

Opens a file to read.

# 11.19 /home/christian/qmd-progress/src/prg\_parallel\_mod.F90 File Reference

## **Data Types**

type prg\_parallel mod::rankreducedata\_t

Data structure for rection over MPI ranks.

#### **Modules**

· module prg\_parallel\_mod

The parallel module.

#### **Functions/Subroutines**

- integer function, public prg\_parallel\_mod::getnranks ()
- integer function, public prg\_parallel\_mod::getmyrank ()
- integer function, public prg\_parallel mod::printrank ()
- subroutine, public prg\_parallel mod::prg\_initparallel()
- subroutine, public prg\_parallel\_mod::prg\_shutdownparallel ()
- integer function prg\_parallel mod::saverequest (irequest)
- subroutine, public prg\_parallel\_mod::prg\_barrierparallel ()
- subroutine, public prg\_parallel\_mod::sendreceiveparallel (sendBuf, sendLen, dest, recvBuf, recvLen, source, nreceived)
- subroutine, public prg\_parallel mod::isendparallel (sendBuf, sendLen, dest)
- subroutine, public prg\_parallel\_mod::sendparallel (sendBuf, sendLen, dest)
- subroutine, public prg\_parallel\_mod::prg\_iprg\_recvparallel (recvBuf, recvLen, rind)
- subroutine, public prg\_parallel\_mod::prg\_recvparallel (recvBuf, recvLen)
- subroutine, public prg\_parallel mod::sumintparallel (sendBuf, recvBuf, icount)
- subroutine, public prg\_parallel\_mod::sumrealparallel (sendBuf, recvBuf, icount)
- subroutine, public prg\_parallel\_mod::maxintparallel (sendBuf, recvBuf, icount)
- subroutine, public prg\_parallel\_mod::maxrealparallel (sendBuf, recvBuf, icount)
- subroutine, public prg\_parallel\_mod::minintparallel (sendBuf, recvBuf, icount)
- subroutine, public prg\_parallel\_mod::minrealparallel (sendBuf, recvBuf, icount)
- subroutine, public prg\_parallel mod::prg\_minrealreduce (rvalue)
- subroutine, public prg\_parallel\_mod::prg\_maxrealreduce (rvalue)
- subroutine, public prg parallel mod::prg maxintreduce2 (value1, value2)
- subroutine, public prg\_parallel\_mod::prg\_sumintreduce2 (value1, value2)
- subroutine, public prg\_parallel\_mod::prg\_sumrealreduce (value1)
- subroutine, public prg\_parallel\_mod::prg\_sumrealreduce2 (value1, value2)
- subroutine, public prg\_parallel\_mod::prg\_sumrealreduce3 (value1, value2, value3)
- subroutine, public prg\_parallel\_mod::prg\_sumrealreducen (valueVec, N)
- subroutine, public prg\_parallel\_mod::prg\_sumintreducen (valueVec, N)
- subroutine, public prg\_parallel mod::minrankrealparallel (sendBuf, recvBuf, icount)
- subroutine, public prg\_parallel\_mod::maxrankrealparallel (sendBuf, recvBuf, icount)
- subroutine, public prg\_parallel\_mod::prg\_bcastparallel (buf, blen, root)
- subroutine, public prg\_parallel\_mod::allgatherrealparallel (sendBuf, sendLen, recvBuf, recvLen)
- subroutine, public prg\_parallel\_mod::allgatherintparallel (sendBuf, sendLen, recvBuf, recvLen)
- subroutine, public prg\_parallel\_mod::allgathervrealparallel (sendBuf, sendLen, recvBuf, recvLen, recvDispl)
- subroutine, public prg\_parallel\_mod::allgathervintparallel (sendBuf, sendLen, recvBuf, recvLen, recvDispl)
- subroutine, public prg\_parallel\_mod::prg\_allsumrealreduceparallel (buf, buflen)
- subroutine, public prg\_parallel\_mod::prg\_allsumintreduceparallel (buf, buflen)
- subroutine, public prg parallel mod::prg allgatherparallel (a)
- subroutine, public prg\_parallel\_mod::prg\_wait ()

## **Variables**

- integer, parameter prg\_parallel mod::dp = kind(1.0d0)
- integer prg\_parallel\_mod::myrank
- integer prg\_parallel mod::nranks
- integer prg parallel mod::ierr
- integer prg\_parallel\_mod::reqcount
- integer, dimension(:), allocatable prg\_parallel mod::requestlist
- integer, dimension(:), allocatable prg\_parallel\_mod::rused

# 11.20 /home/christian/qmd-progress/src/prg\_partition\_mod.F90 File Reference

#### **Modules**

· module prg\_partition\_mod

The partition module.

## **Functions/Subroutines**

• subroutine, public prg\_partition\_mod::prg\_metispartition (gp, ngroups, nnodes, xadj, adjncy, nparts, part, core\_count, CH\_count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm)

Create graph partitions minizing number of cut edges.

• subroutine, public prg\_partition\_mod::prg\_costpartition (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm)

Compute cost of a partition.

• subroutine, public prg\_partition\_mod::update\_prg\_costpartition (gp, xadj, adjncy, partNumber, core\_count, CH count, Halo count, sumCubes, maxCH, smooth maxCH, pnorm, node, new part)

Update cost of partition and the different parameters node is moves into new\_part For each neighbor of node, the following cases hold: Case 1: neighbor is in old\_part Case 2: neighbor is in new\_part Case 3: neighbor is neither in old\_or new\_part.

• subroutine prg partition mod::prg accept prob (it, prg delta, r)

Compute acceptance probability for simulated annealing.

• subroutine prg\_partition\_mod::prg\_costindex (cost, sumCubes, maxCH, smooth\_maxCH, obj\_fun)

Choose objective function to work with.

• subroutine prg\_partition\_mod::prg\_rand\_node (gp, node, seed)

Pick a random node.

• subroutine, public prg\_partition\_mod::prg\_simannealing (gp, xadj, adjncy, partNumber, core\_count, CH\_← count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm, niter, seed)

Graph partitioning based on Simulated Annealing.

• subroutine, public prg\_partition\_mod::prg\_kernlin (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm, nconverg, seed)

Graph partitioning based on inspired by Kernighan-Lin Review METiS manual for description of k-way implementation of KL Pick a core together with its halos Place free vertices on a priority queue with (key, value) =(prg\_delta, best\_← part), with prg\_delta = change in obj\_value Dequeue and allow hill climbing.

- subroutine, public prg partition mod::prg update gp (gp, partNumber, core count)
- subroutine prg\_partition\_mod::prg\_rand\_shuffle (array, seed)

Randomly shuffle array.

• subroutine, public prg partition mod::prg check arrays (gp, core count, CH count, Halo count)

Error checking Checking that core\_count, CH\_count, Halo\_count match.

subroutine, public prg\_partition\_mod::prg\_kernlin\_queue (gp, xadj, adjncy, partNumber, core\_count, CH\_←
count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm)

Greedy algorithm. At each step it chooses the (vertex, new\_part) pair with highest gain Currently implementation is very slow.

• subroutine prg\_partition\_mod::prg\_find\_best\_move (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm, best\_node, best\_part)

For kerlin\_queue to find (vertex, new\_part) pair with highest gain.

- subroutine, public prg\_partition\_mod::prg\_kernlin2 (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo count, sumCubes, maxCH, smooth maxCH, pnorm)
- subroutine <a href="mailto:prg\_get\_largest\_hedge\_in\_part">part</a> (gp, xadj, adjncy, partNumber, core\_count, CH\_count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm, search\_part, largest\_Hedge)
- subroutine, public prg\_partition\_mod::prg\_simannealing\_old (gp, xadj, adjncy, partNumber, core\_count, C←
   H\_count, Halo\_count, sumCubes, maxCH, smooth\_maxCH, pnorm, niter, seed)

# **Variables**

- integer, parameter prg\_partition\_mod::dp = kind(1.0d0)
- integer, parameter prg\_partition\_mod::metis\_index\_kind = METIS\_INDEX\_KIND From /usr/include/metis.h.
- integer, parameter prg\_partition\_mod::metis\_real\_kind = kind(METIS\_REAL\_KIND)

  From /usr/include/metis.h.

# 11.21 /home/christian/qmd-progress/src/prg progress mod.F90 File Reference

## Modules

• module prg\_progress\_mod

The progress module.

## **Functions/Subroutines**

- subroutine, public prg\_progress\_mod::prg\_progress\_init ()
   Initialize progress.
- subroutine, public prg\_progress\_mod::prg\_progress\_shutdown () Shutdown progress.

## **Variables**

• integer, parameter prg\_progress mod::dp = kind(1.0d0)

# 11.22 /home/christian/qmd-progress/src/prg\_ptable\_mod.F90 File Reference

# **Modules**

• module prg\_ptable\_mod

Periodic table of elements.

## **Functions/Subroutines**

- integer function, public prg\_ptable\_mod::element\_atomic\_number (symbol)
- integer function prg\_ptable\_mod::element\_atomic\_number\_upper (symbol)

## **Variables**

- integer, parameter prg\_ptable\_mod::nz = 103
- integer, parameter, private prg\_ptable\_mod::dp = kind(1.0d0)
- character(2), dimension(nz), parameter prg\_ptable\_mod::element\_symbol = [character(2) :: "H", "He", "Li", "Be", "B", "C", "N", "O", "F", "Ne", "Na", "Mg", "Al", "Si", "P", "S", "Cl", "Ar", "K", "Ca", "Sc", "Ti", "V", "Cr", "Mn", "Fe", "Co", "Ni", "Cu", "Zn", "Ga", "Ge", "As", "Se", "Br", "Kr", "Rb", "Sr", "Y", "Zr", "Nb", "Mo", "Tc", "Ru", "Rh", "Pd", "Ag", "Cd", "In", "Sn", "Sb", "Te", "I", "Xe", "Cs", "Ba", "La", "Ce", "Pr", "Nd", "Pm", "Sm", "Eu", "Gd", "Tb", "Dy", "Ho", "Er", "Tm", "Yb", "Lu", "Hf", "Ta", "W", "Re", "Os", "Ir", "Pt", "Au", "Hg", "TI", "Pb", "Bi", "Po", "At", "Rn", "Fr", "Ra", "Ac", "Th", "Pa", "U", "Np", "Pu", "Am", "Cm", "Bk", "Cf", "Es", "Fm", "Md", "No", "Lr"]

#### Element symbol.

character(2), dimension(nz), parameter prg\_ptable\_mod::element\_symbol\_upper = [character(2) :: "H", "HE", "LI", "BE", "B", "C", "N", "O", "F", "NE", "NA", "MG", "AL", "SI", "P", "S", "CL", "AR", "K", "CA", "SC", "TI", "V", "CR", "MN", "FE", "CO", "NI", "CU", "ZN", "GA", "GE", "AS", "SE", "BR", "KR", "RB", "SR", "Y", "ZR", "NB", "MO", "TC", "RU", "RH", "PD", "AG", "CD", "IN", "SN", "SB", "TE", "II", "XE", "CS", "BA", "LA", "CE", "PR", "ND", "PM", "SM", "EU", "GD", "TB", "DY", "HO", "ER", "TM", "YB", "LU", "HF", "TA", "W", "RE", "OS", "IR", "PT", "AU", "HG", "TL", "PB", "BI", "PO", "AT", "RN", "FR", "RA", "AC", "TH", "PA", "U", "NP", "PU", "AM", "CM", "BK", "CF", "ES", "FM", "MD", "NO", "LR"]

Element symbol upper.

character(20), dimension(nz), parameter prg\_ptable\_mod::element\_name = [character(20) :: "Hydrogen", "Helium", "Lithium", "Beryllium", "Boron", "Carbon", "Nitrogen", "Oxygen", "Fluorine", "Neon", "Sodium", "Magnesium", "Aluminium", "Silicon", "Phosphorus", "Sulfur", "Chlorine", "Argon", "Potassium", "Calcium", "Scandium", "Titanium", "Vanadium", "Chromium", "Manganese", "Iron", "Cobalt", "Nickel", "Copper", "Zinc", "Gallium", "Germanium", "Arsenic", "Selenium", "Bromine", "Krypton", "Rubidium", "Strontium", "Yttrium", "Zirconium", "Niobium", "Molybdenum", "Technetium", "Ruthenium", "Rhodium", "Palladium", "Silver", "Cadmium", "Indium", "Tin", "Antimony", "Tellurium", "Iodine", "Xenon", "Caesium", "Barium", "Lanthanum", "Cerium", "Praseodymium", "Neodymium", "Promethium", "Samarium", "Europium", "Gadolinium", "Terbium", "Dysprosium", "Holmium", "Erbium", "Thulium", "Ytterbium", "Lutetium", "Hafnium", "Tantalum", "Tungsten", "Rhenium", "Osmium", "Iridium", "Platinum", "Gold", "Mercury", "Thallium", "Lead", "Bismuth", "Polonium", "Astatine", "Radon", "Francium", "Radium", "Actinium", "Thorium", "Protactinium", "Uranium", "Neptunium", "Plutonium", "Americium", "Curium", "Berkelium", "Californium", "Einsteinium", "Fermium", "Mendelevium", "Nobelium", "Lawrencium"]

# Element name.

• real(dp), dimension(nz), parameter prg\_ptable\_mod::element\_mass = (/ 1.007825032 , 4.002603254 , 7.  $\leftarrow$  01600455 , 9.0121822 , 11.0093054 , 12.0 , 14.003074005 , 15.99491462 , 18.99840322 , 19.992440175 , 22.989769281 , 23.9850417 , 26.98153863 , 27.976926532 , 30.97376163 , 31.972071 , 34.96885268 , 39.962383123 , 38.96370668 , 39.96259098 , 44.9559119 , 47.9479463 , 50.9439595 , 51.9405075 , 54.  $\leftarrow$  9380451 , 55.9349375 , 58.933195 , 57.9353429 , 62.9295975 , 63.929142 , 68.925573 , 73.921177 , 74.  $\leftarrow$  921596 , 79.916521 , 78.918337 , 83.911507 , 84.911789 , 87.905612 , 88.905848 , 89.904704 , 92.906378 , 97.905408 , 97.907216 , 101.904349 , 102.905504 , 105.903486 , 106.905097 , 113.903358 , 114.903878 , 119.902194 , 120.903815 , 129.906224 , 126.904473 , 131.904153 , 132.905451 , 137.905247 , 138.  $\leftarrow$  906353 , 139.905438 , 140.907652 , 141.907723 , 144.912749 , 151.919732 , 152.92123 , 157.924103 , 158.925346 , 163.929174 , 164.930322 , 165.930293 , 168.934213 , 173.938862 , 174.940771 , 179.94655 , 180.947995 , 183.950931 , 186.955753 , 191.96148 , 192.962926 , 194.964791 , 196.966568 , 201.970643 , 204.974427 , 207.976652 , 208.980398 , 208.98243 , 209.987148 , 222.017577 , 223.019735 , 226.025409 , 227.027752 , 232.038055 , 231.035884 , 238.050788 , 237.048173 , 244.064204 , 243.061381 , 247.070354 , 247.070307 , 251.079587 , 252.08298 , 257.095105 , 258.098431 , 259.10103 , 262.10963 /)

Element mass in atomic mass units (1.66 x 10-27 kg)

• real(dp), dimension(nz), parameter prg\_ptable\_mod::element\_vdwr = (/ 1.1 , 1.4 , 1.81 , 1.53 , 1.92 , 1.7 , 1.55 , 1.52 , 1.47 , 1.54 , 2.27 , 1.73 , 1.84 , 2.1 , 1.8 , 1.8 , 1.75 , 1.88 , 2.75 , 2.31 , 2.3 , 2.15 , 2.05 , 2.05 , 2.05 , 2.05 , 2.0 , 2.0 , 2.0 , 2.1 , 1.87 , 2.11 , 1.85 , 1.9 , 1.83 , 2.02 , 3.03 , 2.49 , 2.4 , 2.3 , 2.15 , 2.1 , 2.05 , 2.05 , 2.0 , 2.05 , 2.1 , 2.2 , 2.2 , 1.93 , 2.17 , 2.06 , 1.98 , 2.16 , 3.43 , 2.68 , 2.5 , 2.48 , 2.47 , 2.45 , 2.43 , 2.42 , 2.4 , 2.38 , 2.37 , 2.35 , 2.33 , 2.32 , 2.3 , 2.28 , 2.27 , 2.25 , 2.2 , 2.1 , 2.05 , 2.0 , 2

van der Waals radius (in Angstroms)

• real(dp), dimension(nz), parameter prg\_ptable\_mod::element\_covr = (/ 0.31 , 0.28 , 1.28 , 0.96 , 0.84 , 0.76 , 0.71 , 0.66 , 0.57 , 0.58 , 1.66 , 1.41 , 1.21 , 1.11 , 1.07 , 1.05 , 1.02 , 1.06 , 2.03 , 1.76 , 1.7 , 1.6 , 1.53 , 1.39 , 1.39 , 1.32 , 1.26 , 1.24 , 1.32 , 1.22 , 1.22 , 1.2 , 1.19 , 1.2 , 1.2 , 1.16 , 2.2 , 1.95 , 1.9 , 1.75 , 1.64 , 1.54 , 1.47 , 1.46 , 1.42 , 1.39 , 1.45 , 1.44 , 1.42 , 1.39 , 1.39 , 1.38 , 1.39 , 1.4 , 2.44 , 2.15 , 2.07 , 2.04 , 2.03 , 2.01 , 1.99 , 1.98 , 1.98 , 1.96 , 1.94 , 1.92 , 1.92 , 1.89 , 1.9 , 1.87 , 1.87 , 1.75 , 1.7 , 1.62 , 1.51 , 1.44 , 1.41 , 1.36 , 1.36 , 1.32 , 1.45 , 1.46 , 1.48 , 1.4 , 1.5 , 1.5 , 2.6 , 2.21 , 2.15 , 2.06 , 2.0 , 1.96 , 1.9 , 1.87 , 1.8 , 1.69 , 1.6 ,

Covalent radius (in Angstroms)

• real(dp), dimension(nz), parameter prg\_ptable\_mod::element\_ip = (/ 13.5984 , 24.5874 , 5.3917 , 9.3227 , 8.298 , 11.2603 , 14.5341 , 13.6181 , 17.4228 , 21.5645 , 5.1391 , 7.6462 , 5.9858 , 8.1517 , 10.4867 , 10.36 , 12.9676 , 15.7596 , 4.3407 , 6.1132 , 6.5615 , 6.8281 , 6.7462 , 6.7665 , 7.434 , 7.9024 , 7.881 , 7.6398 , 7.7264 , 9.3942 , 5.9993 , 7.8994 , 9.7886 , 9.7524 , 11.8138 , 13.9996 , 4.1771 , 5.6949 , 6.2173 , 6.6339 , 6.7589 , 7.0924 , 7.28 , 7.3605 , 7.4589 , 8.3369 , 7.5762 , 8.9938 , 5.7864 , 7.3439 , 8.6084 , 9.0096 , 10.4513 , 12.1298 , 3.8939 , 5.2117 , 5.5769 , 5.5387 , 5.473 , 5.525 , 5.582 , 5.6437 , 5.6704 , 6.1498 , 5.8638 , 5.9389 , 6.0215 , 6.1077 , 6.1843 , 6.2542 , 5.4259 , 6.8251 , 7.5496 , 7.864 , 7.8335 , 8.4382 , 8.967 , 8.9588 , 9.2255 , 10.4375 , 6.1082 , 7.4167 , 7.2855 , 8.414 , 0.0 , 10.7485 , 4.0727 , 5.2784 , 5.17 , 6.3067 , 5.89 , 6.1941 , 6.2657 , 6.026 , 5.9738 , 5.9914 , 6.1979 , 6.2817 , 6.42 , 6.5 , 6.58 , 6.65 , 4.9 /)

Ionization energy (in eV)

• real(dp), dimension(nz), parameter prg\_ptable\_mod::element\_ea = (/ 0.75420375 , 0.0 , 0.618049 , 0.0 , 0.279723 , 1.262118 , -0.07 , 1.461112 , 3.4011887 , 0.0 , 0.547926 , 0.0 , 0.43283 , 1.389521 , 0.7465 , 2.0771029 , 3.612724 , 0.0 , 0.501459 , 0.02455 , 0.188 , 0.084 , 0.525 , 0.67584 , 0.0 , 0.151 , 0.6633 , 1.15716 , 1.23578 , 0.0 , 0.41 , 1.232712 , 0.814 , 2.02067 , 3.363588 , 0.0 , 0.485916 , 0.05206 , 0.307 , 0.426 , 0.893 , 0.7472 , 0.55 , 1.04638 , 1.14289 , 0.56214 , 1.30447 , 0.0 , 0.404 , 1.112066 , 1.047401 , 1.970875 , 3.059038 , 0.0 , 0.471626 , 0.14462 , 0.47 , 0.5 , 0

Electron affprg\_inity (in eV)

• real(dp), dimension(nz), parameter prg\_ptable\_mod::atom\_en = (/ 2.2 , 0.0 , 0.98 , 1.57 , 2.04 , 2.55 , 3.04 , 3.44 , 3.98 , 0.0 , 0.93 , 1.31 , 1.61 , 1.9 , 2.19 , 2.58 , 3.16 , 0.0 , 0.82 , 1.0 , 1.36 , 1.54 , 1.63 , 1.66 , 1.55 , 1.83 , 1.88 , 1.91 , 1.9 , 1.65 , 1.81 , 2.01 , 2.18 , 2.55 , 2.96 , 3.0 , 0.82 , 0.95 , 1.22 , 1.33 , 1.6 , 2.16 , 1.9 , 2.2 , 2.28 , 2.2 , 1.93 , 1.69 , 1.78 , 1.96 , 2.05 , 2.1 , 2.66 , 2.6 , 0.79 , 0.89 , 1.1 , 1.12 , 1.13 , 1.14 , 0.0 , 1.17 , 0.0 , 1.2 , 0.0 , 1.22 , 1.23 , 1.24 , 1.25 , 0.0 , 1.27 , 1.3 , 1.5 , 2.36 , 1.9 , 2.2 , 2.2 , 2.28 , 2.54 , 2.0 , 1.62 , 2.33 , 2.02 , 2.0 , 2.2 , 0.0 , 0.7 , 0.9 , 1.1 , 1.3 , 1.5 , 1.38 , 1.36 , 1.28 , 1.3 , 1.3 , 1.3 , 1.3 , 1.3 , 1.3 , 1.3 , 1.3 , 1.3 , 1.3 , 0.0 /)

The Pauling electronegativity for this element.

The maximum expected number of bonds to this element.

• integer, dimension(nz), parameter prg\_ptable\_mod::element\_numel = (/ 1, 2, 1, 2, 3, 4, 5, 6, 7, 8, 1, 2, 3, 4, 5, 6, 7, 8, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17/)

Last shell number of electrons.

character(50), dimension(nz), parameter prg\_ptable\_mod::element\_econf = [character(50) :: "1s" , "1s2" , "1s22s" , "1s22s2" , "1s22s22p" , "1s22s22p2" , "1s22s22p3" , "1s22s22p4" , "1s22s22p5" , "1s22s22p6" , "[Ne]3s" , "[Ne]3s23p" , "[Ne]3s23p7" , "[Ne]3s23p5" , "[\times Ne]3s23p6" , "[Ar]4s2" , "[Ar]3d4s2" , "[Ar]3d24s2" , "[Ar]3d34s2" , "[Ar]3d54s2" , "[Ar]3d54s2" , "[Ar]3d54s2" , "[Ar]3d54s2" , "[Ar]3d54s2" , "[Ar]3d104s24p7" , "[Ar]3d104s24p7" , "[Ar]3d104s24p7" , "[Ar]3d104s24p6" , "[Ar]3d104s24p6" , "[Ar]3d104s24p6" , "[Kr]4d5s2" , "[Xe]4f16s2" , "[Xe]4f16s2" , "[Xe]4f16s2" , "[Xe]4f16s2" , "[Xe]4f16s2" , "[Xe]4f16s2" , "[Xe]4f145d6s2" , "[Xe]4f145d106s2" , "[Xe]4f145d1

The electronic configuration.

# 11.23 /home/christian/qmd-progress/src/prg\_pulaycomponent\_mod.F90 File Reference

#### **Modules**

· module prg\_pulaycomponent\_mod

Produces a matrix to get the Pulay Component of the forces.

## **Functions/Subroutines**

subroutine, public prg\_pulaycomponent\_mod::prg\_pulaycomponent0 (rho\_bml, ham\_bml, pcm\_bml, threshold, M, bml type, verbose)

At 
$$T = 0K$$
,  $P = \rho H \rho$ .

• subroutine, public prg\_pulaycomponent\_mod::prg\_pulaycomponentt (rho\_bml, ham\_bml, zmat\_bml, pcm\_← bml, threshold, M, bml\_type, verbose)

At 
$$T > 0K$$
,  $P = \rho H S^{-1} + S^{-1} H \rho$ .

subroutine, public prg\_pulaycomponent\_mod::prg\_get\_pulayforce (nats, zmat\_bml, ham\_bml, rho\_bml, d
 Sx\_bml, dSy\_bml, dSz\_bml, hindex, FPUL, threshold)

Pulay Force FPUL from  $2Tr[ZZ'HD\frac{dS}{dB}]$ .

## **Variables**

• integer, parameter prg\_pulaycomponent\_mod::dp = kind(1.0d0)

# 11.24 /home/christian/qmd-progress/src/prg\_pulaymixer\_mod.F90 File Reference

## **Data Types**

type prg\_pulaymixer\_mod::mx\_type

## **Modules**

· module prg\_pulaymixer\_mod

Pulay mixer mode.

## **Functions/Subroutines**

• subroutine, public prg\_pulaymixer\_mod::prg\_parse\_mixer (input, filename)

The parser for the mixer routines.

subroutine, public prg\_pulaymixer\_mod::prg\_qmixer (charges, oldcharges, dqin, dqout, scferror, piter, pulay-coef, mpulay, verbose)

Mixing the charges to acelerate scf convergence.

• subroutine, public prg\_pulaymixer\_mod::prg\_linearmixer (charges, oldcharges, scferror, linmixcoef, verbose)

Routine to perform linear mixing.

## **Variables**

integer, parameter prg\_pulaymixer\_mod::dp = kind(1.0d0)

# 11.25 /home/christian/qmd-progress/src/prg\_quantumdynamics\_mod.F90 File Reference

## **Modules**

module prg\_quantumdynamics\_mod

A module to add in common quantum dynamical operations.

## **Functions/Subroutines**

• subroutine, public prg\_quantumdynamics\_mod::prg\_kick\_density (kick\_direc, kick\_mag, dens, norbs, mdim, S, SINV, which atom, r, bmltype, thresh)

Provides perturbation to initial density matrix in the form of an electric field kick. This routine does:  $\rho_{\hat{k}\hat{i}ck} = \exp{\frac{-i}{\hbar}\hat{V}\hat{\rho}\hat{S}}\exp{\frac{i}{\hbar}\hat{V}\hat{S}^{-1}}$  where  $\hat{V}$  is the field disturbance.

• subroutine, public prg\_quantumdynamics\_mod::prg\_get\_sparsity\_cplxmat (matrix\_type, element\_type, thresh, a dense)

This computes the sparsity of a complex matrix given a threshold value This routine does:  $f = \frac{N_0}{N_{tot}}$  where f is the sparsity,  $N_0$  is the number of values less than the threshold, and  $N_{tot}$  is the total number of values. The sparsity and threshold are printed to the screen.

• subroutine, public prg\_quantumdynamics\_mod::prg\_get\_sparsity\_realmat (matrix\_type, element\_type, thresh, a\_dense)

This computes the sparsity of a real matrix given a threshold value This routine does:  $f = \frac{N_0}{N_{tot}}$  where f is the sparsity,  $N_0$  is the number of values less than the threshold, and  $N_{tot}$  is the total number of values. The sparsity and threshold are printed to the screen.

subroutine, public prg\_quantumdynamics\_mod::prg\_kick\_density\_bml (kick\_direc, kick\_mag, rho\_bml, s\_← bml, sinv bml, mdim, which atom, r, matrix type, thresh)

Provides perturbation to initial density matrix in the form of an electric field kick given input matricies in BML format. This routine does:  $\rho_{\hat{k}\hat{i}ck} = \exp{\frac{-i}{\hbar}\hat{V}\hat{\rho}\hat{S}} \exp{\frac{i}{\hbar}\hat{V}\hat{S}^{-1}}$  where  $\hat{V}$  is the field disturbance.

subroutine, public prg\_quantumdynamics\_mod::prg\_lvni\_bml (h1\_bml, sinv\_bml, dt, hbar, rhoold\_bml, rho
 \_bml, aux\_bml, matrix\_type, mdim, thresh)

Performs Liouville-von Neumann integration using leap-frog method. This routine does:  $\hat{\rho}(t+\Delta t)=\hat{\rho}(t-\Delta t)+2\Delta t \frac{\partial \hat{\rho}(t)}{\partial t}$  where the time derivative of the density matrix is defined as follows:  $\frac{\partial \hat{\rho}(t)}{\partial t}=\frac{i}{\hbar}\left(S^{-1}\hat{H}(t)\hat{\rho}(t)-\hat{\rho}(t)\hat{H}(t)S^{-1}\right)$ .

• subroutine, public prg\_quantumdynamics\_mod::prg\_getcharge (rho\_bml, s\_bml, charges, aux\_bml, z, spin-dex, N, nats, thresh)

Constructs the charges from the density matrix.

• subroutine, public prg\_quantumdynamics\_mod::prg\_getdipole (charges, r, mu)

This routine computes the dipole moment of the system with units determined by the units of the coordinate matrix and charges given.

• subroutine, public prg\_quantumdynamics\_mod::prg\_excitation (fill\_mat, orbit\_orig, orbit\_exci)

Produce an excitation in the initially calculated density matrix to.

#### **Variables**

• integer, parameter prg\_quantumdynamics\_mod::dp = kind(1.0d0)

# 11.26 /home/christian/qmd-progress/src/prg\_response\_mod.F90 File Reference

## **Data Types**

type prg\_response\_mod::respdata\_type

## Modules

• module prg\_response\_mod

Module to compute the density matrix response and related quantities.

# **Functions/Subroutines**

- subroutine, public prg\_response\_mod::prg\_parse\_response (RespData, filename)
  - The parser for the calculation of the DM response.
- subroutine, public prg\_response\_mod::prg\_compute\_dipole (charges, coordinate, dipoleMoment, factor, verbose)

To compute the dipole moment of the system. The units of the dipole moment are determined by the units of the coordinates and charges that are given.

• subroutine, public prg\_response\_mod::prg\_write\_dipole\_tcl (dipoleMoment, file, factor, verbose)

To visualize a dipole moment using VMD. This will prg\_generate a .tcl script that could be run using VMD To visualize with VMD: \$ vmd -e dipole.tcl.

subroutine, public prg\_response\_mod::prg\_compute\_polarizability (rsp\_bml, prt\_bml, polarizability, factor, verbose)

To compute the polarizability of the system. The units of the directional polarizability are determined by the units of the perturbation and Hamiltonian. This equation can be found in [5] equation 4a. Note that in equation 4a of the reference there is a 2 that account for the double occupancy which is not present in this case cause the density matrix construction is done by taking the occupancy into account.

• subroutine, public prg response mod::prg pert from file (prt bml, norb)

Read perturbation from file.

• subroutine, public prg\_response\_mod::prg\_compute\_response\_rs (ham\_bml, prt\_bml, rsp\_bml, lambda, bndfil, threshold, verbose)

Computes the first order response density matrix using Rayleigh Schrodinger Perturbation theory The transformation hereby performed are:

• subroutine, public prg\_response\_mod::prg\_compute\_response\_fd (ham\_bml, prt\_bml, rsp\_bml, prg\_delta, bndfil, threshold, verbose)

Computes the first order response density matrix using finite differences. The transformation hereby performed are:

 subroutine, public prg\_response\_mod::prg\_pert\_constant\_field (field, intensity, coordinate, lambda, prt\_bml, threshold, spindex, norbi, verbose, over bml)

Apply a constant field perturbation through the dipole moment operator (  $\hat{\mu}=e\hat{r}$ ). In the matrix representation, this is:  $H^{(1)}=\lambda \frac{1}{2}(S\,er\cdot {\bf E}+\,er\cdot {\bf E}S)$ . The symmetrization is done in order to preserve the Hermiticity of H. In this case the whole system will be affected by the field. In a latter version we will add the possibility of applying this field to a region of the system. In this implementation e=1 and units can be transformed by using the parameter  $\lambda$ .

• subroutine, public prg\_response\_mod::prg\_pert\_sin\_pot (direction, lx, coordinate, lambda, prt\_bml, threshold, spindex, norbi, verbose, over bml)

Apply a sinusoidal length dependent potential  $(\sin(\tilde{r}_x))$  where  $r_x$  is the x coordinate. The Hamiltonian gets modified as follows:  $H^{(1)} = \frac{1}{2}\lambda(S\sin(\tilde{r}_x) + \sin(\tilde{r}_x)S)$ .  $\tilde{r}_x = 2\pi(r/l_x) - \pi$ . The symmetrization is done in order to preserve the Hermiticity of H. Units can be transformed by using the parameter  $\lambda$ .

• subroutine, public prg\_response\_mod::prg\_pert\_cos\_pot (direction, lx, coordinate, lambda, prt\_bml, threshold, spindex, norbi, verbose, over bml)

Apply a cosine length dependent potential  $(\cos(\tilde{\textbf{r}}_x))$  where  $\textbf{r}_x$  is the x coordinate. The Hamiltonian gets modified as follows:  $H^{(1)} = \frac{1}{2}\lambda(S\sin(\tilde{\textbf{r}}_x) + \sin(\tilde{\textbf{r}}_x)S)$ .  $\tilde{\textbf{r}}_x = 2\pi(\textbf{r}/l_x) - \pi$ . The symmetrization is done in order to preserve the Hermiticity of H. Units can be transformed by using the parameter  $\lambda$ .

• subroutine, public prg\_response\_mod::prg\_compute\_response\_sp2 (ham\_bml, prt\_bml, rsp\_bml, rho\_bml, lambda, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, threshold, verbose)

Finds the first order response matrix from a Hamiltonian matrix.

• subroutine, public prg\_response\_mod::prg\_project\_response (rsp\_bml, over\_bml, spindex, norbi, coordinates, rspfunc, verbose)

Project the response onto atomic positions. First order response to the perturbation ( $\rho^{(1)}$ ) projected onto the atomic position. Basically:  $rsp(i) = \sum_{\alpha \in i} \rho_{\alpha\alpha}^{(1)}$ , where orbital  $\alpha$  belong to atom i.

## **Variables**

- integer, parameter prg\_response\_mod::dp = kind(1.0d0)
- real(dp), parameter prg\_response\_mod::pi = 3.14159265358979323846264338327950\_dp

# 11.27 /home/christian/gmd-progress/src/prg\_sp2\_fermi\_mod.F90\_File\_Reference

# **Modules**

· module prg sp2 fermi mod

The SP2 Fermi module.

## **Functions/Subroutines**

• subroutine, public prg\_sp2\_fermi\_mod::prg\_sp2\_fermi\_init (h\_bml, nsteps, nocc, tscale, threshold, occErr 
Limit, traceLimit, x\_bml, mu, beta, h1, hN, sgnlist)

Truncated SP2 prg\_initialization.

• subroutine, public prg\_sp2\_fermi\_mod::prg\_sp2\_fermi\_init\_norecs (h\_bml, nsteps, nocc, tscale, threshold, occErrLimit, traceLimit, x bml, mu, beta, h1, hN, sgnlist, verbose)

Truncated SP2 prg\_initialization. This routine also gives back the Number of SP2 recursive steps that gets a Pseudo-Fermi distribution with a temperature close to the target temperature which is entered using parameter beta = (1/KbT).

• subroutine, public prg\_sp2\_fermi\_mod::prg\_sp2\_fermi (h\_bml, osteps, nsteps, nocc, mu, beta, h1, hN, sgn-list, threshold, eps, traceLimit, x\_bml)

Calculate Truncated SP2.

- subroutine, public prg\_sp2\_fermi\_mod::prg\_sp2\_entropy\_function (mu, h1, hN, nsteps, sgnlist, GG, ee)

  Calculate SP2 entropy function using gaussian quadrature. Note that GG and ee are allocated and returned from this routine.
- real(dp) function, public prg\_sp2\_fermi\_mod::sp2\_entropy\_ts (D0\_bml, GG, ee)

Test SP2 entropy. Get the entropy contribution TS to the total free energy.

- real(dp) function, public prg\_sp2\_fermi\_mod::sp2\_inverse (f, mu, h1, hN, nsteps, sgnlist)
   Calculate the SP2 inverse.
- real(dp) function prg\_sp2\_fermi\_mod::absmaxderivative (func, de)

Gets the absolute maximum of the derivative of a function.

#### **Variables**

• integer, parameter prg\_sp2\_fermi\_mod::dp = kind(1.0d0)

# 11.28 /home/christian/qmd-progress/src/prg\_sp2\_mod.F90 File Reference

#### **Modules**

module prg\_sp2\_mod

The SP2 module.

# **Functions/Subroutines**

• subroutine, public prg\_sp2\_mod::prg\_sp2\_basic (h\_bml, rho\_bml, threshold, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, verbose)

Calculates the density matrix from a Hamiltonian matrix by purification. The method implemented here is the very first verion of the SP2 method.

- subroutine, public prg\_sp2\_mod::prg\_sp2\_alg2 (h\_bml, rho\_bml, threshold, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, verbose)
- subroutine, public prg\_sp2\_mod::prg\_sp2\_alg2\_genseq (h\_bml, rho\_bml, threshold, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, pp, icount, vv, verbose)
- subroutine, public prg\_sp2\_mod::prg\_sp2\_alg2\_seq (h\_bml, rho\_bml, threshold, pp, icount, vv, verbose)
- subroutine, public prg\_sp2\_mod::prg\_prg\_sp2\_alg2\_seq\_inplace (rho\_bml, threshold, pp, icount, vv, mineval, maxeval, verbose)
- subroutine, public prg\_sp2\_mod::prg\_sp2\_alg1 (h\_bml, rho\_bml, threshold, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, verbose)
- subroutine, public prg\_sp2\_mod::prg\_sp2\_alg1\_genseq (h\_bml, rho\_bml, threshold, bndfil, minsp2iter, maxsp2iter, sp2conv, idemtol, pp, icount, vv)
- subroutine, public prg\_sp2\_mod::prg\_sp2\_alg1\_seq (h\_bml, rho\_bml, threshold, pp, icount, vv)
- subroutine, public prg\_sp2\_mod::prg\_prg\_sp2\_alg1\_seq\_inplace (rho\_bml, threshold, pp, icount, vv, mineval, maxeval)
- subroutine, public prg\_sp2\_mod::prg\_sp2\_submatrix (ham\_bml, rho\_bml, threshold, pp, icount, vv, mineval, maxeval, core\_size)

Perform SP2 algorithm using sequence and calculate norm for a submatrix.

• subroutine, public prg\_sp2\_mod::prg\_sp2\_submatrix\_inplace (rho\_bml, threshold, pp, icount, vv, mineval, maxeval, core size)

#### **Variables**

• integer, parameter prg\_sp2\_mod::dp = kind(1.0d0)

# 11.29 /home/christian/qmd-progress/src/prg\_sp2parser\_mod.F90 File Reference

# **Data Types**

type prg\_sp2parser\_mod::sp2data\_type
 General SP2 solver type.

#### **Modules**

 module prg\_sp2parser\_mod SP2 parser.

# **Functions/Subroutines**

subroutine, public prg\_sp2parser\_mod::prg\_parse\_sp2 (sp2data, filename)
 The parser for SP2 solver.

## **Variables**

• integer, parameter prg\_sp2parser\_mod::dp = kind(1.0d0)

# 11.30 /home/christian/qmd-progress/src/prg\_subgraphloop\_mod.F90 File Reference

#### **Modules**

module prg\_subgraphloop\_mod
 The subgraphloop module.

# **Functions/Subroutines**

- subroutine, public prg\_subgraphloop\_mod::prg\_subgraphsp2loop (h\_bml, g\_bml, rho\_bml, gp, threshold)
- subroutine, public prg\_subgraphloop\_mod::prg\_collectmatrixfromparts (gp, rho\_bml) Collect distributed parts into same matrix.
- subroutine, public prg\_subgraphloop\_mod::prg\_balanceparts (gp)
- subroutine, public prg\_subgraphloop\_mod::prg\_partordering (gp)

Set row ordering bases on parts.

- subroutine, public prg\_subgraphloop\_mod::prg\_getgrouppartitionhalosfromgraph (gp, g\_bml, hnode, djflag)

  Get core+halo indeces for all partitions only using the graph.
- subroutine, public prg\_subgraphloop\_mod::prg\_getpartitionhalosfromgraph (gp, g\_bml, djflag)

  Get core+halo indeces for all partitions only using the graph.

## **Variables**

• integer, parameter prg\_subgraphloop\_mod::dp = kind(1.0d0)

# 11.31 /home/christian/qmd-progress/src/prg\_syrotation\_mod.F90 File Reference

# **Data Types**

type prg\_syrotation\_mod::rotation\_type
 Rotation type.

## **Modules**

• module prg\_syrotation\_mod

A module to rotate the coordinates of a sybsystem in chemical systems.

# **Functions/Subroutines**

• subroutine, public prg\_syrotation\_mod::prg\_parse\_rotation (rot, filename)

The parser for rotation.

• subroutine, public prg\_syrotation\_mod::prg\_rotate (rot, r, verbose)

Rotation routine.

# **Variables**

• integer, parameter prg\_syrotation\_mod::dp = kind(1.0d0)

# 11.32 /home/christian/qmd-progress/src/prg\_system\_mod.F90 File Reference

# **Data Types**

• type prg\_system\_mod::estruct\_type

Electronic structure type.

• type prg\_system\_mod::system\_type

System type.

## **Modules**

module prg\_system\_mod

A module to read and handle chemical systems.

## **Functions/Subroutines**

subroutine, public prg\_system\_mod::prg\_get\_nameandext (fullfilename, filename, ext)

Get the name and extension of a file.

• subroutine, public prg\_system\_mod::prg\_parse\_system (system, filename, extin)

The parser for the chemical system.

• subroutine, public prg\_system\_mod::prg\_write\_system (system, filename, extension)

Write system in .xyz, .dat or pdb file.

• subroutine, public prg\_system\_mod::prg\_write\_trajectory (system, iter, each, prg\_deltat, filename, extension)

Write trajectory in .xyz, .dat or pdb file.

• subroutine, public prg\_system\_mod::prg\_write\_trajectoryandproperty (system, iter, each, prg\_deltat, scalarprop, filename, extension)

Write trajectory and atomic properties. Only pdb file.

• subroutine, public prg\_system\_mod::prg\_make\_random\_system (system, nats, seed, lx, ly, lz)

Make random Xx system.

• subroutine prg\_system\_mod::prg\_parameters\_to\_vectors (abc\_angles, lattice\_vector)

Transforms the lattice parameters into lattice vectors.

subroutine prg\_system\_mod::prg\_vectors\_to\_parameters (lattice\_vector, abc\_angles)

Transforms the lattice vectors into lattice parameters.

• subroutine, public prg\_system\_mod::prg\_get\_origin (coords, origin)

Get the origin of the coordinates.

subroutine, public prg\_system\_mod::prg\_get\_distancematrix (coords, dmat)

Get the distance matrix.

• subroutine, public prg\_system\_mod::prg\_translateandfoldtobox (coords, lattice\_vectors, origin, verbose)

Translate and fold to box.

subroutine, public prg\_system\_mod::prg\_centeratbox (coords, lattice\_vectors, verbose)

Translate geometric center to the center of the box.

• subroutine, public prg\_system\_mod::prg\_wraparound (coords, lattice\_vectors, index, verbose)

Wrap around atom i using pbc.

• subroutine, public prg\_system\_mod::prg\_translatetogeomcandfoldtobox (coords, lattice\_vectors, origin)

Translate to geometric center.

• subroutine, public prg\_system\_mod::prg\_replicate (coords, symbols, lattice\_vectors, nx, ny, nz)

Extend/replicate system along lattice vectors.

• subroutine, public prg\_system\_mod::prg\_get\_recip\_vects (lattice\_vectors, recip\_vectors, volr, volk)

Get the volume of the cell and the reciprocal vectors: This soubroutine computes:

subroutine, public prg\_system\_mod::prg\_get\_dihedral (coords, id1, id2, id3, id4, dihedral)

Get the dihedral angle given four atomic positions.

• subroutine, public prg\_system\_mod::prg\_get\_covgraph (sy, nnStructMindist, nnStruct, nrnnstruct, bml\_type, factor, gcov\_bml, mdimin, verbose)

Get the covalency graph in bml format.

- subroutine prg\_system\_mod::prg\_get\_covgraph\_int (sy, nnStructMindist, nnStruct, nrnnstruct, bml\_type, factor, gcov\_bml, mdimin, verbose)
- subroutine, public prg\_system\_mod::prg\_get\_covgraph\_h (sy, nnStructMindist, nnStruct, nrnnstruct, rcut, graph\_h, mdimin, verbose)

Get the covanlency graph.

• subroutine, public prg\_system\_mod::prg\_get\_subsystem (sy, Isize, indices, sbsy, verbose)

Get a subsystem out of the total system.

• subroutine, public prg\_system\_mod::prg\_destroy\_subsystems (sbsy, verbose)

Destroy allocated subsystem.

• subroutine, public prg\_system\_mod::prg\_molpartition (sy, npart, nnStructMindist, nnStruct, nrnnstruct, hetatm, gp, verbose)

Partition by molecule.

subroutine, public prg\_system\_mod::prg\_get\_partial\_atomgraph (rho\_bml, hindex, gch\_bml, threshold, ver-bose)

Get partial subgraph based on the Density matrix.

subroutine, public prg\_system\_mod::prg\_collect\_graph\_p (rho\_bml, nc, nats, hindex, chindex, graph\_←
p, threshold, mdimin, verbose)

Collect the small graph to build the full graph.

subroutine, public prg system mod::prg merge graph (graph p, graph h)

Get partial subgraph based on the Density matrix.

• subroutine, public prg\_system\_mod::prg\_merge\_graph\_adj (graph\_p, graph\_h, xadj, adjncy)

Get partial subgraph based on the Density matrix.

• subroutine, public prg\_system\_mod::prg\_adj2bml (xadj, adjncy, bml\_type, g\_bml)

prg\_adj2bml

• subroutine, public prg\_system\_mod::prg\_graph2bml (graph, bml\_type, g\_bml)

Graph2bml.

subroutine, public prg\_system\_mod::prg\_graph2vector (graph, vector, maxnz)

Vectorize graph.

• subroutine, public prg\_system\_mod::prg\_vector2graph (vector, graph, maxnz)

Back to graph.

subroutine, public prg\_system\_mod::prg\_sortadj (xadj, adjncy)

Sort adj NOTE: this might not be needed anymre since the bml\_get\_adj routine is sorting the values.

#### **Variables**

integer, parameter prg\_system\_mod::dp = kind(1.0d0)

# 11.33 /home/christian/qmd-progress/src/prg\_timer\_mod.F90 File Reference

# **Data Types**

• type prg\_timer\_mod::timer\_status\_t

Timer status type.

#### **Modules**

module prg\_timer\_mod

The timer module.

# **Functions/Subroutines**

• subroutine, public prg\_timer\_mod::timer\_prg\_init()

Initialize timers.

• subroutine prg\_timer\_mod::prg\_timer\_getid ()

Get timer id.

• subroutine, public prg\_timer\_mod::prg\_timer\_shutdown ()

Done with timers.

subroutine, public prg\_timer\_mod::prg\_timer\_start (itimer, tag)

Start Timing.

• subroutine, public prg\_timer\_mod::prg\_timer\_stop (itimer, verbose)

Stop timing.

- subroutine, public prg\_timer\_mod::prg\_timer\_collect ()
- subroutine, public prg\_timer\_mod::prg\_timer\_results ()
- real(8) function, public prg\_timer\_mod::time2milliseconds ()
- subroutine, public prg\_timer\_mod::prg\_print\_date\_and\_time (tag)
- character(2) function, private prg\_timer\_mod::int2char (ival)

## **Variables**

- integer, parameter prg\_timer\_mod::dp = kind(1.0d0)
- integer, public prg\_timer\_mod::loop\_timer
- integer, public prg\_timer\_mod::sp2\_timer
- · integer, public prg\_timer\_mod::genx\_timer
- integer, public prg\_timer\_mod::part\_timer
- integer, public prg\_timer\_mod::subgraph\_timer
- integer, public prg\_timer\_mod::deortho\_timer
- integer, public prg\_timer\_mod::ortho\_timer
- integer, public prg\_timer\_mod::zdiag\_timer
- integer, public prg\_timer\_mod::graphsp2\_timer
- integer, public prg\_timer\_mod::subind\_timer
- integer, public prg\_timer\_mod::subext\_timer
- integer, public prg\_timer\_mod::subsp2\_timer
- · integer, public prg\_timer\_mod::suball\_timer
- integer, public prg\_timer\_mod::bmult\_timer
- integer, public prg\_timer\_mod::badd\_timer
- integer, public prg\_timer\_mod::dyn\_timer
- · integer, public prg\_timer\_mod::mdloop\_timer
- integer, public prg\_timer\_mod::buildz\_timer
- integer, public prg\_timer\_mod::realcoul\_timer
- integer, public prg\_timer\_mod::recipcoul\_timer
- · integer, public prg\_timer\_mod::pairpot\_timer
- integer, public prg\_timer\_mod::halfverlet\_timer
- integer, public prg\_timer\_mod::pos\_timer
- integer, public prg\_timer\_mod::nlist\_timer
- integer prg\_timer\_mod::tstart\_clock
- integer prg\_timer\_mod::tstop\_clock
- integer prg\_timer\_mod::tclock\_rate
- integer prg\_timer\_mod::tclock\_max
- integer prg\_timer\_mod::num\_timers
- type(timer\_status\_t), dimension(:), allocatable prg\_timer\_mod::ptimer

# 11.34 /home/christian/qmd-progress/src/prg\_xlbo\_mod.F90 File Reference

# **Data Types**

type prg\_xlbo\_mod::xlbo\_type

General xlbo solver type.

#### **Modules**

module prg\_xlbo\_mod

A module to perform XLBO integration.

## **Functions/Subroutines**

- subroutine, public prg\_xlbo\_mod::prg\_parse\_xlbo (xlbo, filename)
  - The parser for XLBO parser.
- subroutine, public prg\_xlbo\_mod::prg\_xlbo\_nint (charges, n, n\_0, n\_1, n\_2, n\_3, n\_4, n\_5, mdstep, xl)
  - This routine integrates the dynamical variable "n".
- subroutine, public prg\_xlbo\_mod::prg\_xlbo\_fcoulupdate (fcoul, charges, n)
  - Adjust forces for the linearized XLBOMD functional.

#### **Variables**

- integer, parameter prg\_xlbo\_mod::dp = kind(1.0d0)
- real(dp), parameter prg\_xlbo\_mod::c0 = -6.0\_dp

Coefficients for modified Verlet integration.

- real(dp), parameter prg\_xlbo mod::c1 = 14.0 dp
- real(dp), parameter prg xlbo mod::c2 = -8.0 dp
- real(dp), parameter prg\_xlbo\_mod::c3 = -3.0\_dp
- real(dp), parameter prg\_xlbo\_mod::c4 = 4.0\_dp
- real(dp), parameter prg\_xlbo\_mod::c5 = -1.0\_dp
- real(dp), parameter prg\_xlbo\_mod::kappa = 1.82\_dp

Coefficients for modified Verlet integration.

- real(dp), parameter prg\_xlbo mod::alpha = 0.018 dp
- real(dp), parameter prg\_xlbo mod::cc = 0.9 dp

# 11.35 /home/christian/qmd-progress/src/prg\_xlkernel\_mod.F90 File Reference

## **Data Types**

type prg\_xlkernel\_mod::xlk\_type

# Modules

• module prg\_xlkernel\_mod

Add name.

## **Functions/Subroutines**

- subroutine, public prg\_xlkernel\_mod::prg\_parse\_xlkernel (input, filename)

  The parser for the mixer routines.
- subroutine, public prg\_xlkernel\_mod::prg\_fermi (D0, QQ, ee, gap, Fe\_vec, mu0, H, Z, Nocc, T, OccErrLim, MaxIt, HDIM)
- subroutine, public prg\_xlkernel\_mod::prg\_kernel\_fermi\_full (KK, JJ, D0, mu0, mu1, T, RX, RY, RZ, LBox, Hubbard\_U, Element\_Type, Nr\_atoms, MaxIt, eps, m, HDIM, Max\_Nr\_Neigh, Coulomb\_acc, TIMERAT ← IO, nnRx, nnRy, nnRz, nrnnlist, nnType, H\_INDEX\_START, H\_INDEX\_END, H, S, Z, Nocc, Znuc, QQ, ee, Fe vec)
- subroutine, public prg\_xlkernel\_mod::prg\_v\_kernel\_fermi (D0, dq\_dv, v, mu0, mu1, T, RX, RY, RZ, LBox, Hubbard\_U, Element\_Type, Nr\_atoms, MaxIt, eps, m, HDIM, Max\_Nr\_Neigh, Coulomb\_acc, TIMERAT ← IO, nnRx, nnRy, nnRz, nrnnlist, nnType, H\_INDEX\_START, H\_INDEX\_END, H, S, Z, Nocc, Znuc, QQ, ee, Fe\_vec)
- subroutine, private prg\_xlkernel\_mod::prg\_get\_deriv\_finite\_temp (P1, H0, H1, Nocc, T, Q, ev, fe, mu0, eps, HDIM)
- subroutine, private prg\_xlkernel mod::prg\_mmult (alpha, A, B, beta, C, TA, TB, HDIM)
- subroutine, private prg\_xlkernel\_mod::prg\_eig (A, Q, ee, type, HDIM)
- subroutine, private prg\_xlkernel mod::prg\_inv (X, XI, HDIM)
- subroutine, public prg\_xlkernel\_mod::prg\_rank1 (verbose)

Rank1 kernel ....

# Variables

• integer, parameter prg\_xlkernel\_mod::dp = kind(1.0d0)

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