# SPEC

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# 1 The Stepped Pressure Equilibrium Code

All relevant publications and presentations are given on the MRxMHD website.

A BibTex file is available: spec\_refs.bib.

- There is also a brief description of SPEC on the Princeton Plasma Physics Laboratory Theory Department website.
- An overview of the algorithm is given in a Flow Chart.
- Please direct questions/suggestions to Stuart Hudson.
- Online documentation for SPEC is available on Github pages.
   A PDF version of this manual is available: SPEC\_manual.pdf

# 2 Compilation hints for SPEC

In order to run SPEC, you need a copy of the HDF5 libraries installed which has both the Fortran interface and the parallel (MPI I/O) enabled.

# 2.1 Installation with CMake

Using cmake, SPEC can be built as a stand-alone executable or as a python extension, where SPEC can be run directly from python, with all variables passed directly in memory.

Download the package from git. And change to the root directory of SPEC source code by running  $_{\text{cd}}$   $_{\text{SPEC\_ROOT}}$ 

# 2.1.1 Stand-alone Executable Compiling

Compiling SPEC requires MPI, HDF5, and numerical libraries such as BLAS, LAPACK, FFTW. For numerical libraries, you could use system supplied libraries or you could use intel math kernel library (MKL).

#### **2.1.1.1 CentOS** Here instructions are given for CentOS 7

#### 2.1.1.1 Dependencies Install OpenBLAS, FFTW3, and hdf5 using the command

```
yum install -y gcc-gfortran openmpi openmpi-devel hdf5 hdf5-devel fftw3 fftw3-devel openblas openblas-devel python3 python3-devel cmake ninja-build
```

If you don't have the latest version of cmake available on your system, you can create a python virtual environment ( instructions are here), activate it, and then install cmake in that virtual environment using pip pip install cmake ninja

**2.1.1.1.2 Configure** When using cmake to build SPEC, the first step is to configure compilers and the locations of libraries. Cmake can detect compilers and libraries at standard locations easily but needs hand-holding when the required libraries are non-standard locations.

The following command was used to configure cmake build setup for SPEC on Centos

There are few points to note on the above command

- · All the build related files will be in build folder.
- Ninja build system is used. If your system doesn't have ninja installed, remove the -G option. The default is the standard make tool.
- We ae using OpenBLAS for BLAS and LAPACK and MPI fortran compiler
- Since most of the libraries are in standard location, we don't have to specify them. We are giving couple of options related to HDF5 libraries.
- The installation path is install subfolder location within SPEC folder.
- We are interested in a verbose output and also want to store the output in log file.

#### 2.1.1.1.3 Build After successful completion of cmake configuration step, building is trivial

```
cmake --build build
""
#### Install
The last step is to install the executable by running
"bash
cmake --install build
```

That's it! If all the above steps completed without errors, you have the SPEC executable xspec installed at install/bin folder

# 2.1.2 Python Extension Compiling

Building the SPEC python extension will also build the SPEC executable. In the SPEC root folder, edit the  $cmake\_config.json$  as necessary for your system. Few example .json files are provided in the  $cmake\_config.json$  machines folder.

**2.1.2.1 Dependencies** It is strongly suggested to use a python virtual environment either conda or python venv. After virtual environment is installed and activated, install the python related dependencies. Please note that these are in addition to the dependencies listed earlier in stand-alone installation steps. If you are using conda virtual environment try installing the dependencies using conda install command

conda install -n <your\_venv> numpy f90nml scikit-build cmake ninja

#### If you are using venv virtual environment, run

pip install numpy f90nml scikit-build cmake ninja

Now install f90wrap. Please keep in mind that numpy has to be installed before installing f90wrap.

pip install -U git+https://github.com/zhucaoxiang/f90wrap

Now install the SPEC extension by running the setup.py script present in the SPEC root folder.

python setup.py bdist\_wheel; cd dist/; pip install \*.whl

in succession. At this point, you should be able to import the spec module in python. To test this, you can try the following command from the shell:

python -c "import spec; print('success')"

#### If you want editable install, run

python setup.py develop

#### 2.1.3 Stellar cluster at PPPL

- 2.1.3.1 Python wrapper Below are the steps to build python wrappers for SPEC on stellar.
  - 1. Needed modules are

i. hdf5/gcc/1.10.6 ii. intel-mkl/2021.1.1 iii. openmpi/gcc/4.1.0

#### 2.1.4 iv. anaconda3/2021.5.

# Note

# FFTW is supplied as part of Intel MKL and we just need to link against MKL.

Load the modules by running

module load hdf5/gcc/1.10.6 intel-mkl/2021.1.1 openmpi/gcc/4.1.0 anaconda3/2021.5

1. Create conda virtual environment.

conda create -n spec\_ve python=3.8

You have to press enter twice. Here a conda virtual environment named <code>spec\_ve</code> is created with python version 3.8 and lot of packages are installed. Activate by running <code>conda activate spec\_ve</code>

2. Install cmake, ninja, scikit-build, numpy using either conda or pip.

conda install cmake ninja scikit-build numpy

or

pip install cmake ninja scikit-build numpy

3. Install f90wrap by running

pip install git+https://github.com/zhucaoxiang/f90wrap.git

4. Clone the spec repo from github

git clone https://github.com/PrincetonUniversity/SPEC.git

Change the working directory by running cd SPEC.

5. Edit the cmake\_config.json to populate correct cmake\_flags. For stellar, cmake\_config.json should look like

```
"cmake_args": [
   "-DCMAKE_C_COMPILER=mpicc",
   "-DCMAKE_CXX_COMPILER=mpicxx",
   "-DCMAKE_CXX_COMPILER=mpifort",
   "-DBLA_VENDOR=Intel10_641p",
   "-DHDF5_ROOT=/usr/local/hdf5/gcc/1.10.6",
   "-DHDF5_PREFER_PARALLEL=False"]
```

6. Then build the python wheel for SPEC wrapper using

```
python setup.py bdist_wheel
```

The resulting wheel is located in dist folder. Install SPEC python wrapper by running pip install dist/spec\*.whl

 Install mpi4py using pip/conda. If using pip, don't forget to use --no-cache-dir flag pip install --no-cache-dir mpi4py

```
Or conda install mpi4py
```

- **2.1.4.1 SPEC executable.** The python wrapper builds spec executable but it gets installed at an obscure location. If you mainly want SPEC executable xspec, the steps are similar.
  - 1. Load the required modules. Refer to the first step in the python wrapper instructions.
  - 2. Clone the SPEC repo and make SPEC as working directory. Refer to the 5th step above.
  - 3. Run the cmake configuration by running

```
cmake -Bbuild -S . -DCMAKE_C_COMPILER=mpicc -DCMAKE_CXX_COMPILER=mpicxx
-DCMAKE_Fortran_COMPILER=mpifort -DBLA_VENDOR=Intel10_641p \
-DHDF5_ROOT=/usr/local/hdf5/gcc/1.10.6 -DHDF5_PREFER_PARALLEL=False
-DCMAKE_INSTALL_PREFIX=<SPEC_install_location>
```

Please note SPEC gets installed at <SPEC\_install\_location>/bin, where <SPEC\_install $\leftarrow$  \_location> is the folder of your choice. Building of SPEC library will be done in the folder build, where all the intermediary compilation files will be located.

4. Compile the code by running

```
cmake --build build
```

This command will invoke make build generator. Alternatively, you can switch to build folder and run make utility manually.

```
cd build
```

5. Install the SPEC executable by running

```
cmake --install build
```

SPEC library gets installed <SPEC\_install\_location>/lib and SPEC executable get installed at <SPEC\_install\_location>/bin

#### 2.1.5 Mac

Here is how to build the HDF5 library:

- 1. download hdf5-1.10.5.tar.gz from https://www.hdfgroup.org/downloads/hdf5/source-code/
- 2. extract: tar xzf hdf5-1.10.5.tar.gz
- 3. cd into source folder: cd hdf5-1.10.5
- 4. make a build folder: mkdir build
- 5. cd into build folder: cd build
- 6. run cmake with options for the Fortran interface:  $cmake DHDF5\_BUILD\_FORTRAN:BOOL=ON$  ..
- 7. actually build the HDF5 library: make

The compilation of SPEC itself then proceeds as usual. You then only need to specify the HDF5 folder in the Makefile, which will likely be  $\protect\end{Applications/HDF\_Group/HDF5/1.10.5}$ .

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# 3 Todo List

#### Subprogram bnorml (mn, Ntz, efmn, ofmn)

There is a very clumsy attempt to parallelize this which could be greatly improved.

# Subprogram casing (teta, zeta, gBn, icasing)

It would be MUCH faster to only require the tangential field on a regular grid!!!

Please check why  $B_s$  is not computed. Is it because  $B_s \nabla s \times \mathbf{n} = 0$ ?

This needs to be revised.

# Subprogram curent (Ivol, mn, Nt, Nz, iflag, IdltGp)

Perhaps this can be proved analytically; in any case it should be confirmed numerically.

#### Subprogram inputlist::lconstraint

if Lconstraint==2, under reconstruction.

# Subprogram inputlist::wbuild\_vector\_potential

: what is this?

# Type intghs\_module::intghs\_workspace

Zhisong might need to update the documentation of this type.

# Subprogram ma02aa (Ivol, NN)

If Lconstraint = 2, then  $\mu=\mu_1$  is varied in order to satisfy the helicity constraint, and  $\Delta\psi_p=\mu_2$  is not varied, and Nxdof=1. (under re-construction)

#### Subprogram pc00aa (NGdof, position, Nvol, mn, ie04dgf)

Unfortunately, E04DGF seems to require approximately 3N function evaluations before proceeding to minimize the energy functional, where there are N degrees of freedom. I don't know how to turn this off!

# Subprogram pc00ab (mode, NGdof, Position, Energy, Gradient, nstate, iuser, ruser)

IT IS VERY LIKELY THAT FFTs WOULD BE FASTER!!!

#### Subprogram spec

If Lminimize.eq.1, call pc00aa() to find minimum of energy functional using quasi-Newton, preconditioned conjugate gradient method, E04DGF

# Subprogram stzxyz (Ivol, stz, RpZ)

Please see co01aa() for documentation.

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	g.f90 tructs the field created by the plasma currents, at an arbitrary, external location using I casing	171
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src/spsint.f90 Calculates volume integrals of Chebyshev-polynomials and metric elements for preconditioner	225
src/spsmat.f90 Constructs matrices for the precondtioner	225
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# 7 Module Documentation

# 7.1 Diagnostics to check the code

# **Functions/Subroutines**

• subroutine bfield (zeta, st, Bst)

Compute the magnetic field.

• subroutine hesian (NGdof, position, Mvol, mn, LGdof)

Computes eigenvalues and eigenvectors of derivative matrix,  $\nabla_{\xi} \mathbf{F}$ .

• subroutine jo00aa (Ivol, Ntz, Iquad, mn)

Measures error in Beltrami equation,  $\nabla \times \mathbf{B} - \mu \mathbf{B}$ .

• subroutine pp00aa

Constructs Poincaré plot and "approximate" rotational-transform (driver).

• subroutine pp00ab (Ivol, sti, Nz, nPpts, poincaredata, fittedtransform, utflag)

Constructs Poincaré plot and "approximate" rotational-transform (for single field line).

• subroutine stzxyz (Ivol, stz, RpZ) Calculates coordinates,  $\mathbf{x}(s, \theta, \zeta) \equiv R \, \mathbf{e}_R + Z \, \mathbf{e}_Z$ , and metrics, at given  $(s, \theta, \zeta)$ .

#### 7.1.1 Detailed Description

#### 7.1.2 Function/Subroutine Documentation

# 

Compute the magnetic field.

Returns the magnetic field line equations,  $d\mathbf{x}/d\phi = \mathbf{B}/B^{\phi}$ .

# **Equations of field line flow**

• The equations for the fieldlines are normalized to the toroidal field, i.e.

$$\dot{s} \equiv \frac{B^s}{B^{\zeta}}, \qquad \dot{\theta} \equiv \frac{B^{\theta}}{B^{\zeta}}. \tag{1}$$

#### Representation of magnetic field

• The components of the vector potential,  $\mathbf{A} = A_{\theta} \nabla + A_{\zeta} \nabla \zeta$ , are

$$A_{\theta}(s,\theta,\zeta) = \sum_{i,l} A_{\theta,e,i,l} \, \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} A_{\theta,o,i,l} \, \overline{T}_{l,i}(s) \sin \alpha_i, \tag{2}$$

$$A_{\zeta}(s,\theta,\zeta) = \sum_{i,l} A_{\zeta,e,i,l} \, \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} A_{\zeta,o,i,l} \, \overline{T}_{l,i}(s) \sin \alpha_i, \tag{3}$$

where  $\overline{T}_{l,i}(s) \equiv \overline{s}^{m_i/2} T_l(s)$ ,  $T_l(s)$  is the Chebyshev polynomial, and  $\alpha_j \equiv m_j \theta - n_j \zeta$ . The regularity factor,  $\overline{s}^{m_i/2}$ , where  $\overline{s} \equiv (1+s)/2$ , is only included if there is a coordinate singularity in the domain (i.e. only in the innermost volume, and only in cylindrical and toroidal geometry.)

• The magnetic field,  $\sqrt{g}\,{f B}=\sqrt{g}B^s{f e}_s+\sqrt{g}B^\theta{f e}_\theta+\sqrt{g}B^\zeta{f e}_\ell$ , is

$$\sqrt{g} \mathbf{B} = \mathbf{e}_{s} \sum_{i,l} \left[ (-m_{i} A_{\zeta,e,i,l} - n_{i} A_{\theta,e,i,l}) \overline{T}_{l,i} \sin \alpha_{i} + (+m_{i} A_{\zeta,o,i,l} + n_{i} A_{\theta,o,i,l}) \overline{T}_{l,i} \cos \alpha_{i} \right] 
+ \mathbf{e}_{\theta} \sum_{i,l} \left[ (-m_{i} A_{\zeta,e,i,l} - n_{i} A_{\theta,e,i,l}) \overline{T}'_{l,i} \cos \alpha_{i} + (-m_{i} A_{\zeta,o,i,l} + n_{i} A_{\theta,o,i,l}) \overline{T}'_{l,i} \sin \alpha_{i} \right] 
+ \mathbf{e}_{\zeta} \sum_{i,l} \left[ (-m_{i} A_{\zeta,e,i,l} - n_{i} A_{\theta,e,i,l}) \overline{T}'_{l,i} \cos \alpha_{i} + (-m_{i} A_{\zeta,o,i,l} + n_{i} A_{\theta,o,i,l}) \overline{T}'_{l,i} \sin \alpha_{i} \right]$$

$$(4)$$

• In Eqn. (1), the coordinate Jacobian,  $\sqrt{g}$ , cancels. No coordinate metric information is required to construct the fieldline equations from the magnetic vector potential.

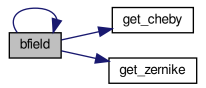
IT IS REQUIRED TO SET IVOL THROUGH GLOBAL MEMORY BEFORE CALLING BFIELD. The format of this subroutine is constrained by the NAG ode integration routines.

#### **Parameters**

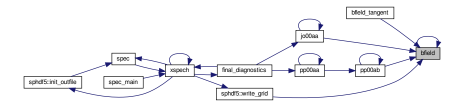
in	$zeta$ toroidal angle $\zeta$	
in	st	radial coordinate $s$ and poloidal angle $\theta$
out	Bst	tangential magnetic field directions $B_s, B_{ heta}$

References allglobal::ate, allglobal::ate, allglobal::aze, allglobal::aze, bfield(), allglobal::cpus, allglobal::gbzeta, get\_cheby(), get\_zernike(), constants::half, allglobal::halfmm, allglobal::im, allglobal::in, allglobal::ivol, allglobal::lcoordinatesingularity, inputlist::lrad, allglobal::mn, allglobal::mpi\_comm\_spec, inputlist::mpol, allglobal::myid, allglobal::ncpu, allglobal::node, allglobal::notstellsym, constants::one, fileunits::ounit, allglobal::regumm, numerical::small, constants::two, numerical::vsmall, inputlist::wmacros, and constants::zero.

Referenced by bfield(), bfield\_tangent(), jo00aa(), pp00ab(), and sphdf5::write\_grid(). Here is the call graph for this function:



Here is the caller graph for this function:



```
7.1.2.2 hesian() subroutine hesian (
    integer, intent(in) NGdof,
    real, dimension(0:ngdof) position,
    integer, intent(in) Mvol,
    integer, intent(in) mn,
    integer, intent(in) LGdof)
```

Computes eigenvalues and eigenvectors of derivative matrix,  $\nabla_{\varepsilon} \mathbf{F}$ .

#### **Parameters**

in	NGdof	number of global degrees of freedom
in,out	position	internal geometrical degrees of freedom
in	Mvol	total number of volumes in computation
in	mn	number of Fourier harmonics
in	LGdof	what is this?

# construction of Hessian matrix

- The routine dforce() is used to compute the derivatives, with respect to interface geometry, of the force imbalance harmonics,  $[[p+B^2/2]]_j$ , which may be considered to be the "physical" constraints, and if Igeometry==3 then also the derivatives of the "artificial" spectral constraints,  $I_j \equiv (R_\theta X + Z_\theta Y)_j$ .
- The input variable Lconstraint determines how the enclosed fluxes,  $\Delta \psi_t$  and  $\Delta \psi_p$ , and the helicity multiplier,  $\mu$ , vary as the geometry is varied; see global.f90 and mp00ac() for more details.

#### construction of eigenvalues and eigenvectors

• If LHevalues==T then the eigenvalues of the Hessian are computed using the NAG routine F02EBF.

- If LHevectors==T then the eigenvalues and the eigenvectors of the Hessian are computed.
- Note that if Igeometry==3, then the derivative-matrix also contains information regarding how the "artificial" spectral constraints vary with geometry; so, the eigenvalues and eigenvectors are not purely "physical".
- The eigenvalues and eigenvectors (if required) are written to the file .ext.GF.ev as follows:

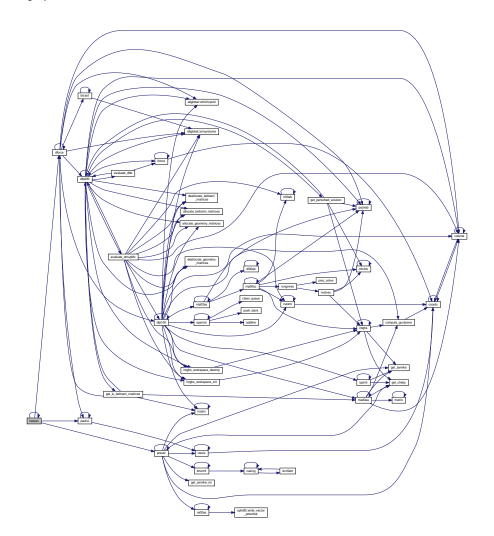
```
open(hunit,file="."//trim(ext)//".GF.ev",status="unknown",form="unformatted")
write(hunit)ngdof,ldvr,ldvi ! integers; if only the eigenvalues were computed then Ldvr=Ldvi=1;
write(hunit)evalr(1:ngdof) ! reals ; real part of eigenvalues;
write(hunit)evalr(1:ngdof) ! reals ; imaginary part of eigenvalues;
write(hunit)evecr(1:ngdof,1:ngdof) ! reals ; real part of eigenvalues; only if Ldvr=NGdof;
write(hunit)eveci(1:ngdof,1:ngdof) ! reals ; imaginary part of eigenvalues; only if Ldvi=NGdof;
close(hunit)
```

 The eigenvectors are saved in columns of evecr, eveci, as described by the NAG documentation for F02EBF.

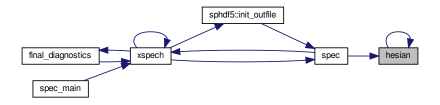
References allglobal::cpus, allglobal::dbbdmp, allglobal::dbbdrz, allglobal::dessian, allglobal::dffdrz, dforce(), allglobal::dmupfdx, inputlist::dpp, inputlist::dqq, allglobal::drbc, allglobal::drbs, allglobal::dzbc, allglobal::dzbc, allglobal::dzbc, allglobal::dzbc, allglobal::dzbc, allglobal::dzbc, allglobal::iputlist::ipermetry, allglobal::in, allglobal::irbc, allglobal::irbs, allglobal::izbc, allglobal::izbs, allglobal::lbbintegral, inputlist::lfindzero, inputlist::lfreebound, allglobal::lhessianallocated, inputlist::lhevalues, inputlist::lhevactors, inputlist::lhmatrix, allglobal::ncpu, allglobal::ncpu, allglobal::notstellsym, inputlist::nvol, constants::one, fileunits::ounit, packxi(), inputlist::pflux, preset(), allglobal::psifactor, numerical::small, numerical::sqrtmachprec, constants::ten, constants::two, numerical::vsmall, inputlist::wmacros, allglobal::yesstellsym, and constants::zero.

Referenced by hesian(), and spec().

Here is the call graph for this function:



Here is the caller graph for this function:



# 

Measures error in Beltrami equation,  $\nabla \times \mathbf{B} - \mu \mathbf{B}$ .

integer, intent(in) mn )

This routine is called by xspech() as a post diagnostic and only if Lcheck==1.

construction of current,  $\mathbf{j} \equiv \nabla \times \nabla \times \mathbf{A}$ 

• The components of the vector potential,  $\mathbf{A} = A_{\theta} \nabla + A_{\zeta} \nabla \zeta$ , are

$$A_{\theta}(s,\theta,\zeta) = \sum_{i,l} A_{\theta,e,i,l} \, \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} A_{\theta,o,i,l} \, \overline{T}_{l,i}(s) \sin \alpha_i, \tag{5}$$

$$A_{\zeta}(s,\theta,\zeta) = \sum_{i,l} A_{\zeta,e,i,l} \, \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} A_{\zeta,o,i,l} \, \overline{T}_{l,i}(s) \sin \alpha_i, \tag{6}$$

where  $\overline{T}_{l,i}(s) \equiv \overline{s}^{m_i/2} T_l(s)$ ,  $T_l(s)$  is the Chebyshev polynomial, and  $\alpha_j \equiv m_j \theta - n_j \zeta$ . The regularity factor,  $\overline{s}^{m_i/2}$ , where  $\overline{s} \equiv (1+s)/2$ , is only included if there is a coordinate singularity in the domain (i.e. only in the innermost volume, and only in cylindrical and toroidal geometry.)

• The magnetic field,  $\sqrt{g}\,{f B}=\sqrt{g}B^s{f e}_s+\sqrt{g}B^\theta{f e}_\theta+\sqrt{g}B^\zeta{f e}_\zeta$ , is

$$\sqrt{g} \mathbf{B} = \mathbf{e}_{s} \sum_{i,l} \left[ (-m_{i} A_{\zeta,e,i,l} - n_{i} A_{\theta,e,i,l}) \overline{T}_{l,i} \sin \alpha_{i} + (+m_{i} A_{\zeta,o,i,l} + n_{i} A_{\theta,o,i,l}) \overline{T}_{l,i} \cos \alpha_{i} \right] 
+ \mathbf{e}_{\theta} \sum_{i,l} \left[ (-m_{i} A_{\zeta,e,i,l} - n_{i} A_{\theta,e,i,l}) \overline{T}'_{l,i} \cos \alpha_{i} + (-m_{i} A_{\zeta,o,i,l} + n_{i} A_{\theta,o,i,l}) \overline{T}'_{l,i} \sin \alpha_{i} \right] 
+ \mathbf{e}_{\zeta} \sum_{i,l} \left[ (-m_{i} A_{\zeta,e,i,l} - n_{i} A_{\theta,e,i,l}) \overline{T}'_{l,i} \cos \alpha_{i} + (-m_{i} A_{\zeta,o,i,l} + n_{i} A_{\theta,o,i,l}) \overline{T}'_{l,i} \sin \alpha_{i} \right] 
+ \mathbf{e}_{\zeta} \sum_{i,l} \left[ (-m_{i} A_{\zeta,e,i,l} - n_{i} A_{\theta,e,i,l}) \overline{T}'_{l,i} \cos \alpha_{i} + (-m_{i} A_{\zeta,o,i,l} + n_{i} A_{\theta,o,i,l}) \overline{T}'_{l,i} \sin \alpha_{i} \right]$$

· The current is

$$\sqrt{g}\,\mathbf{j} = (\partial_{\theta}B_{\zeta} - \partial_{\zeta}B_{\theta})\,\mathbf{e}_{s} + (\partial_{\zeta}B_{s} - \partial_{s}B_{\zeta})\,\mathbf{e}_{\theta} + (\partial_{s}B_{\theta} - \partial_{\theta}B_{s})\,\mathbf{e}_{\zeta},\tag{8}$$

where (for computational convenience) the covariant components of  $\boldsymbol{B}$  are computed as

$$B_s = (\sqrt{g}B^s) g_{ss} / \sqrt{g} + (\sqrt{g}B^\theta) g_{s\theta} / \sqrt{g} + (\sqrt{g}B^\zeta) g_{s\zeta} / \sqrt{g}, \tag{9}$$

$$B_{\theta} = (\sqrt{g}B^{s}) g_{s\theta} / \sqrt{g} + (\sqrt{g}B^{\theta}) g_{\theta\theta} / \sqrt{g} + (\sqrt{g}B^{\zeta}) g_{\theta\zeta} / \sqrt{g}, \tag{10}$$

$$B_{\zeta} = (\sqrt{g}B^{s}) g_{s\zeta} / \sqrt{g} + (\sqrt{g}B^{\theta}) g_{\theta\zeta} / \sqrt{g} + (\sqrt{g}B^{\zeta}) g_{\zeta\zeta} / \sqrt{g}. \tag{11}$$

# quantification of the error

· The measures of the error are

$$||(\mathbf{j} - \mu \mathbf{B}) \cdot \nabla s|| = \int ds \oint d\theta d\zeta ||\nabla g \mathbf{j} \cdot \nabla s - \mu \sqrt{g} \mathbf{B} \cdot \nabla s|, \qquad (12)$$

$$||(\mathbf{j} - \mu \mathbf{B}) \cdot \nabla \theta|| \equiv \int ds \oint d\theta d\zeta ||\sqrt{g} \mathbf{j} \cdot \nabla \theta - \mu \sqrt{g} \mathbf{B} \cdot \nabla \theta|,$$
(13)

$$||(\mathbf{j} - \mu \mathbf{B}) \cdot \nabla \zeta|| \equiv \int ds \oint d\theta d\zeta ||\sqrt{g} \,\mathbf{j} \cdot \nabla \zeta - \mu \,\sqrt{g} \,\mathbf{B} \cdot \nabla \zeta|. \tag{14}$$

#### comments

- Is there a better definition and quantification of the error? For example, should we employ an error measure that is dimensionless?
- If the coordinate singularity is in the domain, then  $|\nabla \theta| \to \infty$  at the coordinate origin. What then happens to  $||(\mathbf{j} \mu \mathbf{B}) \cdot \nabla \theta||$  as defined in Eqn. (13)?
- What is the predicted scaling of the error in the Chebyshev-Fourier representation scale with numerical resolution? Note that the predicted error scaling for  $E^s$ ,  $E^\theta$  and  $E^\zeta$  may not be standard, as various radial derivatives are taken to compute the components of  ${\bf j}$ . (See for example the discussion in Sec.IV.C in Hudson et al. (2011) [4], where the expected scaling of the error for a finite-element implementation is confirmed numerically.)
- Instead of using Gaussian integration to compute the integral over s, an adaptive quadrature algorithm may be preferable.

#### **Parameters**

in	Ivol	in which volume should the Beltrami error be computed
in $\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \$		number of grid points in $\theta$ and $\zeta$
in	in Iquad degree of Gaussian quadrature	
in <i>mn</i>		number of Fourier harmonics

#### details of the numerics

- The integration over s is performed using Gaussian integration, e.g.,  $\int f(s)ds \approx \sum_k \omega_k f(s_k)$ ; with the abscissae,  $s_k$ , and the weights,  $\omega_k$ , for k=1, Iquad v, determined by CDGQF. The resolution, N  $\equiv$  Iquad v, is determined by Nquad (see global.f90 and preset()). A fatal error is enforced by jo00aa() if CDGQF returns an ifail  $\neq 0$ .
- Inside the Gaussian quadrature loop, i.e. for each  $s_k$ ,
  - The metric elements,  $g_{\mu,\nu} \equiv \text{gij} (1:6,0,1:\text{Ntz})$ , and the Jacobian,  $\sqrt{g} \equiv \text{sg}(0,1:\text{Ntz})$ , are calculated on a regular angular grid,  $(\theta_i,\zeta_j)$ , in coords(). The derivatives  $\partial_i g_{\mu,\nu} \equiv \text{gij} (1:6,i,1\leftrightarrow i,1\leftrightarrow i,1)$  and  $\partial_i \sqrt{g} \equiv \text{sg}(i,1:\text{Ntz})$ , with respect to  $i \in \{s,\theta,\zeta\}$  are also returned.
  - The Fourier components of the vector potential given in Eqn. (5) and Eqn. (6), and their first and second radial derivatives, are summed.
  - The quantities  $\sqrt{g}B^s$ ,  $\sqrt{g}B^\theta$  and  $\sqrt{g}B^\zeta$ , and their first and second derivatives with respect to  $(s, \theta, \zeta)$ , are computed on the regular angular grid (using FFTs).
  - The following quantities are then computed on the regular angular grid

$$\sqrt{g}j^{s} = \sum_{u} \left[ \partial_{\theta}(\sqrt{g}B^{u}) g_{u,\zeta} + (\sqrt{g}B^{u}) \partial_{\theta}g_{u,\zeta} - (\sqrt{g}B^{u}) g_{u,\zeta} \partial_{\theta}\sqrt{g}/\sqrt{g} \right] / \sqrt{g}$$

$$- \sum_{u} \left[ \partial_{\zeta}(\sqrt{g}B^{u}) g_{u,\theta} + (\sqrt{g}B^{u}) \partial_{\zeta}g_{u,\theta} - (\sqrt{g}B^{u}) g_{u,\theta} \partial_{\zeta}\sqrt{g}/\sqrt{g} \right] / \sqrt{g}, \qquad (15)$$

$$\sqrt{g}j^{\theta} = \sum_{u} \left[ \partial_{\zeta}(\sqrt{g}B^{u}) g_{u,s} + (\sqrt{g}B^{u}) \partial_{\zeta}g_{u,s} - (\sqrt{g}B^{u}) g_{u,s} \partial_{\zeta}\sqrt{g}/\sqrt{g} \right] / \sqrt{g}$$

$$- \sum_{u} \left[ \partial_{s}(\sqrt{g}B^{u}) g_{u,\zeta} + (\sqrt{g}B^{u}) \partial_{s}g_{u,\zeta} - (\sqrt{g}B^{u}) g_{u,\zeta} \partial_{s}\sqrt{g}/\sqrt{g} \right] / \sqrt{g}, \qquad (16)$$

$$\sqrt{g}j^{\zeta} = \sum_{u} \left[ \partial_{s}(\sqrt{g}B^{u}) g_{u,\theta} + (\sqrt{g}B^{u}) \partial_{s}g_{u,\theta} - (\sqrt{g}B^{u}) g_{u,\theta} \partial_{s}\sqrt{g}/\sqrt{g} \right] / \sqrt{g}$$

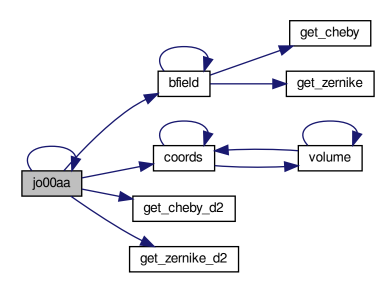
$$- \sum_{u} \left[ \partial_{\theta}(\sqrt{g}B^{u}) g_{u,s} + (\sqrt{g}B^{u}) \partial_{\theta}g_{u,s} - (\sqrt{g}B^{u}) g_{u,s} \partial_{\theta}\sqrt{g}/\sqrt{g} \right] / \sqrt{g}. \qquad (17)$$

• The error is stored into an array called beltramierror which is then written to the HDF5 file in hdfint().

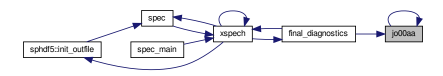
References allglobal::ate, allglobal::ato, allglobal::aze, allglobal::aze, allglobal::beltramierror, bfield(), allglobal::cfmn, allglobal::cheby, coords(), allglobal::cpus, allglobal::dpflux, allglobal::dtflux, allglobal::efmn, allglobal::gbzeta, get\_cheby\_d2(), get\_zernike\_d2(), allglobal::guvij, constants::half, inputlist::igeometry, allglobal::im, allglobal::im, allglobal::in, allglobal::ivol, jo00aa(), allglobal::lcoordinatesingularity, inputlist::lerrortype, inputlist::lrad, allglobal::mpi\_comm\_spec, inputlist::mpol, inputlist::mu, allglobal::myid, inputlist::nfp, allglobal::node, allglobal::notstellsym, allglobal::nt, inputlist::nvol, allglobal::nz, allglobal::ofmn, constants::one, fileunits::ounit, constants::pi2, allglobal::regumm, allglobal::rij, allglobal::rtt, allglobal::sfmn, allglobal::sg, allglobal::tt, constants::two, inputlist::wmacros, allglobal::zernike, constants::zero, and allglobal::zij.

Referenced by final diagnostics(), and jo00aa().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.1.2.4 pp00aa() subroutine pp00aa

Constructs Poincaré plot and "approximate" rotational-transform (driver).

# relevant input variables

- · The resolution of Poincaré plot is controlled by
  - nPtraj trajectories will be located in each volume;
  - nPpts iterations per trajectory;
  - odetol o.d.e. integration tolerance;

- The magnetic field is given by bfield() .
- The approximate rotational transform is determined, in pp00ab(), by fieldline integration.

#### format of output: Poincaré

 The Poincaré data is written to .ext.poincare:xxxx , where xxxx is an integer indicating the volume. The format of this file is as follows:

```
write (svol,'(i4.4)') lvol! lvol labels volume; open (lunit+myid, file="."//trim(ext)//".poincare."//svol, status="unknown", form="unformatted") do until end of file write (lunit+myid) nz, nppts ! integers write (lunit+myid) data (1:4,0:nz-1,1:nppts)! doubles enddo close (lunit+myid) where  -\theta \equiv \text{data}(1,k,j) \text{ is the poloidal angle,} \\ -s \equiv \text{data}(2,k,j) \text{ is the radial coordinate,} \\ -R \equiv \text{data}(3,k,j) \text{ is the cylindrical } R, \\ -Z \equiv \text{data}(4,k,j) \text{ is the cylindrical } Z,
```

- The integer k=0,Nz-1 labels toroidal planes, so that  $\phi = (2\pi/\text{Nfp})(k/\text{Nz})$ ,
- The integer j=1,nPpts labels toroidal iterations.
- Usually (if no fieldline integration errors are encountered) the number of fieldlines followed in volume lvol is given by N+1, where the radial resolution,  $N \equiv Ni \ (lvol)$ , is given on input. This will be over-ruled by if  $nPtrj \ (lvol)$ , given on input, is non-negative.
- The starting location for the fieldline integrations are equally spaced in the radial coordinate  $s_i = s_{l-1} + i(s_l s_{l-1})/N$  for i = 0, N, along the line  $\theta = 0, \zeta = 0$ .

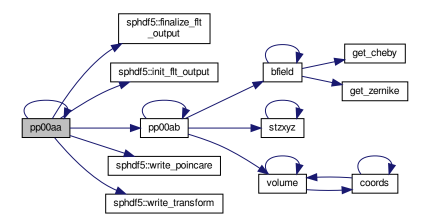
# format of output: rotational-transform

• The rotational-transform data is written to .ext.transform:xxxx , where xxxx is an integer indicating the volume. The format of this file is as follows:

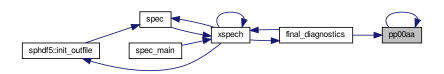
References allglobal::cpus, allglobal::diotadxup, sphdf5::finalize\_flt\_output(), constants::half, inputlist::igeometry, sphdf5::init\_flt\_output(), inputlist::iota, allglobal::ivol, inputlist::lconstraint, allglobal::lcoordinatesingularity, allglobal::lplasmaregion, inputlist::lrad, allglobal::lvacuumregion, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ncpu, inputlist::nppts, inputlist::nptrj, inputlist::nvol, allglobal::nz, inputlist::odetol, inputlist::oita, constants::one, fileunits::ounit, constants::pi, pp00aa(), pp00ab(), inputlist::ppts, constants::two, inputlist::wmacros, sphdf5::write\_poincare(), sphdf5::write\_transform(), and constants::zero.

Referenced by final diagnostics(), and pp00aa().

Here is the call graph for this function:



Here is the caller graph for this function:



```
7.1.2.5 pp00ab() subroutine pp00ab (
    integer, intent(in) lvol,
    real, dimension(1:2) sti,
    integer, intent(in) Nz,
    integer, intent(in) nPpts,
    real, dimension(1:4,0:nz-1,1:nppts) poincaredata,
    real, dimension(1:2) fittedtransform,
    integer, intent(out) utflag)
```

Constructs Poincaré plot and "approximate" rotational-transform (for single field line). **relevant input variables** 

- · The resolution of Poincaré plot is controlled by
  - nPpts iterations per trajectory;
  - odetol o.d.e. integration tolerance;

The magnetic field is given by bfield().

#### rotational-transform

• The approximate rotational transform is determined by field line integration. This is constructed by fitting a least squares fit to the field line trajectory.

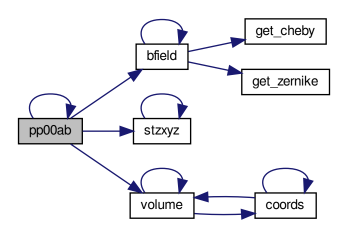
#### **Parameters**

in	Ivol	
	sti	
in	Nz	
in	nPpts	
	poincaredata	
	fittedtransform	
out	utflag	

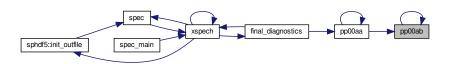
References bfield(), allglobal::cpus, allglobal::ivol, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ncpu, allglobal::node, inputlist::nvol, inputlist::odetol, constants::one, fileunits::ounit, constants::pi2, pp00ab(), numerical::small, stzxyz(), constants::two, volume(), and constants::zero.

Referenced by pp00aa(), and pp00ab().

Here is the call graph for this function:



Here is the caller graph for this function:



• This routine is a "copy" of co01aa(), which calculates the coordinate information on a regular, discrete grid in  $\theta$  and  $\zeta$  at given s whereas stzxyz() calculates the coordinate information at a single point  $(s, \theta, \zeta)$ .

• Todo Please see co01aa() for documentation.

#### **Parameters**

in	Ivol	
in	stz	
out	RpZ	

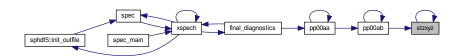
References allglobal::cpus, constants::half, allglobal::halfmm, inputlist::igeometry, allglobal::im, allglobal::in, allglobal::irbc, allglobal::irbc, allglobal::izbc, allglobal::izbc, allglobal::izbc, allglobal::notstellsym, inputlist::ntor, inputlist::nvol, constants::one, fileunits::ounit, stzxyz(), numerical::vsmall, and constants::zero.

Referenced by pp00ab(), and stzxyz().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.2 Free-Boundary Computation

# **Functions/Subroutines**

- subroutine bnorml (mn, Ntz, efmn, ofmn)
  - Computes  $\mathbf{B}_{Plasma} \cdot \mathbf{e}_{\theta} \times \mathbf{e}_{\zeta}$  on the computational boundary,  $\partial \mathcal{D}$ .
- subroutine casing (teta, zeta, gBn, icasing)

Constructs the field created by the plasma currents, at an arbitrary, external location using virtual casing.

• subroutine dvcfield (Ndim, tz, Nfun, vcintegrand)

Differential virtual casing integrand.

### 7.2.1 Detailed Description

#### 7.2.2 Function/Subroutine Documentation

```
real, dimension(1:mn), intent(out) efmn,
real, dimension(1:mn), intent(out) ofmn)
```

Computes  $\mathbf{B}_{Plasma} \cdot \mathbf{e}_{\theta} \times \mathbf{e}_{\zeta}$  on the computational boundary,  $\partial \mathcal{D}$ .

#### free-boundary constraint

- The normal field at the computational boundary,  $\partial \mathcal{D}$ , should be equal to  $(\mathbf{B}_P + \mathbf{B}_C) \cdot \mathbf{e}_{\theta} \times \mathbf{e}_{\zeta}$ , where  $\mathbf{B}_P$  is the "plasma" field (produced by internal plasma currents) and is computed using virtual casing, and  $\mathbf{B}_C$  is the "vacuum" field (produced by the external coils) and is given on input.
- The plasma field,  $\mathbf{B}_P$ , can only be computed after the equilibrium is determined, but this information is required to compute the equilibrium to begin with; and so there is an iteration involved.
- Suggested values of the vacuum field can be self generated; see xspech() for more documentation on this.

# compute the normal field on a regular grid on the computational boundary

- For each point on the computational boundary, casing() is called to compute the normal field produced by the plasma currents.
- Todo There is a very clumsy attempt to parallelize this which could be greatly improved.
- · An FFT gives the required Fourier harmonics.

#### See also

# casing.f90

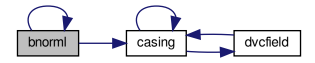
#### **Parameters**

in	mn	total number of Fourier harmonics
in	Ntz	total number of grid points in $\boldsymbol{\theta}$ and $zeta$
out	efmn	even Fourier coefficients
out	ofmn	odd Fouier coefficients

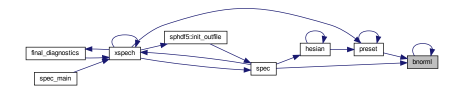
References allglobal::ate, allglobal::ato, allglobal::aze, allglobal::azo, bnorml(), casing(), allglobal::cfmn, allglobal::cpus, allglobal::dxyz, allglobal::globaljk, allglobal::gteta, allglobal::guvij, allglobal::gzeta, constants::half, inputlist::igeometry, allglobal::jiimag, allglobal::jiireal, allglobal::im, allglobal::jiimag, allglobal::jiimag, allglobal::jiimag, allglobal::jiimag, allglobal::jiimag, allglobal::jiimag, allglobal::myid, allglobal::myid, allglobal::ncpu, allglobal::notstellsym, allglobal::nt, allglobal::nxyz, allglobal::nz, constants::one, fileunits::ounit, constants::pi, constants::pi2, allglobal::rij, allglobal::sfmn, allglobal::sg, numerical::small, constants::ten, allglobal::tetazeta, allglobal::tt, constants::two, inputlist::vcasingper, inputlist::vcasingtol, allglobal::virtualcasingfactor, inputlist::wmacros, constants::zero, and allglobal::zij.

Referenced by bnorml(), preset(), and spec().

Here is the call graph for this function:



Here is the caller graph for this function:



Constructs the field created by the plasma currents, at an arbitrary, external location using virtual casing. Compute the external magnetic field using virtual casing.

### Theory and numerics

· Required inputs to this subroutine are the geometry of the plasma boundary,

$$\mathbf{x}(\theta,\zeta) \equiv x(\theta,\zeta)\mathbf{i} + y(\theta,\zeta)\mathbf{j} + z(\theta,\zeta)\mathbf{k},\tag{18}$$

and the tangential field on this boundary,

$$\mathbf{B}_{s} = B^{\theta} \mathbf{e}_{\theta} + B^{\zeta} \mathbf{e}_{\zeta},\tag{19}$$

where  $\theta$  and  $\zeta$  are arbitrary poloidal and toroidal angles, and  $\mathbf{e}_{\theta} \equiv \partial \mathbf{x}/\partial \theta$ ,  $\mathbf{e}_{\zeta} \equiv \partial \mathbf{x}/\partial \zeta$ . This routine assumes that the plasma boundary is a flux surface, i.e.  $\mathbf{B} \cdot \mathbf{e}_{\theta} \times \mathbf{e}_{\zeta} = 0$ .

• The virtual casing principle (Shafranov & Zakharov (1972) [7], Lazerson (2012) [5] and Hanson (2015) [1]) shows that the field outside/inside the plasma arising from plasma currents inside/outside the boundary is equivalent to the field generated by a surface current,

$$\mathbf{j} = \mathbf{B}_s \times \mathbf{n},\tag{20}$$

where n is normal to the surface.

- The field at some arbitrary point,  $\bar{\mathbf{x}},$  created by this surface current is given by

$$\mathbf{B}(\bar{\mathbf{x}}) = -\frac{1}{4\pi} \int_{\mathcal{S}} \frac{(\mathbf{B}_s \times d\mathbf{s}) \times \hat{\mathbf{r}}}{r^2},\tag{21}$$

where  $d\mathbf{s} \equiv \mathbf{e}_{\theta} \times \mathbf{e}_{\zeta} \ d\theta d\zeta$ .

· For ease of notation introduce

$$\mathbf{J} \equiv \mathbf{B}_s \times d\mathbf{s} = \alpha \, \mathbf{e}_{\theta} - \beta \, \mathbf{e}_{\zeta}, \tag{22}$$

where  $\alpha \equiv B_{\zeta} = B^{\theta}g_{\theta\zeta} + B^{\zeta}g_{\zeta\zeta}$  and  $\beta \equiv B_{\theta} = B^{\theta}g_{\theta\theta} + B^{\zeta}g_{\theta\zeta}$ .

• We may write in Cartesian coordinates  ${f J}=j_x\ {f i}+j_y\ {f j}+j_z\ {f k},$  where

$$j_x = \alpha x_\theta - \beta x_\zeta \tag{23}$$

$$j_y = \alpha y_\theta - \beta y_\zeta \tag{24}$$

$$j_z = \alpha z_\theta - \beta z_\zeta. \tag{25}$$

· Requiring that the current,

$$\mathbf{j} \equiv \nabla \times \mathbf{B} = \sqrt{g}^{-1} (\partial_{\theta} B_{\zeta} - \partial_{\zeta} B_{\theta}) \mathbf{e}_{s} + \sqrt{g}^{-1} (\partial_{\zeta} B_{s} - \partial_{s} B_{\zeta}) \mathbf{e}_{\theta} + \sqrt{g}^{-1} (\partial_{s} B_{\theta} - \partial_{\theta} B_{s}) \mathbf{e}_{\zeta}$$
(26)

has no normal component to the surface, i.e.  $\mathbf{j} \cdot \nabla s = 0$ , we obtain the condition  $\partial_{\theta} B_{\zeta} = \partial_{\zeta} B_{\theta}$ , or  $\partial_{\theta} \alpha = \partial_{\zeta} \beta$ . In axisymmetric configurations, where  $\partial_{\zeta} \beta = 0$ , we must have  $\partial_{\theta} \alpha = 0$ .

• The displacement from an arbitrary point, (X, Y, Z), to a point, (x, y, z), that lies on the surface is given

$$\mathbf{r} \equiv r_x \,\mathbf{i} + r_y \,\mathbf{j} + r_z \,\mathbf{k} = (X - x) \,\mathbf{i} + (Y - y) \,\mathbf{j} + (Z - z) \,\mathbf{k}. \tag{27}$$

· The components of the magnetic field produced by the surface current are then

$$B^{x} = \oint \!\! \oint \! d\theta d\zeta \ (j_{y}r_{z} - j_{z}r_{y})/r^{3}, \tag{28}$$

$$B^{y} = \oint \!\! \oint \! d\theta d\zeta \ (j_{z}r_{x} - j_{x}r_{z})/r^{3}, \tag{29}$$

$$B^{z} = \oint \!\! \oint \! d\theta d\zeta \ (j_{x}r_{y} - j_{y}r_{x})/r^{3} \tag{30}$$

up to a scaling factor virtual casing factor  $=-1/4\pi$  that is taken into account at the end.

· When all is said and done, this routine calculates

$$\int_{0}^{2\pi} \int_{0}^{2\pi} \text{vcintegrand } d\theta d\zeta \tag{31}$$

for a given (X, Y, Z), where vcintegrand is given in Eqn. (33).

The surface integral is performed using DCUHRE, which uses an adaptive subdivision strategy and also computes absolute error estimates. The absolute and relative accuracy required are provided by the inputvar vcasingtol. The minimum number of function evaluations is provided by the inputvar vcasingits.

#### Calculation of integrand

• An adaptive integration is used to compute the integrals. Consequently, the magnetic field tangential to the plasma boundary is required at an arbitrary point. This is computed, as always, from  $\mathbf{B} = \nabla \times \mathbf{A}$ , and this provides  $\mathbf{B} = B^{\theta} \mathbf{e}_{\theta} + B^{\zeta} \mathbf{e}_{\zeta}$ . Recall that  $B^{s} = 0$  by construction on the plasma boundary.

Todo It would be MUCH faster to only require the tangential field on a regular grid!!!

• Then, the metric elements  $g_{\theta\theta}$ ,  $g_{\theta\zeta}$  and  $g_{\zeta\zeta}$  are computed. These are used to "lower" the components of the magnetic field,  $\mathbf{B} = B_{\theta} \nabla \theta + B_{\zeta} \nabla \zeta$ .

**Todo** Please check why  $B_s$  is not computed. Is it because  $B_s \nabla s \times \mathbf{n} = 0$ ?

- The distance between the "evaluate" point, (X,Y,Z), and the given point on the surface, (x,y,z) is computed.
- If the computational boundary becomes too close to the plasma boundary, the distance is small and this causes problems for the numerics. I have tried to regularize this problem by introducing  $\epsilon$  =inputvar vcasingeps. Let the "distance" be

$$D \equiv \sqrt{(X-x)^2 + (Y-y)^2 + (Z-Z)^2} + \epsilon^2.$$
 (32)

• On taking the limit that  $\epsilon \to 0$ , the virtual casing integrand is

vcintegrand 
$$\equiv (B_x n_x + B_y n_y + B_z n_z)(1 + 3\epsilon^2/D^2)/D^3$$
, (33)

where the normal vector is  $\mathbf{n} \equiv n_x \mathbf{i} + n_y \mathbf{j} + n_z \mathbf{k}$ . The normal vector, Nxyz, to the computational boundary (which does not change) is computed in preset().

Todo This needs to be revised.

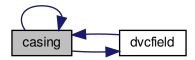
#### **Parameters**

in	teta	$\theta$
in	zeta	ζ
out	gBn	$\sqrt{g}\mathbf{B}\cdot\mathbf{n}$
out	icasing	return flag from dcuhre()

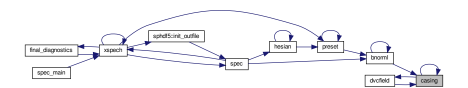
References casing(), allglobal::cpus, dvcfield(), allglobal::dxyz, allglobal::globaljk, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ncpu, allglobal::nxyz, fileunits::ounit, constants::pi, constants::pi2, inputlist::vcasingits, inputlist::vcasingper, inputlist::vcasingtol, fileunits::vunit, inputlist::wmacros, and constants::zero.

Referenced by bnorml(), casing(), and dvcfield().

Here is the call graph for this function:



Here is the caller graph for this function:



Differential virtual casing integrand.

Differential virtual casing integrand

### **Parameters**

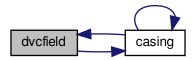
_			
	in	Ndim	number of parameters (==2)
	in	tz	$ heta$ and $\zeta$
	in	Nfun	number of function values (==3)
	out	vcintegrand	cartesian components of magnetic field

References allglobal::ate, allglobal::ate, allglobal::aze, allglobal::aze, allglobal::aze, casing(), allglobal::cpus, allglobal::dxyz, allglobal::first\_free\_bound, constants::four, allglobal::global;, constants::half, inputlist::igeometry, allglobal::im,

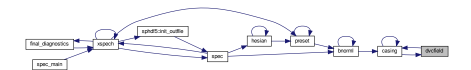
allglobal::in, allglobal::irbc, allglobal::irbc, allglobal::izbc, allglobal::izbc, inputlist::lrad, allglobal::mn, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ncpu, allglobal::notstellsym, inputlist::nvol, allglobal::nxyz, constants::one, fileunits::ounit, numerical::small, constants::three, allglobal::tt, inputlist::vcasingeps, fileunits::vunit, allglobal::yesstellsym, and constants::zero.

Referenced by casing().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.3 Parallelization

## **Functions/Subroutines**

• subroutine breast (Ivol)

Broadcasts Beltrami fields, profiles, . . .

# 7.3.1 Detailed Description

# 7.3.2 Function/Subroutine Documentation

# 

# broadcasting

- The construction of the Beltrami fields is distributed on separate cpus.
- All "local" information needs to be broadcast so that the "global" force vector,

$$\mathbf{F}_i \equiv [[p + B^2/2]]_i = (p + B^2/2)_{v,i} - (p + B^2/2)_{v-1,i}$$
(34)

can be constructed, and so that restart and output files can be saved to file.

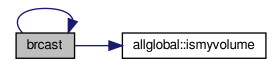
# Parameters

in	Ivol	index of nested volume

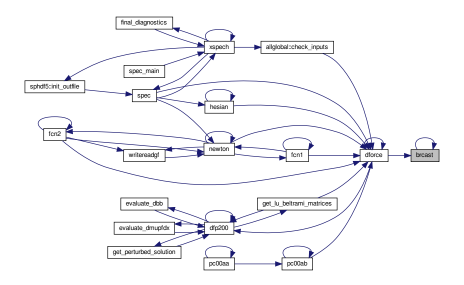
7.4 Geometry 25

References allglobal::ate, allglobal::ato, allglobal::aze, allglobal::azo, allglobal::bemn, allglobal::bemn, brcast(), allglobal::cpus, inputlist::curpol, inputlist::curtor, allglobal::dbbdmp, allglobal::dffdrz, allglobal::diotadxup, allglobal::ditgpdxtp, allglobal::dmupfdx, allglobal::dpflux, allglobal::dtflux, inputlist::helicity, allglobal::iemn, allglobal::imagneticok, allglobal::iomn, allglobal::ismyvolume(), allglobal::ismyvolumevalue, allglobal::labintegral, allglobal::lbbintegral, inputlist::lconstraint, inputlist::lfindzero, allglobal::lgdof, allglobal::lhessianallocated, allglobal::localconstraint, inputlist::lrad, allglobal::mn, inputlist::mnvol, allglobal::mpi\_comm\_spec, inputlist::mu, allglobal::myid, allglobal::notstellsym, allglobal::ntz, inputlist::nvol, fileunits::ounit, allglobal::pemn, allglobal::pomn, allglobal::semn, allglobal::somn, allglobal::vvolume, inputlist::wmacros, and constants::zero. Referenced by brcast(), and dforce().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.4 Geometry

### **Functions/Subroutines**

• subroutine coords (Ivol, Iss, Lcurvature, Ntz, mn) Calculates coordinates,  $\mathbf{x}(s,\theta,\zeta) \equiv R \, \mathbf{e}_R + Z \, \mathbf{e}_Z$ , and metrics, using FFTs.

#### 7.4.1 Detailed Description

# 7.4.2 Function/Subroutine Documentation

integer, intent(in) mn )

Calculates coordinates,  $\mathbf{x}(s, \theta, \zeta) \equiv R \mathbf{e}_R + Z \mathbf{e}_Z$ , and metrics, using FFTs.

#### Coordinates

- We work in coordinates,  $(s, \theta, \zeta)$ , which are be defined *inversely* via a transformation *to* Cartesian coordinates, (x, y, z).
- The toroidal angle,  $\zeta$ , is identical to the cylindrical angle,  $\zeta \equiv \phi$ .
- The radial coordinate, s, is *not* a global variable: it only needs to be defined in each volume, and in each volume  $s \in [-1, 1]$ .
- The choice of poloidal angle,  $\theta$ , does not affect the following.

#### Geometry

- The geometry of the "ideal"-interfaces,  $\mathbf{x}_v(\theta,\zeta)$ , is given by  $R(\theta,\zeta)$  and  $Z(\theta,\zeta)$  as follows:
  - Igeometry=1: Cartesian

$$\mathbf{x} \equiv r_{pol}\theta \,\,\hat{\mathbf{i}} + r_{tor}\zeta \,\,\hat{\mathbf{j}} + R \,\,\hat{\mathbf{k}} \tag{35}$$

where  $r_{pol}$  and  $r_{tor}$  are inputs and  $r_{pol} = r_{tor} = 1$  by default.

- Igeometry=2: Cylindrical

$$\mathbf{x} = R \cos \theta \,\hat{\mathbf{i}} + R \sin \theta \,\hat{\mathbf{j}} + \zeta \,\hat{\mathbf{k}} \tag{36}$$

- Igeometry=3: Toroidal

$$\mathbf{x} \equiv R \,\hat{\mathbf{r}} + Z \,\hat{\mathbf{k}} \tag{37}$$

where  $\hat{\mathbf{r}} \equiv \cos \phi \, \hat{\mathbf{i}} + \sin \phi \, \hat{\mathbf{j}}$  and  $\hat{\phi} \equiv -\sin \phi \, \hat{\mathbf{i}} + \cos \phi \, \hat{\mathbf{j}}$ .

• The geometry of the ideal interfaces is given as Fourier summation: e.g., for stellarator-symmetry

$$R_v(\theta,\zeta) \equiv \sum_j R_{j,v} \cos \alpha_j,$$
 (38)

$$Z_v(\theta,\zeta) \equiv \sum_j Z_{j,v} \sin \alpha_j,$$
 (39)

where  $\alpha_i \equiv m_i \theta - n_i \zeta$ .

### interpolation between interfaces

- The "coordinate" functions,  $R(s,\theta,\zeta)$  and  $Z(s,\theta,\zeta)$ , are constructed by radially interpolating the Fourier representations of the ideal-interfaces.
- The v-th volume is bounded by  $\mathbf{x}_{v-1}$  and  $\mathbf{x}_v$ .
- In each *annular* volume, the coordinates are constructed by linear interpolation:

$$R(s,\theta,\zeta) \equiv \sum_{j} \left[ \frac{(1-s)}{2} R_{j,v-1} + \frac{(1+s)}{2} R_{j,v} \right] \cos \alpha_{j},$$

$$Z(s,\theta,\zeta) \equiv \sum_{j} \left[ \frac{(1-s)}{2} Z_{j,v-1} + \frac{(1+s)}{2} Z_{j,v} \right] \sin \alpha_{j},$$

$$(40)$$

coordinate singularity: regularized extrapolation

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• For cylindrical or toroidal geometry, in the innermost, "simple-torus" volume, the coordinates are constructed by an interpolation that "encourages" the interpolated coordinate surfaces to not intersect.

• Introduce  $\bar{s} \equiv (s+1)/2$ , so that in each volume  $\bar{s} \in [0,1]$ , then

$$R_j(s) = R_{j,0} + (R_{j,1} - R_{j,0})f_j,$$
 (41)

$$Z_i(s) = Z_{i,0} + (Z_{i,1} - Z_{i,0})f_i,$$
 (42)

where, in toroidal geometry,

$$f_j \equiv \left\{ \begin{array}{ll} \bar{s} & , & \text{for } m_j = 0, \\ \bar{s}^{m_j} & , & \text{otherwise.} \end{array} \right\}. \tag{43}$$

• Note: The location of the coordinate axis, i.e. the  $R_{j,0}$  and  $Z_{j,0}$ , is set in the coordinate "packing" and "unpacking" routine, packxi().

#### Jacobian

- · The coordinate Jacobian (and some other metric information) is given by
  - Igeometry=1: Cartesian

$$\mathbf{e}_{\theta} \times \mathbf{e}_{\zeta} = -r_{tor} R_{\theta} \,\, \hat{\mathbf{i}} - r_{pol} R_{\zeta} \,\, \hat{\mathbf{j}} + r_{pol} r_{tor} \hat{\mathbf{k}}$$
(44)

$$\boldsymbol{\xi} \cdot \mathbf{e}_{\theta} \times \mathbf{e}_{\zeta} = \delta R \tag{45}$$

$$\sqrt{g} = R_s \, r_{pol} \, r_{tor} \tag{46}$$

- Igeometry=2: Cylindrical

$$\mathbf{e}_{\theta} \times \mathbf{e}_{\zeta} = (R_{\theta} \sin \theta + R \cos \theta) \, \hat{\mathbf{i}} + (R \sin \theta - R_{\theta} \cos \theta) \, \hat{\mathbf{j}} - RR_{\zeta} \, \hat{\mathbf{k}}$$
(47)

$$\boldsymbol{\xi} \cdot \mathbf{e}_{\theta} \times \mathbf{e}_{\zeta} = \delta R R \tag{48}$$

$$\sqrt{g} = R_s R \tag{49}$$

- Igeometry=3: Toroidal

$$\mathbf{e}_{\theta} \times \mathbf{e}_{\zeta} = -R Z_{\theta} \,\hat{r} + (Z_{\theta} R_{\zeta} - R_{\theta} Z_{\zeta}) \hat{\phi} + R R_{\theta} \,\hat{z} \tag{50}$$

$$\boldsymbol{\xi} \cdot \mathbf{e}_{\theta} \times \mathbf{e}_{\zeta} = R(\delta Z R_{\theta} - \delta R Z_{\theta}) \tag{51}$$

$$\sqrt{g} = R(Z_s R_\theta - R_s Z_\theta) \tag{52}$$

# cartesian metrics

· The cartesian metrics are

$$g_{ss}=R_sR_s, \quad g_{s\theta}=R_sR_{\theta}, \quad g_{s\zeta}=R_sR_{\zeta}, \quad g_{\theta\theta}=R_{\theta}R_{\theta}+r_{nol}^2, \quad g_{\theta\zeta}=R_{\theta}R_{\zeta}, \quad g_{\zeta\zeta}=R_{\zeta}R_{\zeta}+r_{tor}^2$$
 (53)

#### cylindrical metrics

· The cylindrical metrics are

$$g_{ss} = R_s R_s$$
,  $g_{s\theta} = R_s R_{\theta}$ ,  $g_{s\zeta} = R_s R_{\zeta}$ ,  $g_{\theta\theta} = R_{\theta} R_{\theta} + R^2$ ,  $g_{\theta\zeta} = R_{\theta} R_{\zeta}$ ,  $g_{\zeta\zeta} = R_{\zeta} R_{\zeta} + 1$  (54)

# logical control

- The logical control is provided by Lcurvature as follows:
  - Lcurvature=0 : only the coordinate transformation is computed, i.e. only R and Z are calculated, e.g. global()
  - Lcurvature=1 : the Jacobian,  $\sqrt{g}$ , and "lower" metrics,  $g_{\mu,\nu}$ , are calculated, e.g. bnorml(), lforce(), curent(), metrix(), sc00aa()
  - Lcurvature=2: the "curvature" terms are calculated, by which I mean the second derivatives of the position vector; this information is required for computing the current,  $\mathbf{j} = \nabla \times \nabla \times \mathbf{A}$ , e.g. jo00aa()
  - Lcurvature=3 : the derivative of the  $g_{\mu,\nu}/\sqrt{g}$  w.r.t. the interface boundary geometry is calculated, e.g. metrix(), curent()
  - Lcurvature=4: the derivative of the  $g_{\mu,\nu}$  w.r.t. the interface boundary geometry is calculated, e.g. dforce()
  - Lcurvature=5 : the derivative of  $\sqrt{g}$  w.r.t. the interface boundary geometry is calculated, e.g. rzaxis()

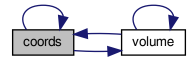
#### **Parameters**

in	Ivol	specified in which volume to compute coordinates
in	lss	radial coordinate $s$
in	Lcurvature	logical control flag
in	Ntz	number of points in $\theta$ and $\zeta$
in	mn	number of Fourier harmonics

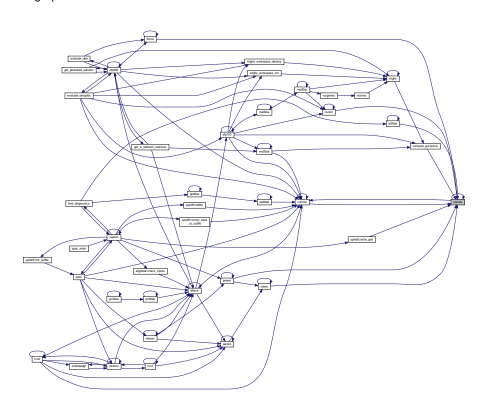
References coords(), allglobal::cosi, allglobal::cpus, allglobal::dbdx, allglobal::drodr, allglobal::drodz, allglobal::dzodr, allglobal::dzodz, allglobal::dzodz, allglobal::dzodz, allglobal::ipuij, constants::half, allglobal::halfmm, inputlist::igeometry, allglobal::im, allglobal::im, allglobal::irbs, allglobal::irbs, allglobal::izbs, allglobal::lcoordinatesingularity, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::notstellsym, allglobal::nt, inputlist::ntor, allglobal::nz, constants::one, fileunits::ounit, constants::pi2, allglobal::rji, inputlist::rpol, inputlist::rtor, allglobal::sg, allglobal::sini, numerical::small, constants::two, volume(), numerical::vsmall, inputlist::zbc, inputlist::zbs, constants::zero, and allglobal::zij.

Referenced by compute\_guvijsave(), coords(), curent(), jo00aa(), lforce(), preset(), rzaxis(), volume(), and sphdf5::write\_grid().

Here is the call graph for this function:



Here is the caller graph for this function:



7.5 Plasma Currents 29

## 7.5 Plasma Currents

#### **Functions/Subroutines**

• subroutine curent (Ivol, mn, Nt, Nz, iflag, IdItGp)

Computes the plasma current,  $I \equiv \int B_{\theta} d\theta$ , and the "linking" current,  $G \equiv \int B_{\zeta} d\zeta$ .

#### 7.5.1 Detailed Description

#### 7.5.2 Function/Subroutine Documentation

```
7.5.2.1 curent() subroutine curent (
    integer, intent(in) lvol,
    integer, intent(in) mn,
    integer, intent(in) Nt,
    integer, intent(in) Nz,
    integer, intent(in) iflag,
    real, dimension(0:1,-1:2), intent(out) ldItGp )
```

Computes the plasma current,  $I \equiv \int B_{\theta} d\theta$ , and the "linking" current,  $G \equiv \int B_{\zeta} d\zeta$ .

#### enclosed currents

In the vacuum region, the enclosed currents are given by either surface integrals of the current density or line
integrals of the magnetic field,

$$\int_{\mathcal{S}} \mathbf{j} \cdot d\mathbf{s} = \int_{\partial \mathcal{S}} \mathbf{B} \cdot d\mathbf{l},\tag{55}$$

and line integrals are usually easier to compute than surface integrals.

- The magnetic field is given by the curl of the magnetic vector potential, as described in e.g. bfield().
- The toroidal, plasma current is obtained by taking a "poloidal" loop,  $d\mathbf{l}=\mathbf{e}_{\theta}\,d\theta$ , on the plasma boundary, where  $B^s=0$ , to obtain

$$I \equiv \int_0^{2\pi} \mathbf{B} \cdot \mathbf{e}_{\theta} \, d\theta = \int_0^{2\pi} (-\partial_s A_{\zeta} \, \bar{g}_{\theta\theta} + \partial_s A_{\theta} \, \bar{g}_{\theta\zeta}) \, d\theta, \tag{56}$$

where  $\bar{g}_{\mu\nu} \equiv g_{\mu\nu}/\sqrt{g}$ .

• The poloidal, "linking" current through the torus is obtained by taking a "toroidal" loop,  $d\mathbf{l}=\mathbf{e}_{\zeta}\,d\zeta$ , on the plasma boundary to obtain

$$G \equiv \int_{0}^{2\pi} \mathbf{B} \cdot \mathbf{e}_{\zeta} \, d\zeta = \int_{0}^{2\pi} \left( -\partial_{s} A_{\zeta} \, \bar{g}_{\theta\zeta} + \partial_{s} A_{\theta} \, \bar{g}_{\zeta\zeta} \right) \, d\zeta. \tag{57}$$

#### Fourier integration

• Using  $f\equiv -\partial_s A_\zeta \; \bar{g}_{\theta\theta} + \partial_s A_\theta \; \bar{g}_{\theta\zeta}$ , the integral for the plasma current is

$$I = \sum_{i}' f_i \cos(n_i \zeta) 2\pi, \tag{58}$$

where  $\sum_{i=1}^{n} f(x_i)$  includes only the  $m_i = 0$  harmonics.

• Using  $g\equiv -\partial_s A_\zeta \; \bar{g}_{\theta\zeta} + \partial_s A_\theta \; \bar{g}_{\zeta\zeta}$ , the integral for the linking current is

$$G = \sum_{i}' g_i \cos(m_i \zeta) 2\pi, \tag{59}$$

where  $\sum'$  includes only the  $n_i=0$  harmonics.

• The plasma current, Eqn. (58), should be independent of  $\zeta$ , and the linking current, Eqn. (59), should be independent of  $\theta$ .

Todo Perhaps this can be proved analytically; in any case it should be confirmed numerically.

#### **Parameters**

in	Ivol	index of volume	
in	mn number of Fourier harmonics		
in	$Nt$ number of grid points along $\theta$		
in	$Nz$ number of grid points along $\zeta$		
in	iflag	iflag some integer flag	
out	ldltGp	plasma and linking current	

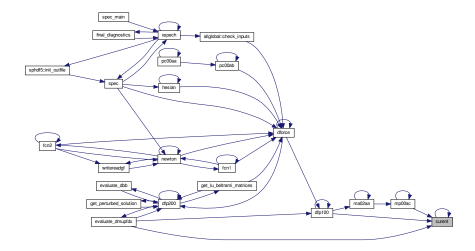
References allglobal::ate, allglobal::ato, allglobal::aze, allglobal::azo, allglobal::cfmn, allglobal::comn, coords(), allglobal::cpus, curent(), allglobal::efmn, allglobal::evmn, allglobal::guvij, allglobal::jiimag, allglobal::jiimag, allglobal::jiimag, allglobal::jiimag, allglobal::jiimag, allglobal::mag, allglobal::ofmn, allglobal::ofmn, constants::one, fileunits::ounit, constants::pi2, allglobal::sfmn, allglobal::sg, allglobal::simn, allglobal::tt, constants::two, inputlist::wmacros, allglobal::yesstellsym, and constants::zero.

Referenced by curent(), dfp100(), evaluate\_dmupfdx(), and mp00ac().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.6 "global" force

## **Functions/Subroutines**

• subroutine dforce (NGdof, position, force, LComputeDerivatives, LComputeAxis) Calculates  $\mathbf{F}(\mathbf{x})$ , where  $\mathbf{x} \equiv \{geometry\} \equiv \{R_{i,v}, Z_{i,v}\}$  and  $\mathbf{F} \equiv [[p+B^2/2]] + \{spectral \ constraints\}$ , and  $\nabla \mathbf{F}$ . 7.6 "global" force 31

## 7.6.1 Detailed Description

### 7.6.2 Function/Subroutine Documentation

Calculates  $\mathbf{F}(\mathbf{x})$ , where  $\mathbf{x} \equiv \{\text{geometry}\} \equiv \{R_{i,v}, Z_{i,v}\}$  and  $\mathbf{F} \equiv [[p+B^2/2]] + \{\text{spectral constraints}\}$ , and  $\nabla \mathbf{F}$ . **unpacking** 

• The geometrical degrees of freedom are represented as a vector,  $\mathbf{x} \equiv \{R_{i,v}, Z_{i,v}\}$ , where  $i=1, \, \text{mn}$  labels the Fourier harmonic and  $v=1, \, \text{Mvol} \, -1$  is the interface label. This vector is "unpacked" using packxi(). (Note that packxi() also sets the coordinate axis, i.e. the  $R_{i,0}$  and  $Z_{i,0}$ .)

### **Matrices computation**

- the volume-integrated metric arrays, DToocc, etc. are evaluated in each volume by calling ma00aa()
- the energy and helicity matrices, dMA (0:NN, 0:NN), dMB (0:NN, 0:2), etc. are evaluated in each volume by calling matrix()

#### parallelization over volumes

Two different cases emerge: either a local constraint or a global constraint is considered. This condition is determined by the flag LocalConstraint.

- · Local constraint
  - In each volume, vvol=1,Mvol,
    - \* the logical array ImagneticOK (vvol) is set to .false.
    - \* The MPI node associated to the volume calls dfp100(). This routine calls ma02aa() (and might iterate on mp00ac()) and computes the field solution in each volume consistent with the constraint.
    - \* The MPI node associated to the volume calls dfp200(). This computes  $p + B^2/2$  (and the spectral constraints if required) at the interfaces in each volumes, as well as the derivatives of the force-balance if LComputeDerivatives=1.
  - After the parallelization loop over the volumes, brcast() is called to broadcast the required information.
- · Global constraint

The MPI node 0 minimizes the constraint with HYBRID1() by iterating on dfp100() until the field matches the constraint. Other MPI nodes enter the subroutine loop\_dfp100(). In loop\_dfp100(), each MPI node

- calls dfp100(),
- solves the field in its associated volumes,
- communicates the field to the node  $\boldsymbol{0}$  and
- repeats this loop until the node 0 sends a flag iflag=5.

### broadcasting

The required quantities are broadcast by brcast().

### construction of force

• The force vector,  $\mathbf{F}(\mathbf{x})$ , is a combination of the pressure-imbalance Fourier harmonics,  $[[p+B^2/2]]_{i,v}$ , where i labels Fourier harmonic and v is the interface label:

$$F_{i,v} \equiv \left[ (p_{v+1} + B_{i,v+1}^2/2) - (p_v + B_{i,v}^2/2) \right] \times \text{BBweight}_i,$$
 (60)

where BBweight\_i is defined in preset(); and the spectral condensation constraints,

$$F_{i,v} \equiv I_{i,v} \times \text{epsilon} + S_{i,v,1} \times \text{sweight}_v - S_{i,v+1,0} \times \text{sweight}_{v+1},$$
 (61)

where the spectral condensation constraints,  $I_{i,v}$ , and the "star-like" poloidal angle constraints,  $S_{i,v,\pm 1}$ , are calculated and defined in Iforce(); and the <code>sweight</code>  $_v$  are defined in <code>preset()</code>. All quantities local to a volume are computed in <code>dfp200()</code>, information is then broadcasted to the MPI node 0 in <code>dforce()</code> and the global force is evaluated.

## construct derivatives of matrix equation

• Matrix perturbation theory is used to compute the derivatives of the solution, i.e. the Beltrami fields, as the geometry of the interfaces changes:

#### **Parameters**

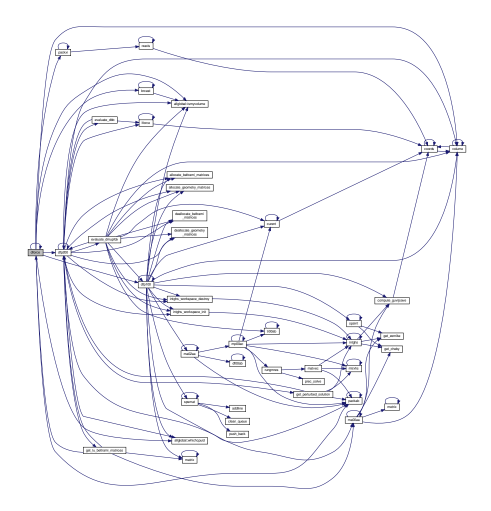
in	NGdof	number of global degrees of freedom
in	position	
out	force	
in	LComputeDerivatives	
in,out	LComputeAxis	

References allglobal::ate, allglobal::ato, allglobal::aze, allglobal::aze, allglobal::bbe, allglobal::bbo, allglobal::bbweight, allglobal::bemn, allglobal::bomn, brcast(), allglobal::cpus, allglobal::dbdmp, allglobal::dbdx, allglobal::dessian, allglobal::dffdrz, dforce(), dfp100(), dfp200(), allglobal::diotadxup, allglobal::ditgpdxtp, allglobal::dmupfdx, allglobal::dpflux, inputlist::drz, allglobal::dtflux, allglobal::energy, inputlist::epsilon, allglobal::forceerr, constants::half, allglobal::hessian, allglobal::iemn, inputlist::igeometry, allglobal::iie, allglobal::iio, allglobal::ima, allglobal::imagneticok, allglobal::in, allglobal::iomn, allglobal::ipdtdpf, allglobal::iquad, allglobal::irbc, allglobal::irbs, allglobal::ismyvolume(), allglobal::ismyvolumevalue, allglobal::izbc, allglobal::izbs, allglobal::ilbbintegral, inputlist::lcheck, inputlist::lconstraint, allglobal::localconstraint, numerical::logtolerance, allglobal::lplasmaregion, inputlist::lrad, allglobal::lhessianallocated, allglobal::mn, allglobal::mpi\_comm\_spec, inputlist::mu, inputlist::mupftol, allglobal::myid, allglobal::nadof, allglobal::ncpu, allglobal::notstellsym, inputlist::ntor, inputlist::nvol, constants::one, fileunits::ounit, packab(), packxi(), constants::pi, constants::pi2, allglobal::psifactor, allglobal::semn, allglobal::solution, allglobal::somn, allglobal::seweight, constants::two, volume(), allglobal::whichcpuid(), inputlist::wmacros, allglobal::xoffset, allglobal::yesstellsym, and constants::zero.

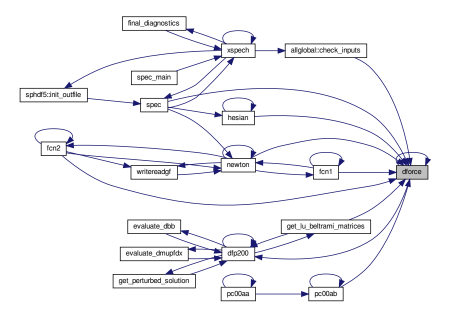
Referenced by allglobal::check\_inputs(), dforce(), fcn1(), fcn2(), get\_lu\_beltrami\_matrices(), hesian(), newton(), pc00ab(), and spec().

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Here is the call graph for this function:

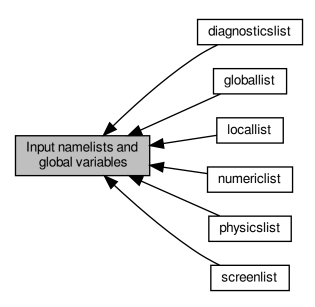


Here is the caller graph for this function:



# 7.7 Input namelists and global variables

Collaboration diagram for Input namelists and global variables:



# Modules

· physicslist

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The namelist physicslist controls the geometry, profiles, and numerical resolution.

· numericlist

The namelist numericlist controls internal resolution parameters that the user rarely needs to consider.

· locallist

The namelist locallist controls the construction of the Beltrami fields in each volume.

· globallist

The namelist globallist controls the search for global force-balance.

· diagnosticslist

The namelist diagnosticslist controls post-processor diagnostics, such as Poincaré plot resolution, etc.

· screenlist

The namelist screenlist controls screen output. Every subroutine, e.g. xy00aa.h, has its own write flag, Wxy00aa.h

#### **Modules**

· module constants

some constants used throughout the code

· module numerical

platform-dependant numerical resolution

· module fileunits

central definition of file units to avoid conflicts

· module cputiming

timing variables

module typedefns

type definitions for custom datatypes

## **Functions/Subroutines**

• subroutine inputlist::initialize\_inputs

### **Variables**

• integer, parameter inputlist::mnvol = 256

The maximum value of Nvol is MNvol=256.

• integer, parameter inputlist::mmpol = 128

The maximum value of Mpol is MNpol=64.

• integer, parameter inputlist::mntor = 128

The maximum value of Ntor is MNtor=64.

# 7.7.1 Detailed Description

Input namelists.

## 7.8 "local" force

## **Functions/Subroutines**

subroutine Iforce (Ivol, iocons, ideriv, Ntz, dBB, XX, YY, length, DDI, MMI, iflag)
 Computes B<sup>2</sup>, and the spectral condensation constraints if required, on the interfaces, \( \mathcal{I}\_i \).

# 7.8.1 Detailed Description

## 7.8.2 Function/Subroutine Documentation

```
7.8.2.1 Iforce() subroutine lforce (
    integer, intent(in) lvol,
    integer, intent(in) iocons,
    integer, intent(in) ideriv,
    integer, intent(in) Ntz,
    real, dimension(1:ntz, -1:2) dBB,
    real, dimension(1:ntz) XX,
    real, dimension(1:ntz) YY,
    real, dimension(1:ntz) length,
    real DDl,
    real MMl,
    integer, intent(in) iflag)
```

Computes  $B^2$ , and the spectral condensation constraints if required, on the interfaces,  $\mathcal{I}_i$ . field strength

- The field strength is given by  $B^2 = B^s B_s + B^\theta B_\theta + B^\zeta B_\zeta$ , and on the interfaces  $B^s = 0$  by construction.
- The magnetic field is  $\sqrt{g} \mathbf{B} = (\partial_{\theta} A_{\zeta} \partial_{\zeta} A_{\theta}) \mathbf{e}_s \partial_s A_{\zeta} \mathbf{e}_{\theta} + \partial_s A_{\theta} \mathbf{e}_{\zeta}.$
- The covariant components of the field are computed via  $B_{\theta}=B^{\theta}g_{\theta\theta}+B^{\zeta}g_{\theta\zeta}$  and  $B_{\zeta}=B^{\theta}g_{\theta\zeta}+B^{\zeta}g_{\zeta\zeta}$ .
- The expression for  ${\cal B}^2$  is

$$(\sqrt{g})^2 B^2 = A'_{\zeta} A'_{\zeta} g_{\theta\theta} - 2 A'_{\zeta} A'_{\theta} g_{\theta\zeta} + A'_{\theta} A'_{\theta} g_{\zeta\zeta}, \tag{62}$$

where the " $\prime$ " denotes derivative with respect to s.

· The quantity returned is

$$F \equiv \text{pscale} \times \frac{P}{V^{\gamma}} + \frac{B^2}{2},\tag{63}$$

where  $P \equiv \text{adiabatic}$  and  $V \equiv \text{volume}$ .

# spectral constraints

- In addition to the physical-force-balance constraints, namely that  $[[p + B^2/2]] = 0$  across the interfaces, additional angle constraints are required to obtain a unique Fourier representation of the interface geometry.
- Introducing the angle functional: a weighted combination of the "polar" constraint; the normalized, poloidal, spectral width (Hirshman & Meier (1985) [3], Hirshman & Breslau (1998) [2]) the poloidal-angle origin constraint; and the "length" of the angle curves

where i labels the interfaces, and

$$\Theta_{i,\theta} \equiv \frac{x y_{\theta} - x_{\theta} y}{x^2 + y^2},\tag{65}$$

$$M_i \equiv \frac{\sum_j m_j^p (R_{j,i}^2 + Z_{j,i}^2)}{\sum_j (R_{j,i}^2 + Z_{j,i}^2)},$$
(66)

$$L_i \equiv \sqrt{[R_i(\theta,\zeta) - R_{i-1}(\theta,\zeta)]^2 + [Z_i(\theta,\zeta) - Z_{i-1}(\theta,\zeta)]^2},$$
(67)

and where j labels the Fourier harmonics. The  $\alpha_i$ ,  $\beta_i$ ,  $\gamma_i$  and  $\delta_i \equiv \text{sweight}$  are user-supplied weight factors.

• The polar constraint is derived from defining  $\tan\Theta\equiv y/x$ , where

$$x(\theta,\zeta) \equiv R_i(\theta,\zeta) - R_{i,0}(\zeta),$$
 (68)

$$y(\theta,\zeta) \equiv Z_i(\theta,\zeta) - Z_{i,0}(\zeta), \tag{69}$$

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and where the geometric center of each interface is given by the arc-length weighted integrals, see rzaxis(),

$$R_{i,0} \equiv \int_0^{2\pi} d\theta \ R_i(\theta,\zeta) \sqrt{R_{i,\theta}(\theta,\zeta)^2 + Z_{i,\theta}(\theta,\zeta)^2}, \tag{70}$$

$$Z_{i,0} \equiv \int_0^{2\pi} d\theta \ Z_i(\theta,\zeta) \sqrt{R_{i,\theta}(\theta,\zeta)^2 + Z_{i,\theta}(\theta,\zeta)^2},\tag{71}$$

and  $\cos\Theta=x/\sqrt{x^2+y^2}$  has been used to simplify the expressions and to avoid divide-by-zero.

Only "poloidal tangential" variations will be allowed to find the extremum of F, which are described by

$$\delta R_i(\theta, \zeta) \equiv R_{i,\theta}(\theta, \zeta) \, \delta u_i(\theta, \zeta), \tag{72}$$

$$\delta Z_i(\theta,\zeta) \equiv Z_{i,\theta}(\theta,\zeta) \, \delta u_i(\theta,\zeta),$$
 (73)

from which it follows that the variation in each Fourier harmonic is

$$\delta R_{j,i} = \oint \!\! \int \!\! d\theta d\zeta \ R_{i,\theta}(\theta,\zeta) \, \delta u_i(\theta,\zeta) \, \cos(m_j \theta - n_j \zeta), \tag{74}$$

$$\delta Z_{j,i} = \oint \!\! \int \!\! d\theta d\zeta \ Z_{i,\theta}(\theta,\zeta) \, \delta u_i(\theta,\zeta) \, \sin(m_j \theta - n_j \zeta), \tag{75}$$

and

$$\delta R_{i,\theta}(\theta,\zeta) \equiv R_{i,\theta\theta}(\theta,\zeta) \, \delta u_i(\theta,\zeta) + R_{i,\theta}(\theta,\zeta) \, \delta u_{i,\theta}(\theta,\zeta) \tag{76}$$

$$\delta Z_{i,\theta}(\theta,\zeta) \equiv Z_{i,\theta\theta}(\theta,\zeta) \,\delta u_i(\theta,\zeta) + Z_{i,\theta}(\theta,\zeta) \,\delta u_{i,\theta}(\theta,\zeta) \tag{77}$$

• The variation in F is

$$\delta F = \sum_{i=1}^{N-1} \alpha_{i} \oint d\theta d\zeta \left(\frac{-2\Theta_{i,\theta\theta}}{\Theta_{i,\theta}^{2}}\right) \delta u_{i} 
+ \sum_{i=1}^{N-1} \beta_{i} \oint d\theta d\zeta \left(R_{i,\theta}X_{i} + Z_{i,\theta}Y_{i}\right) \delta u_{i} 
+ \sum_{i=1}^{N-1} \gamma_{i} \int d\zeta \left(Z_{i}(0,\zeta) - Z_{i,0}\right) Z_{i,\theta} \delta u_{i} 
+ \sum_{i=1}^{N-1} \delta_{i} \oint d\theta d\zeta \left(\frac{\Delta R_{i}R_{i,\theta} + \Delta Z_{i}Z_{i,\theta}}{L_{i}}\right) \delta u_{i} 
- \sum_{i=1}^{N-1} \delta_{i+1} \oint d\theta d\zeta \left(\frac{\Delta R_{i+1}R_{i,\theta} + \Delta Z_{i+1}Z_{i,\theta}}{L_{i+1}}\right) \delta u_{i}$$
(78)

where, for the stellarator symmetric case,

$$X_i \equiv \sum_{j} (m_j^p - M_i) R_{j,i} \cos(m_j \theta - n_j \zeta), \tag{79}$$

$$Y_i \equiv \sum_{j} (m_j^p - M_i) Z_{j,i} \sin(m_j \theta - n_j \zeta), \tag{80}$$

and

$$\Delta R_i \equiv R_i(\theta, \zeta) - R_{i-1}(\theta, \zeta), \tag{81}$$

$$\Delta Z_i \equiv Z_i(\theta, \zeta) - Z_{i-1}(\theta, \zeta), \tag{82}$$

• The spectral constraints derived from Eqn. (78) are

$$I_{i}(\theta,\zeta) \equiv -2\alpha_{i} \frac{\Theta_{i,\theta\theta}}{\Theta_{i,\theta}^{2}} + \beta_{i} \left( R_{i,\theta} X_{i} + Z_{i,\theta} Y_{i} \right) + \gamma_{i} \left( Z_{i}(0,\zeta) - Z_{i,0} \right) Z_{i,\theta}(0,\zeta)$$

$$+ \delta_{i} \frac{\Delta R_{i} R_{i,\theta} + \Delta Z_{i} Z_{i,\theta}}{L_{i}} - \delta_{i+1} \frac{\Delta R_{i+1} R_{i,\theta} + \Delta Z_{i+1} Z_{i,\theta}}{L_{i+1}}$$
(83)

- Note that choosing p=2 gives  $X=-R_{\theta\theta}$  and  $Y=-Z_{\theta\theta}$ , and the spectrally condensed angle constraint,  $R_{\theta}X+Z_{\theta}Y=0$ , becomes  $\partial_{\theta}(R_{\theta}^2+Z_{\theta}^2)=0$ , which defines the equal arc length angle.
- The poloidal-angle origin term, namely  $\gamma_i\left(Z_i(0,\zeta)-Z_{i,0}\right)Z_{i,\theta}(0,\zeta)$  is only used to constrain the  $m_j=0$  harmonics.
- The construction of the angle functional was influenced by the following considerations:
  - The minimal spectral width constraint is very desirable as it reduces the required Fourier resolution, but it does not constrain the m=0 harmonics and the minimizing spectral-width poloidal-angle may not be consistent with the poloidal angle used on adjacent interfaces.
  - The regularization of the vector potential and the coordinate interpolation near the coordinate origin (see elsewhere) assumes that the poloidal angle is the polar angle.
  - The user will provide the Fourier harmonics of the boundary, and thus the user will implicitly define the poloidal angle used on the boundary.
  - Minimizing the length term will ensure that the poloidal angle used on each interface is smoothly connected to the poloidal angle used on adjacent interfaces.
- A suitable choice of the weight factors,  $\alpha_i$ ,  $\beta_i$ ,  $\gamma_i$  and  $\delta_i$ , will ensure that the polar constraint dominates for the innermost surfaces and that this constraint rapidly becomes insignificant away from the origin; that the minimal spectral constraint dominates in the "middle"; and that the minimizing length constraint will be significant near the origin and dominant near the edge, so that the minimizing spectral width angle will be continuously connected to the polar angle on the innermost surfaces and the user-implied angle at the plasma boundary. The length constraint should not be insignificant where the spectral constraint is dominant (so that the m=0 harmonics are constrained).
- The polar constraint does not need normalization. The spectral width constraint has already been normalized. The length constraint is not yet normalized, but perhaps it should be.
- The spectral constraints given in Eqn. (83) need to be differentiated with respect to the interface Fourier harmonics,  $R_{j,i}$  and  $Z_{j,i}$ . The first and second terms lead to a block diagonal hessian, and the length term leads to a block tri-diagonal hessian.
- Including the poloidal-angle origin constraint means that the polar angle constraint can probably be ignored, i.e.  $\alpha_i=0$ .

### **Parameters**

Ivol	
iocons	
ideriv	
Ntz	
dBB	
XX	
YY	
length	
DDI	
MMI	
iflag	
	iocons ideriv Ntz dBB XX YY length DDI MMI

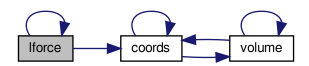
References inputlist::adiabatic, allglobal::ate, allglobal::ato, allglobal::aze, allglobal::azo, allglobal::bemn, allglobal::cfmn, allglobal::cfmn, allglobal::comn, coords(), allglobal::cpus, allglobal::drij, allglobal::dzij, allglobal::efmn, allglobal::efmn, inputlist::gamma, allglobal::guvij, constants::half, allglobal::iemn, inputlist::igeometry, allglobal::ijimag, allglobal::ijreal, allglobal::irij, allglobal::irij, allglobal::izbc, allglobal::izbs, allglobal::izij, allglobal::jimag, allglobal::jireal, inputlist::lcheck, allglobal::lcoordinatesingularity, lforce(), inputlist::lrad, allglobal::mmpp, allglobal::mn, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ncpu, allglobal::notstellsym, allglobal::nt, inputlist::nvol, allglobal::nz, allglobal::odmn, allglobal::ofmn, constants::one, fileunits::ounit, allglobal::pemn, allglobal::pemn, inputlist::pscale, allglobal::regumm, allglobal::rtt, allglobal::semn,

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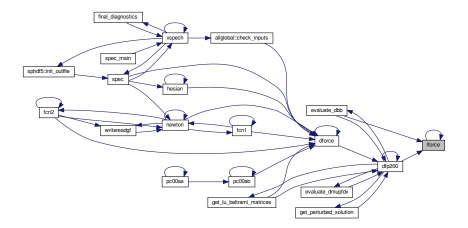
allglobal::sfmn, allglobal::sg, allglobal::simn, allglobal::trij, allglobal::trij, allglobal::trij, allglobal::tvolume, allglobal::yesstellsym, and constants::zero.

Referenced by dfp200(), evaluate\_dbb(), and lforce().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.9 Integrals

## **Functions/Subroutines**

- subroutine df00ab (pNN, xi, Fxi, DFxi, Ldfjac, iflag)
  - Evaluates volume integrals, and their derivatives w.r.t. interface geometry, using "packed" format.
- subroutine ma00aa (Iquad, mn, Ivol, Irad)
  - Calculates volume integrals of Chebyshev polynomials and metric element products.
- subroutine spsint (Iquad, mn, Ivol, Irad)

Calculates volume integrals of Chebyshev-polynomials and metric elements for preconditioner.

## 7.9.1 Detailed Description

## 7.9.2 Function/Subroutine Documentation

```
7.9.2.1 df00ab() subroutine df00ab (
    integer, intent(in) pNN,
    real, dimension(0:pnn-1), intent(in) xi,
    real, dimension(0:pnn-1), intent(out) Fxi,
    real, dimension(0:ldfjac-1,0:pnn-1), intent(out) DFxi,
```

```
integer, intent(in) Ldfjac,
integer, intent(in), value iflag)
```

Evaluates volume integrals, and their derivatives w.r.t. interface geometry, using "packed" format.

#### **Parameters**

in	pNN	
in	xi	
out	Fxi	
out	DFxi	
in	Ldfjac	
in	iflag	

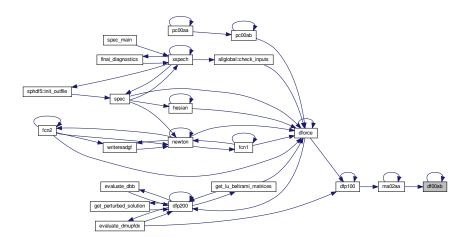
References allglobal::cpus, df00ab(), allglobal::dma, allglobal::dmd, constants::half, inputlist::helicity, allglobal::ivol, allglobal::mbpsi, allglobal::mpi\_comm\_spec, allglobal::myid, inputlist::nvol, constants::one, fileunits::ounit, numerical::small, constants::two, and constants::zero.

Referenced by df00ab(), and ma02aa().

Here is the call graph for this function:



Here is the caller graph for this function:



```
7.9.2.2 ma00aa() subroutine ma00aa (
integer, intent(in) lquad,
integer, intent(in) mn,
integer, intent(in) lvol,
integer, intent(in) lrad)
```

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Calculates volume integrals of Chebyshev polynomials and metric element products. **Chebyshev-metric information** 

· The following quantities are calculated:

DToocc(l,p,i,j) 
$$\equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}_{p,j} \oint \!\!\!\!\! \oint \!\!\!\! d\theta d\zeta \, \cos \alpha_i \cos \alpha_j$$
 (84)

DToocs (l,p,i,j) 
$$\equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}_{p,j} \, \oint \!\!\!\!\! \oint d\theta d\zeta \, \cos \alpha_i \sin \alpha_j$$
 (85)

DToosc(l,p,i,j) 
$$\equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}_{p,j} \, \oint \!\!\!\! \oint \!\!\! d\theta d\zeta \, \sin \alpha_i \cos \alpha_j$$
 (86)

DTooss(1,p,i,j) 
$$\equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}_{p,j} \, \oint \!\!\!\!\! \oint d\theta d\zeta \, \sin \alpha_i \sin \alpha_j$$
 (87)

TDstcc(l,p,i,j) 
$$\equiv \int ds \, \overline{T}_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \int \!\!\!\! d\theta d\zeta \, \cos \alpha_i \cos \alpha_j \, \overline{g}_{s\theta}$$
 (92)

TDstcs(l,p,i,j) 
$$\equiv \int ds \, \overline{T}_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\!\! \oint d\theta d\zeta \, \cos \alpha_i \sin \alpha_j \, \bar{g}_{s\theta}$$
 (93)

$$\mathsf{TDstsc}(\mathsf{l},\mathsf{p},\mathsf{i},\mathsf{j}) \equiv \int ds \, \overline{T}_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\!\! \oint \!\!\! d\theta d\zeta \, \sin\alpha_i \cos\alpha_j \, \overline{g}_{s\theta} \tag{94}$$

TDstss(l,p,i,j) 
$$\equiv \int ds \, \overline{T}_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \oint \!\!\! d\theta d\zeta \, \sin \alpha_i \sin \alpha_j \, \bar{g}_{s\theta}$$
 (95)

TDstcc(l,p,i,j) 
$$\equiv \int ds \, \overline{T}_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \int d\theta \, d\zeta \, \cos \alpha_i \cos \alpha_j \, \overline{g}_{s\zeta}$$
 (96)

TDstcs(l,p,i,j) 
$$\equiv \int ds \, \overline{T}_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \oint d\theta d\zeta \, \cos \alpha_i \sin \alpha_j \, \overline{g}_{s\zeta}$$
 (97)

TDstsc(l,p,i,j) 
$$\equiv \int ds \, \overline{T}_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \oint \!\!\! d\theta d\zeta \, \sin \alpha_i \cos \alpha_j \, \overline{g}_{s\zeta}$$
 (98)

TDstss(l,p,i,j) 
$$\equiv \int ds \, \overline{T}_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \oint \!\!\! d\theta d\zeta \, \sin \alpha_i \sin \alpha_j \, \bar{g}_{s\zeta}$$
 (99)

$$\text{DDstcc}(1,p,i,j) \equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \oint \!\! d\theta d\zeta \, \cos\alpha_i \cos\alpha_j \, \overline{g}_{\theta\theta} \tag{100}$$

DDstcs(l,p,i,j) 
$$\equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \oint \!\!\! d\theta d\zeta \, \cos \alpha_i \sin \alpha_j \, \overline{g}_{\theta\theta}$$
 (101)

$$\text{DDstsc}(1,p,i,j) \equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \oint \!\! d\theta d\zeta \, \sin \alpha_i \cos \alpha_j \, \bar{g}_{\theta\theta} \tag{102}$$

$$\mathsf{DDstcc}(\mathsf{l},\mathsf{p},\mathsf{i},\mathsf{j}) \equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\!\! \oint \!\!\! d\theta d\zeta \, \cos\alpha_i \cos\alpha_j \, \overline{g}_{\theta\zeta} \tag{104}$$

DDstcs(l,p,i,j) 
$$\equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\!\! \oint \!\!\!\! d\theta d\zeta \, \cos \alpha_i \sin \alpha_j \, \bar{g}_{\theta\zeta}$$
 (105)

$$\text{DDstsc}(1,p,i,j) \equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \oint \!\!\! d\theta d\zeta \, \sin\alpha_i \cos\alpha_j \, \overline{g}_{\theta\zeta} \tag{106}$$

DDstss(l,p,i,j) 
$$\equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \oint \!\!\! d\theta d\zeta \, \sin \alpha_i \sin \alpha_j \, \bar{g}_{\theta\zeta}$$
 (107)

$$\text{DDstcc}(1, p, i, j) \equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \oint \!\!\! d\theta d\zeta \, \cos \alpha_i \cos \alpha_j \, \overline{g}_{\zeta\zeta} \tag{108}$$

DDstcs(l,p,i,j) 
$$\equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}'_{p,j} \, \phi \!\!\!\!/ \, d\theta d\zeta \, \cos \alpha_i \sin \alpha_j \, \overline{g}_{\zeta\zeta}$$
 (109)

$$\text{DDstsc}(l,p,i,j) \equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\! \oint \!\!\! d\theta d\zeta \, \sin\alpha_i \cos\alpha_j \, \bar{g}_{\zeta\zeta} \tag{110}$$

DDstss(l,p,i,j) 
$$\equiv \int ds \, \overline{T}'_{l,i} \, \overline{T}'_{p,j} \, \oint \!\!\!\!\! \oint \!\!\!\! d\theta d\zeta \, \sin \alpha_i \sin \alpha_j \, \bar{g}_{\zeta\zeta}$$
 (111)

where  $\overline{T}_{l,i} \equiv T_l \, \bar{s}^{m_i/2}$  if the domain includes the coordinate singularity, and  $\overline{T}_{l,i} \equiv T_l$  if not; and  $\bar{g}_{\mu\nu} \equiv g_{\mu\nu}/\sqrt{g}$ .

• The double-angle formulae are used to reduce the above expressions to the Fourier harmonics of  $\bar{g}_{\mu\nu}$ : see kija and kijs, which are defined in preset.f90 .

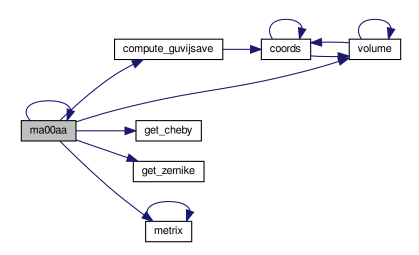
#### **Parameters**

	in	Iquad   degree of quadrature	
in mn number of Fourier harmonics		number of Fourier harmonics	
in <i>Ivol</i> index of nested volume		index of nested volume	
ĺ	in	Irad	order of Chebychev polynomials

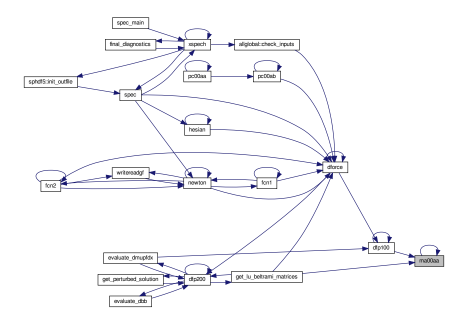
Referenced by dfp100(), get lu beltrami matrices(), and ma00aa().

7.9 Integrals 43

Here is the call graph for this function:



Here is the caller graph for this function:



Calculates volume integrals of Chebyshev-polynomials and metric elements for preconditioner.

Computes the integrals needed for spsmat.f90. Same as ma00aa.f90, but only compute the relevant terms that are non-zero.

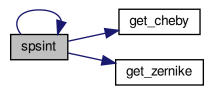
#### **Parameters**

Iquad	
mn	
Ivol	
Irad	

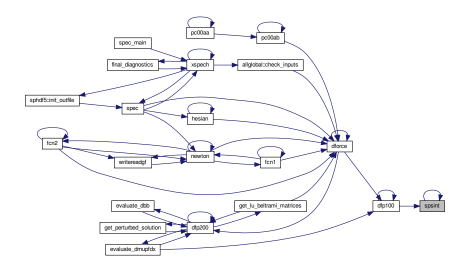
References allglobal::cpus, allglobal::ddttcc, allglobal::ddttcs, allglobal::ddttcs, allglobal::ddttcs, allglobal::ddttcc, allglobal::ddttcc, allglobal::ddttcc, allglobal::ddttcc, allglobal::ddttcc, allglobal::ddttcc, allglobal::ddttcc, allglobal::ddttcc, allglobal::ddttccc, allglobal::ddttccc, allglobal::ddttccc, allglobal::ddttccc, allglobal::ddttccc, allglobal::dtcccc, allglobal::dtcccc, allglobal::dtcccc, allglobal::dtcccc, allglobal::dtcccc, allglobal::dtcccc, allglobal::dtcccc, allglobal::im, allglobal::im, allglobal::im, allglobal::im, allglobal::kija, allglobal::kija, allglobal::kija, allglobal::lccoordinatesingularity, allglobal::mne, allglobal::mpi\_comm\_spec, inputlist::mpol, allglobal::myid, allglobal::ncpu, allglobal::notstellsym, allglobal::ntt, constants::one, fileunits::ounit, constants::pi, constants::pi2, allglobal::regumm, numerical::small, spsint(), numerical::sqrtmachprec, allglobal::tdstcc, allglobal::tdstcs, allglobal::tdstsc, allglobal::tdstsc, allglobal::tdstsc, allglobal::tdstsc, allglobal::tdstsc, allglobal::tdstsc, allglobal::tdstsc, allglobal::ttsscc, allglobal::ttsscc,

Referenced by dfp100(), and spsint().

Here is the call graph for this function:



Here is the caller graph for this function:



7.10 Solver/Driver 45

## 7.10 Solver/Driver

#### **Functions/Subroutines**

• subroutine ma02aa (Ivol, NN)

Constructs Beltrami field in given volume consistent with flux, helicity, rotational-transform and/or parallel-current constraints.

## 7.10.1 Detailed Description

#### 7.10.2 Function/Subroutine Documentation

Constructs Beltrami field in given volume consistent with flux, helicity, rotational-transform and/or parallel-current constraints.

#### **Parameters**

	in	Ivol	index of nested volume for which to run this
Ī	in	NN	number of degrees of freedom in the (packed format) vector potential;

## sequential quadratic programming

- Only relevant if LBsequad=T . See LBeltrami for details.
- Documentation on the implementation of E04UFF is under construction.

## **Newton method**

• Only relevant if LBnewton=T . See LBeltrami for details.

## linear method

- Only relevant if LBlinear=T . See LBeltrami for details.
- The quantity  $\mu$  is *not* not treated as a "magnetic" degree-of-freedom equivalent to in the degrees-of-freedom in the magnetic vector potential (as it strictly should be, because it is a Lagrange multiplier introduced to enforce the helicity constraint).
- In this case, the Beltrami equation,  $\nabla \times \mathbf{B} = \mu \mathbf{B}$ , is *linear* in the magnetic degrees-of-freedom.
- · The algorithm proceeds as follows:

### plasma volumes

- In addition to the enclosed toroidal flux,  $\Delta\psi_t$ , which is held constant in the plasma volumes, the Beltrami field in a given volume is assumed to be parameterized by  $\mu$  and  $\Delta\psi_p$ . (Note that  $\Delta\psi_p$  is not defined in a torus.)
- These are "packed" into an array, e.g.  $\mu \equiv (\mu, \Delta \psi_p)^T$ , so that standard library routines , e.g. C05PCF, can be used to (iteratively) find the appropriately-constrained Beltrami solution, i.e.  $\mathbf{f}(\mu) = 0$ .
- The function  $f(\mu)$ , which is computed by mp00ac(), is defined by the input parameter Lconstraint:
  - \* If Lconstraint = -1, 0, then  $\mu$  is not varied and Nxdof=0.
  - \* If Lconstraint = 1, then  $\mu$  is varied to satisfy the transform constraints; and Nxdof=1 in the simple torus and Nxdof=2 in the annular regions. (Note that in the "simple-torus" region, the enclosed poloidal flux  $\Delta \psi_p$  is not well-defined, and only  $\mu=\mu_1$  is varied in order to satisfy the transform constraint on the "outer" interface of that volume.)
  - \* Todo If Lconstraint = 2, then  $\mu=\mu_1$  is varied in order to satisfy the helicity constraint, and  $\Delta\psi_p=\mu_2$  is *not* varied, and Nxdof=1. (under re-construction)

#### vacuum volume

- In the vacuum,  $\mu=0$ , and the enclosed fluxes,  $\Delta\psi_t$  and  $\Delta\psi_p$ , are considered to parameterize the family of solutions. (These quantities may not be well-defined if  ${\bf B}\cdot{\bf n}\neq 0$  on the computational boundary.)
- These are "packed" into an array,  $\mu \equiv (\Delta \psi_t, \Delta \psi_p)^T$ , so that, as above, standard routines can be used to iteratively find the appropriately constrained solution, i.e.  $\mathbf{f}(\mu) = 0$ .
- The function  $f(\mu)$ , which is computed by mp00ac(), is defined by the input parameter Lconstraint:
  - \* If Lconstraint = -1, then  $\mu$  is not varied and Nxdof=0.
  - \* If Lconstraint = 0,2, then  $\mu$  is varied to satisfy the enclosed current constraints, and Nxdof=2.
  - \* If Lconstraint = 1, then  $\mu$  is varied to satisfy the constraint on the transform on the inner boundary  $\equiv$  plasma boundary and the "linking" current, and Nxdof=2.
- The Beltrami fields, and the rotational-transform and helicity etc. as required to determine the function  $f(\mu)$  are calculated in mp00ac().
- This routine, mp00ac(), is called iteratively if Nxdof>1 via C05PCF to determine the appropriately constrained Beltrami field,  $\mathbf{B}_{\mu}$ , so that  $\mathbf{f}(\mu)=0$ .
- The input variables mupftol and mupfits control the required accuracy and maximum number of iterations.
- If Nxdof=1, then mp00ac() is called only once to provide the Beltrami fields with the given value of  $\mu$ .

### debugging: finite-difference confirmation of the derivatives of the rotational-transform

- Note that the rotational-transform (if required) is calculated by tr00ab(), which is called by mp00ac().
- If Lconstraint=1, then mp00ac() will ask tr00ab() to compute the derivatives of the transform with respect to variations in the helicity-multiplier,  $\mu$ , and the enclosed poloidal-flux,  $\Delta\psi_p$ , so that C05PCF may more efficiently find the solution.
- · The required derivatives are

$$\frac{\partial \, \iota}{\partial \mu}$$
 (112)

$$\frac{\partial +}{\partial \Delta \psi_p} \tag{113}$$

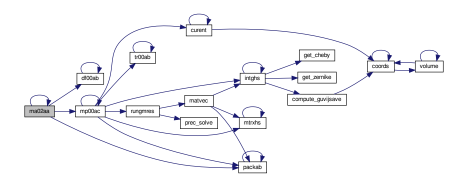
to improve the efficiency of the iterative search. A finite difference estimate of these derivatives is available; need DEBUG, Lcheck=2 and Lconstraint=1.

References allglobal::ate, allglobal::cpus, df00ab(), allglobal::dma, allglobal::dmb, allglobal::dmd, allglobal::dpflux, allglobal::dtflux, constants::half, inputlist::helicity, allglobal::im, allglobal::imagneticok, allglobal::in, allglobal::ivol, allglobal::lbbintegral, allglobal::lbbintegra

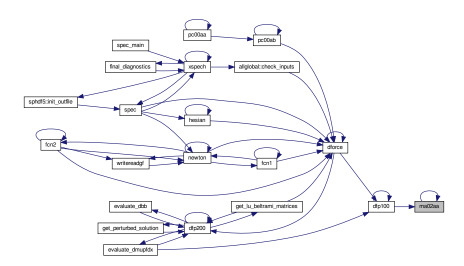
Referenced by dfp100(), and ma02aa().

7.11 Build matrices 47

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.11 Build matrices

## Functions/Subroutines

• subroutine matrix (Ivol, mn, Irad)

Constructs energy and helicity matrices that represent the Beltrami linear system. gauge conditions

• subroutine mtrxhs (Ivol, mn, Irad, resultA, resultD, idx)

Constructs matrices that represent the Beltrami linear system, matrix-free.

• subroutine spsmat (Ivol, mn, Irad)

Constructs matrices for the precondtioner.

# 7.11.1 Detailed Description

# 7.11.2 Function/Subroutine Documentation

integer, intent(in) mn,
integer, intent(in) lrad )

Constructs energy and helicity matrices that represent the Beltrami linear system.

## gauge conditions

• In the v-th annulus, bounded by the (v-1)-th and v-th interfaces, a general covariant representation of the magnetic vector-potential is written

$$\bar{\mathbf{A}} = \bar{A}_s \nabla s + \bar{A}_\theta \nabla \theta + \bar{A}_\zeta \nabla \zeta eta. \tag{114}$$

• To this add  $\nabla g(s, \theta, \zeta)$ , where g satisfies

$$\begin{array}{lcl} \partial_s g(s,\theta,\zeta) & = & - & \bar{A}_s(s,\theta,\zeta) \\ \partial_\theta g(-1,\theta,\zeta) & = & - & \bar{A}_\theta(-1,\theta,\zeta) \\ \partial_\zeta g(-1,0,\zeta) & = & - & \bar{A}_\zeta(-1,0,\zeta). \end{array} \tag{115}$$

• Then  $\mathbf{A} = \bar{\mathbf{A}} + \nabla g$  is given by  $\mathbf{A} = A_{\theta} \nabla \theta + A_{\zeta} \nabla \zeta$  with

$$A_{\theta}(-1,\theta,\zeta) = 0 \tag{116}$$

$$A_{\zeta}(-1,0,\zeta) = 0 \tag{117}$$

- This specifies the gauge: to see this, notice that no gauge term can be added without violating the conditions in Eqn. (116) or Eqn. (117).
- · Note that the gauge employed in each volume is distinct.

## boundary conditions

- The magnetic field is  $\sqrt{g} \mathbf{B} = (\partial_{\theta} A_{\zeta} \partial_{\zeta} A_{\theta}) \mathbf{e}_{s} \partial_{s} A_{\zeta} \mathbf{e}_{\theta} + \partial_{s} A_{\theta} \mathbf{e}_{\zeta}$ .
- In the annular volumes, the condition that the field is tangential to the inner interface,  $\sqrt{g}\mathbf{B}\cdot\nabla s=0$  at s=-1, gives  $\partial_{\theta}A_{\zeta}-\partial_{\zeta}A_{\theta}=0$ . With the above condition on  $A_{\theta}$  given in Eqn. (116), this gives  $\partial_{\theta}A_{\zeta}=0$ , which with Eqn. (117) gives

$$A_{\zeta}(-1,\theta,\zeta) = 0. \tag{118}$$

• The condition at the outer interface, s=+1, is that the field is  $\sqrt{g}\,\mathbf{B}\cdot\nabla s=\partial_{\theta}A_{\zeta}-\partial_{\zeta}A_{\theta}=b$ , where b is supplied by the user. For each of the plasma regions, b=0. For the vacuum region, generally  $b\neq 0$ .

## enclosed fluxes

- In the plasma regions, the enclosed fluxes must be constrained.
- · The toroidal and poloidal fluxes enclosed in each volume are determined using

$$\int_{S} \mathbf{B} \cdot \mathbf{ds} = \int_{\partial S} \mathbf{A} \cdot \mathbf{dl}.$$
 (119)

## **Fourier-Chebyshev representation**

• The components of the vector potential,  $\mathbf{A}=A_{\theta}\nabla+A_{\zeta}\nabla\zeta$ , are

$$A_{\theta}(s,\theta,\zeta) = \sum_{i,l} A_{\theta,e,i,l} \, \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} A_{\theta,o,i,l} \, \overline{T}_{l,i}(s) \sin \alpha_i, \tag{120}$$

$$A_{\zeta}(s,\theta,\zeta) = \sum_{i,l} A_{\zeta,e,i,l} \, \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} A_{\zeta,o,i,l} \, \overline{T}_{l,i}(s) \sin \alpha_i, \tag{121}$$

where  $\overline{T}_{l,i}(s)$  is the **recombined** Chebyshev polynomial in a volume without an axis, or **modified** Zernike polynomial in a volume with an axis (i.e. only in the innermost volume, and only in cylindrical and toroidal geometry.) , and  $\alpha_j \equiv m_j \theta - n_j \zeta$ .

7.11 Build matrices 49

• The magnetic field,  $\sqrt{g} \mathbf{B} = \sqrt{g} B^s \mathbf{e}_s + \sqrt{g} B^\theta \mathbf{e}_\theta + \sqrt{g} B^\zeta \mathbf{e}_\zeta$ , is

$$\sqrt{g} \mathbf{B} = \mathbf{e}_{s} \sum_{i,l} \left[ (-m_{i} A_{\zeta,e,i,l} - n_{i} A_{\theta,e,i,l}) \overline{T}_{l,i} \sin \alpha_{i} + (+m_{i} A_{\zeta,o,i,l} + n_{i} A_{\theta,o,i,l}) \overline{T}_{l,i} \cos \alpha_{i} \right] 
+ \mathbf{e}_{\theta} \sum_{i,l} \left[ (-m_{i} A_{\zeta,e,i,l}) \overline{T}'_{l,i} \cos \alpha_{i} + (-m_{i} A_{\zeta,o,i,l}) \overline{T}'_{l,i} \sin \alpha_{i} \right] (122) 
+ \mathbf{e}_{\zeta} \sum_{i,l} \left[ (-m_{i} A_{\zeta,e,i,l}) \overline{T}'_{l,i} \cos \alpha_{i} + (-m_{i} A_{\zeta,o,i,l}) \overline{T}'_{l,i} \sin \alpha_{i} \right]$$

• The components of the velocity,  $\mathbf{v} \equiv v_s \nabla s + v_\theta \nabla \theta + v_\zeta \nabla \zeta eta$ , are

$$v_s(s,\theta,\zeta) = \sum_{i,l} v_{s,e,i,l} \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} v_{s,o,i,l} \overline{T}_{l,i}(s) \sin \alpha_i, \tag{123}$$

$$v_{\theta}(s,\theta,\zeta) = \sum_{i,l} v_{\theta,e,i,l} \, \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} v_{\theta,o,i,l} \, \overline{T}_{l,i}(s) \sin \alpha_i, \tag{124}$$

$$v_{\zeta}(s,\theta,\zeta) = \sum_{i,l} v_{\zeta,e,i,l} \, \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} v_{\zeta,o,i,l} \, \overline{T}_{l,i}(s) \sin \alpha_i. \tag{125}$$

## constrained energy functional

 The constrained energy functional in each volume depends on the vector potential and the Lagrange multipliers,

 $\mathcal{F} \equiv \mathcal{F}[A_{\theta,e,i,l},A_{\zeta,e,i,l},A_{\theta,o,i,l},v_{s,e,i,l},v_{s,o,i,l},v_{\theta,e,i,l},v_{\theta,o,i,l},v_{\zeta,e,i,l},v_{\zeta,o,i,l},\mu,a_i,b_i,c_i,d_i,e_i,f_i,g_1,h_1], \tag{126}$  and is given by:

$$\mathcal{F} \equiv \int \mathbf{B} \cdot \mathbf{B} \, dv + \int \mathbf{v} \cdot \mathbf{v} \, dv - \mu \left[ \int \mathbf{A} \cdot \mathbf{B} \, dv - K \right]$$

$$+ \sum_{i=1} a_i \left[ \sum_{l} A_{\theta,e,i,l} T_l(-1) - 0 \right]$$

$$+ \sum_{i=1} b_i \left[ \sum_{l} A_{\zeta,e,i,l} T_l(-1) - 0 \right]$$

$$+ \sum_{i=2} c_i \left[ \sum_{l} A_{\theta,o,i,l} T_l(-1) - 0 \right]$$

$$+ \sum_{i=2} d_i \left[ \sum_{l} A_{\zeta,o,i,l} T_l(-1) - 0 \right]$$

$$+ \sum_{i=2} e_i \left[ \sum_{l} (-m_i A_{\zeta,e,i,l} - n_i A_{\theta,e,i,l}) T_l(+1) - b_{s,i} \right]$$

$$+ \sum_{i=2} f_i \left[ \sum_{l} (+m_i A_{\zeta,o,i,l} + n_i A_{\theta,o,i,l}) T_l(+1) - b_{c,i} \right]$$

$$+ \sum_{i=2} A_{\theta,e,1,l} T_l(+1) - \Delta \psi_l$$

$$+ \sum_{l} A_{\zeta,e,1,l} T_l(+1) + \Delta \psi_p$$

where

- $a_i$ ,  $b_i$ ,  $c_i$  and  $d_i$  are Lagrange multipliers used to enforce the combined gauge and interface boundary condition on the inner interface,
- $e_i$  and  $f_i$  are Lagrange multipliers used to enforce the interface boundary condition on the outer interface, namely  $\sqrt{g}\,{f B}\cdot\nabla s=b$ ; and
- $g_1$  and  $h_1$  are Lagrange multipliers used to enforce the constraints on the enclosed fluxes.
- In each plasma volume the boundary condition on the outer interface is b=0.
- In the vacuum volume (only for free-boundary), we may set  $\mu=0$ .

• Note: in SPEC version >3.00, the basis recombination method is used to ensure the boundary condition on the inner side of an interface. The lagrange multipliers  $a_i, b_i, c_i, d_i$  are no longer used in volumes without a coordinate singularity. In a volume with a coordinate singularity, they are used only  $a_i, c_i$  with \$m=0,1\$ are excluded also due to Zernike basis recombination.

## derivatives of magnetic energy integrals

• The first derivatives of  $\int dv \ \mathbf{B} \cdot \mathbf{B}$  with respect to  $A_{\theta,e,i,l}$ ,  $A_{\theta,o,i,l}$ ,  $A_{\zeta,e,i,l}$  and  $A_{\zeta,o,i,l}$  are

$$\begin{split} &\frac{\partial}{\partial A_{\theta,e,i,l}}\int\!\!dv\;\mathbf{B}\cdot\mathbf{B} &= 2\int\!\!dv\;\mathbf{B}\cdot\frac{\partial\mathbf{B}}{\partial A_{\theta,e,i,l}} = 2\int\!\!dv\;\mathbf{B}\cdot\left[-n_i\overline{T}_{l,i}\sin\alpha_i\,\mathbf{e}_s + \overline{T}'_{l,i}\cos\alpha_i\,\mathbf{e}_\zeta\right]/\sqrt{N_c} 8) \\ &\frac{\partial}{\partial A_{\theta,o,i,l}}\int\!\!dv\;\mathbf{B}\cdot\mathbf{B} &= 2\int\!\!dv\;\mathbf{B}\cdot\frac{\partial\mathbf{B}}{\partial A_{\theta,o,i,l}} = 2\int\!\!dv\;\mathbf{B}\cdot\left[+n_i\overline{T}_{l,i}\cos\alpha_i\,\mathbf{e}_s + \overline{T}'_{l,i}\sin\alpha_i\,\mathbf{e}_\zeta\right]/\sqrt{N_c} 9) \\ &\frac{\partial}{\partial A_{\zeta,e,i,l}}\int\!\!dv\;\mathbf{B}\cdot\mathbf{B} &= 2\int\!\!dv\;\mathbf{B}\cdot\frac{\partial\mathbf{B}}{\partial A_{\zeta,e,i,l}} = 2\int\!\!dv\;\mathbf{B}\cdot\left[-m_i\overline{T}_{l,i}\sin\alpha_i\,\mathbf{e}_s - \overline{T}'_{l,i}\cos\alpha_i\,\mathbf{e}_\theta\right]/\sqrt{N_c} 9) \\ &\frac{\partial}{\partial A_{\zeta,o,i,l}}\int\!\!dv\;\mathbf{B}\cdot\mathbf{B} &= 2\int\!\!dv\;\mathbf{B}\cdot\frac{\partial\mathbf{B}}{\partial A_{\zeta,o,i,l}} = 2\int\!\!dv\;\mathbf{B}\cdot\left[+m_i\overline{T}_{l,i}\cos\alpha_i\,\mathbf{e}_s - \overline{T}'_{l,i}\sin\alpha_i\,\mathbf{e}_\theta\right]/\sqrt{N_c} 9) \end{split}$$

• The second derivatives of  $\int dv \mathbf{B} \cdot \mathbf{B}$  with respect to  $A_{\theta,e,i,l}$ ,  $A_{\theta,o,i,l}$ ,  $A_{\zeta,e,i,l}$  and  $A_{\zeta,o,i,l}$  are

• The second derivatives of 
$$\int dv \ \mathbf{B} \cdot \mathbf{B}$$
 with respect to  $A_{\theta,e,i,l}$ ,  $A_{\theta,o,i,l}$ ,  $A_{\zeta,e,i,l}$  and  $A_{\zeta,o,i,l}$  are  $\frac{\partial}{\partial A_{\theta,e,i,l}} \int dv \ \mathbf{B} \cdot \mathbf{B} = 2 \int dv \ (+n_j n_i \overline{T}_{p,j} \overline{T}_{l,i} s_j s_i g_s s_s - n_j \overline{T}_{p,j} \overline{T}_{l,i} s_j c_i g_s \zeta - n_i \overline{T}_{l,i} \overline{T}_{p,j} s_i s_j g_s \zeta + \overline{T}_{p,j} \overline{T}_{l,i} s_j c_i g_\zeta \zeta / \sqrt{2} \frac{\partial}{\partial A_{\theta,e,i,l}} \int dv \ \mathbf{B} \cdot \mathbf{B} = 2 \int dv \ (-n_j n_i \overline{T}_{p,j} \overline{T}_{l,i} c_j s_i g_s s_s + n_j \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_s \zeta - n_i \overline{T}_{l,i} \overline{T}_{p,j} s_i s_j g_s \zeta + \overline{T}_{p,j} \overline{T}_{l,i} s_j c_i g_\zeta \zeta / \sqrt{2} \frac{\partial}{\partial A_{\zeta,e,i,l}} \int dv \ \mathbf{B} \cdot \mathbf{B} = 2 \int dv \ (+m_j n_i \overline{T}_{p,j} \overline{T}_{l,i} c_j s_i g_s s_s - m_j \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_s \zeta + n_i \overline{T}_{l,i} \overline{T}_{p,j} s_i s_j g_s \theta - \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_s \zeta / \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_s \zeta + n_i \overline{T}_{l,i} \overline{T}_{p,j} s_i s_j g_s \theta - \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_s \zeta / \overline{T}_{p,j} \overline{T}_{$ 

$$\frac{\partial}{\partial A_{\theta,o,j,p}} \frac{\partial}{\partial A_{\zeta,o,i,l}} \int dv \, \mathbf{B} \cdot \mathbf{B} = 2 \int dv \, (+n_j m_i \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_{ss} - n_j \overline{T}_{p,j} \overline{T}'_{l,i} c_j s_i g_{s\theta} + m_i \overline{T}_{l,i} \overline{T}'_{p,j} c_i s_j g_{s\zeta} - \overline{T}'_{p,j} \overline{T}'_{l,i} s_j s_i g_{\theta\zeta}) / dv \, \mathbf{B} \cdot \mathbf{B} = 2 \int dv \, (-m_j m_i \overline{T}_{p,j} \overline{T}_{l,i} s_j c_i g_{ss} + m_j \overline{T}_{p,j} \overline{T}'_{l,i} s_j s_i g_{s\theta} - m_i \overline{T}_{l,i} \overline{T}'_{p,j} c_i c_j g_{s\theta} + \overline{T}'_{p,j} \overline{T}'_{l,i} c_j s_i g_{\theta\theta}) / dv \, \mathbf{B} \cdot \mathbf{B} = 2 \int dv \, (+m_j m_i \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_{ss} - m_j \overline{T}_{p,j} \overline{T}'_{l,i} c_j s_i g_{s\theta} - m_i \overline{T}_{l,i} \overline{T}'_{p,j} c_i s_j g_{s\theta} + \overline{T}'_{p,j} \overline{T}'_{l,i} s_j s_i g_{\theta\theta}) / dv \, \mathbf{B} \cdot \mathbf{B} = 2 \int dv \, (+m_j m_i \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_{ss} - m_j \overline{T}_{p,j} \overline{T}'_{l,i} c_j s_i g_{s\theta} - m_i \overline{T}_{l,i} \overline{T}'_{p,j} c_i s_j g_{s\theta} + \overline{T}'_{p,j} \overline{T}'_{l,i} s_j s_i g_{\theta\theta}) / dv \, \mathbf{B} \cdot \mathbf{B} = 2 \int dv \, (+m_j m_i \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_{ss} - m_j \overline{T}_{p,j} \overline{T}'_{l,i} c_j s_i g_{s\theta} - m_i \overline{T}_{l,i} \overline{T}'_{p,j} c_i s_j g_{s\theta} + \overline{T}'_{p,j} \overline{T}'_{l,i} s_j s_i g_{\theta\theta}) / dv \, \mathbf{B} \cdot \mathbf{B} = 2 \int dv \, (+m_j m_i \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_{ss} - m_j \overline{T}_{p,j} \overline{T}'_{l,i} c_j s_i g_{s\theta} - m_i \overline{T}_{l,i} \overline{T}'_{p,j} c_i s_j g_{s\theta} + \overline{T}'_{p,j} \overline{T}'_{l,i} s_j s_i g_{\theta\theta}) / dv \, \mathbf{B} \cdot \mathbf{B} = 2 \int dv \, (+m_j m_i \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_{ss} - m_j \overline{T}_{p,j} \overline{T}'_{l,i} c_j s_i g_{s\theta} - m_i \overline{T}_{l,i} \overline{T}'_{p,j} c_i s_j g_{s\theta} + \overline{T}'_{p,j} \overline{T}'_{l,i} s_j s_i g_{\theta\theta}) / dv \, \mathbf{B} \cdot \mathbf{B} = 2 \int dv \, (+m_j m_i \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_{ss} - m_j \overline{T}_{p,j} \overline{T}'_{l,i} c_j s_i g_{s\theta} - m_i \overline{T}_{l,i} \overline{T}'_{p,j} c_i s_j g_{s\theta} + \overline{T}'_{p,j} \overline{T}'_{l,i} s_j s_i g_{\theta\theta}) / dv \, \mathbf{B} \cdot \mathbf{B} = 2 \int dv \, (+m_j m_i \overline{T}_{p,j} \overline{T}_{l,i} c_j c_i g_{ss} - m_j \overline{T}_{p,j} \overline{T}'_{l,i} c_j s_i g_{s\theta} - m_i \overline{T}_{l,i} c_j c_i g_{s\theta} - m$$

7.11 Build matrices 51

## derivatives of helicity integrals

• The first derivatives of  $\int dv \ \mathbf{A} \cdot \mathbf{B}$  with respect to  $A_{\theta,e,i,l}, A_{\theta,o,i,l}, A_{\zeta,e,i,l}$  and  $A_{\zeta,o,i,l}$  are

$$\begin{split} &\frac{\partial}{\partial A_{\theta,e,i,l}} \int\!\! dv \; \mathbf{A} \cdot \mathbf{B} &= \int\!\! dv \; \left( \frac{\partial \mathbf{A}}{\partial A_{\theta,e,i,l}} \cdot \mathbf{B} + \mathbf{A} \cdot \frac{\partial \mathbf{B}}{\partial A_{\theta,e,i,l}} \right) = \int\!\! dv \; (\overline{T}_{l,i} \cos \alpha_i \nabla \theta \cdot \mathbf{B} + \mathbf{A} \cdot \overline{T}'_{l,i} \cos \alpha_i \, \mathbf{e}_{\zeta} / (\sqrt[3]{2})) \\ &\frac{\partial}{\partial A_{\theta,o,i,l}} \int\!\! dv \; \mathbf{A} \cdot \mathbf{B} &= \int\!\! dv \; \left( \frac{\partial \mathbf{A}}{\partial A_{\theta,o,i,l}} \cdot \mathbf{B} + \mathbf{A} \cdot \frac{\partial \mathbf{B}}{\partial A_{\theta,o,i,l}} \right) = \int\!\! dv \; (\overline{T}_{l,i} \sin \alpha_i \nabla \theta \cdot \mathbf{B} + \mathbf{A} \cdot \overline{T}'_{l,i} \sin \alpha_i \, \mathbf{e}_{\zeta} / (\sqrt[3]{2})) \\ &\frac{\partial}{\partial A_{\zeta,e,i,l}} \int\!\! dv \; \mathbf{A} \cdot \mathbf{B} &= \int\!\! dv \; \left( \frac{\partial \mathbf{A}}{\partial A_{\zeta,e,i,l}} \cdot \mathbf{B} + \mathbf{A} \cdot \frac{\partial \mathbf{B}}{\partial A_{\zeta,e,i,l}} \right) = \int\!\! dv \; (\overline{T}_{l,i} \cos \alpha_i \nabla \zeta \cdot \mathbf{B} - \mathbf{A} \cdot \overline{T}'_{l,i} \cos \alpha_i \, \mathbf{e}_{\theta} / (\sqrt[3]{2})) \\ &\frac{\partial}{\partial A_{\zeta,o,i,l}} \int\!\! dv \; \mathbf{A} \cdot \mathbf{B} &= \int\!\! dv \; \left( \frac{\partial \mathbf{A}}{\partial A_{\zeta,o,i,l}} \cdot \mathbf{B} + \mathbf{A} \cdot \frac{\partial \mathbf{B}}{\partial A_{\zeta,o,i,l}} \right) = \int\!\! dv \; (\overline{T}_{l,i} \sin \alpha_i \nabla \zeta \cdot \mathbf{B} - \mathbf{A} \cdot \overline{T}'_{l,i} \sin \alpha_i \, \mathbf{e}_{\theta} / (\sqrt[3]{2})) \\ &\frac{\partial}{\partial A_{\zeta,o,i,l}} \int\!\! dv \; \mathbf{A} \cdot \mathbf{B} &= \int\!\! dv \; \left( \frac{\partial \mathbf{A}}{\partial A_{\zeta,o,i,l}} \cdot \mathbf{B} + \mathbf{A} \cdot \frac{\partial \mathbf{B}}{\partial A_{\zeta,o,i,l}} \right) = \int\!\! dv \; (\overline{T}_{l,i} \sin \alpha_i \nabla \zeta \cdot \mathbf{B} - \mathbf{A} \cdot \overline{T}'_{l,i} \sin \alpha_i \, \mathbf{e}_{\theta} / (\sqrt[3]{2})) \\ &\frac{\partial}{\partial A_{\zeta,o,i,l}} \int\!\! dv \; \mathbf{A} \cdot \mathbf{B} &= \int\!\! dv \; \left( \frac{\partial \mathbf{A}}{\partial A_{\zeta,o,i,l}} \cdot \mathbf{B} + \mathbf{A} \cdot \frac{\partial \mathbf{B}}{\partial A_{\zeta,o,i,l}} \right) = \int\!\! dv \; (\overline{T}_{l,i} \sin \alpha_i \nabla \zeta \cdot \mathbf{B} - \mathbf{A} \cdot \overline{T}'_{l,i} \sin \alpha_i \, \mathbf{e}_{\theta} / (\sqrt[3]{2}) \right) \\ &\frac{\partial}{\partial A_{\zeta,o,i,l}} \int\!\! dv \; \mathbf{A} \cdot \mathbf{B} = \int\!\! dv \; \left( \frac{\partial \mathbf{A}}{\partial A_{\zeta,o,i,l}} \cdot \mathbf{B} + \mathbf{A} \cdot \frac{\partial \mathbf{B}}{\partial A_{\zeta,o,i,l}} \right) = \int\!\! dv \; (\overline{T}_{l,i} \sin \alpha_i \nabla \zeta \cdot \mathbf{B} - \mathbf{A} \cdot \overline{T}'_{l,i} \sin \alpha_i \, \mathbf{e}_{\theta} / (\sqrt[3]{2}) \right) \\ &\frac{\partial}{\partial A_{\zeta,o,i,l}} \int\!\! dv \; \mathbf{A} \cdot \mathbf{B} = \int\!\! dv \; \left( \frac{\partial \mathbf{A}}{\partial A_{\zeta,o,i,l}} \cdot \mathbf{B} + \mathbf{A} \cdot \frac{\partial \mathbf{B}}{\partial A_{\zeta,o,i,l}} \right) = \int\!\! dv \; (\overline{T}_{l,i} \sin \alpha_i \nabla \zeta \cdot \mathbf{B} - \mathbf{A} \cdot \overline{T}'_{l,i} \sin \alpha_i \, \mathbf{e}_{\theta} / (\sqrt[3]{2}) \right) \\ &\frac{\partial}{\partial A_{\zeta,o,i,l}} \int\!\! dv \; \mathbf{A} \cdot \mathbf{B} = \int\!\!\! dv \; \left( \frac{\partial \mathbf{A}}{\partial A_{\zeta,o,i,l}} \cdot \mathbf{B} + \mathbf{A} \cdot \frac{\partial \mathbf{B}}{\partial A_{\zeta,o,i,l}} \right) = \int\!\!\! dv \; (\overline{T}_{l,i} \cdot \mathbf{B} - \mathbf{A} \cdot \overline{T}'_{l,i} \cdot \mathbf{B} - \mathbf{A} \cdot \overline{T}'_{l,i} \cdot \mathbf{B} \right) \\ &\frac{\partial}{\partial A_{\zeta,o,i,l}} \int\!\!\!\! dv \; \mathbf{A} \cdot \mathbf{B} - \mathbf{A} \cdot \overline{T}'_{l,i} \cdot \mathbf$$

- Note that in the above expressions,  $\mathbf{A} \cdot \mathbf{e}_s = 0$  has been used.
- The second derivatives of  $\int dv \ \mathbf{A} \cdot \mathbf{B}$  with respect to  $A_{\theta,e,i,l}$ ,  $A_{\theta,o,i,l}$ ,  $A_{\zeta,e,i,l}$  and  $A_{\zeta,o,i,l}$  are

$$\begin{array}{lll} \frac{\partial}{\partial A_{\theta,e,i,l}} & \frac{\partial}{\partial A_{\theta,e,i,l}} & \int dv \ \mathbf{A} \cdot \mathbf{B} & = & \int dv \ \left[ +\overline{T}_{l,i} \cos \alpha_i \nabla \theta + \overline{T}_{p,j} \cos \alpha_j \mathbf{e}_\zeta + \overline{T}_{p,j} \cos \alpha_j \nabla \theta + \overline{T}_{l,i} \cos \alpha_i \mathbf{e}_\zeta \right] / \sqrt{39} \\ \frac{\partial}{\partial A_{\theta,o,j,p}} & \frac{\partial}{\partial A_{\theta,e,i,l}} & \int dv \ \mathbf{A} \cdot \mathbf{B} & = & \int dv \ \left[ +\overline{T}_{l,i} \cos \alpha_i \nabla \theta + \overline{T}_{p,j} \sin \alpha_j \mathbf{e}_\zeta + \overline{T}_{p,j} \sin \alpha_j \nabla \theta + \overline{T}_{l,i} \cos \alpha_i \mathbf{e}_\zeta \right] / \sqrt{39} \\ \frac{\partial}{\partial A_{\zeta,e,j,p}} & \frac{\partial}{\partial A_{\theta,e,i,l}} & \int dv \ \mathbf{A} \cdot \mathbf{B} & = & \int dv \ \left[ -\overline{T}_{l,i} \cos \alpha_i \nabla \theta + \overline{T}_{p,j} \sin \alpha_j \mathbf{e}_\theta + \overline{T}_{p,j} \sin \alpha_j \nabla \zeta + \overline{T}_{l,i} \cos \alpha_i \mathbf{e}_\zeta \right] / \sqrt{39} \\ \frac{\partial}{\partial A_{\theta,e,j,p}} & \frac{\partial}{\partial A_{\theta,e,i,l}} & \int dv \ \mathbf{A} \cdot \mathbf{B} & = & \int dv \ \left[ -\overline{T}_{l,i} \cos \alpha_i \nabla \theta + \overline{T}_{p,j} \sin \alpha_j \mathbf{e}_\theta + \overline{T}_{p,j} \sin \alpha_j \nabla \zeta + \overline{T}_{l,i} \cos \alpha_i \mathbf{e}_\zeta \right] / \sqrt{39} \\ \frac{\partial}{\partial A_{\theta,e,j,p}} & \frac{\partial}{\partial A_{\theta,e,i,l}} & \int dv \ \mathbf{A} \cdot \mathbf{B} & = & \int dv \ \left[ +\overline{T}_{l,i} \sin \alpha_i \nabla \theta + \overline{T}_{p,j} \cos \alpha_j \mathbf{e}_\zeta + \overline{T}_{p,j} \cos \alpha_j \nabla \theta + \overline{T}_{l,i} \sin \alpha_i \mathbf{e}_\zeta \right] / \sqrt{39} \\ \frac{\partial}{\partial A_{\theta,e,j,p}} & \frac{\partial}{\partial A_{\theta,e,i,l}} & \int dv \ \mathbf{A} \cdot \mathbf{B} & = & \int dv \ \left[ +\overline{T}_{l,i} \sin \alpha_i \nabla \theta + \overline{T}_{p,j} \sin \alpha_j \mathbf{e}_\zeta + \overline{T}_{p,j} \sin \alpha_j \nabla \theta + \overline{T}_{l,i} \sin \alpha_i \mathbf{e}_\zeta \right] / \sqrt{39} \\ \frac{\partial}{\partial A_{\theta,e,j,p}} & \frac{\partial}{\partial A_{\theta,e,i,l}} & \int dv \ \mathbf{A} \cdot \mathbf{B} & = & \int dv \ \left[ -\overline{T}_{l,i} \sin \alpha_i \nabla \theta + \overline{T}_{p,j} \sin \alpha_j \mathbf{e}_\zeta + \overline{T}_{p,j} \sin \alpha_j \nabla \zeta + \overline{T}_{l,i} \sin \alpha_i \mathbf{e}_\zeta \right] / \sqrt{39} \\ \frac{\partial}{\partial A_{\theta,e,j,p}} & \frac{\partial}{\partial A_{\theta,e,i,l}} & \int dv \ \mathbf{A} \cdot \mathbf{B} & = & \int dv \ \left[ -\overline{T}_{l,i} \sin \alpha_i \nabla \theta + \overline{T}_{p,j} \sin \alpha_j \mathbf{e}_\zeta + \overline{T}_{p,j} \sin \alpha_j \nabla \zeta + \overline{T}_{l,i} \sin \alpha_i \mathbf{e}_\zeta \right] / \sqrt{39} \\ \frac{\partial}{\partial A_{\theta,e,j,p}} & \frac{\partial}{\partial A_{\theta,e,i,l}} & \int dv \ \mathbf{A} \cdot \mathbf{B} & = & \int dv \ \left[ -\overline{T}_{l,i} \sin \alpha_i \nabla \theta + \overline{T}_{p,j} \sin \alpha_j \mathbf{e}_\zeta + \overline{T}_{p,j} \sin \alpha_j \nabla \zeta + \overline{T}_{l,i} \sin \alpha_i \mathbf{e}_\zeta \right] / \sqrt{39} \\ \frac{\partial}{\partial A_{\theta,e,j,p}} & \frac{\partial}{\partial A_{\theta,e,i,l}} & \int dv \ \mathbf{A} \cdot \mathbf{B} & = & \int dv \ \left[ +\overline{T}_{l,i} \cos \alpha_i \nabla \zeta + \overline{T}_{p,j} \sin \alpha_j \mathbf{e}_\zeta - \overline{T}_{p,j} \sin \alpha_j \nabla \zeta + \overline{T}_{l,i} \sin \alpha_i \mathbf{e}_\zeta \right] / \sqrt{39} \\ \frac{\partial}{\partial A_{\theta,e,j,p}} & \frac{\partial}{\partial A_{\xi,e,i,l}} & \int dv \ \mathbf{A} \cdot \mathbf{B} & = & \int dv \ \left[ +\overline{T}_{l,i} \sin \alpha_i \nabla \zeta + \overline{T}_{p,j} \sin \alpha_j \mathbf{e}_\zeta - \overline{T}_{p,j} \sin \alpha_j \nabla$$

• In these expressions the terms  $\nabla \theta \cdot \mathbf{e}_{\theta} = \nabla \zeta \cdot \mathbf{e}_{\zeta} = 1$ , and  $\nabla \theta \cdot \mathbf{e}_{\zeta} = \nabla \zeta \cdot \mathbf{e}_{\theta} = 0$  have been included to show the structure of the derivation.

## derivatives of kinetic energy integrals

• The first derivatives of  $\int dv \, v^2$  with respect to  $v_{s,e,i,l}$  etc. are

$$\frac{\partial}{\partial v_{s,e,i,l}} \int dv \, \mathbf{v} \cdot \mathbf{v} = 2 \int dv \, \mathbf{v} \cdot \overline{T}_{l,i} \cos \alpha_i \nabla s \tag{152}$$

$$\frac{\partial}{\partial v_{s,o,i,l}} \int dv \, \mathbf{v} \cdot \mathbf{v} = 2 \int dv \, \mathbf{v} \cdot \overline{T}_{l,i} \sin \alpha_i \nabla s \tag{153}$$

$$\frac{\partial}{\partial v_{\theta e i} l} \int dv \, \mathbf{v} \cdot \mathbf{v} = 2 \int dv \, \mathbf{v} \cdot \overline{T}_{l,i} \cos \alpha_i \nabla \theta$$
 (154)

$$\frac{\partial}{\partial v_{\theta,o,i,l}} \int dv \, \mathbf{v} \cdot \mathbf{v} = 2 \int dv \, \mathbf{v} \cdot \overline{T}_{l,i} \sin \alpha_i \nabla \theta$$
 (155)

$$\frac{\partial}{\partial v_{\zeta,e,i,l}} \int dv \, \mathbf{v} \cdot \mathbf{v} = 2 \int dv \, \mathbf{v} \cdot \overline{T}_{l,i} \cos \alpha_i \nabla \zeta \tag{156}$$

$$\frac{\partial}{\partial v_{\zeta,\varrho,i,l}} \int dv \, \mathbf{v} \cdot \mathbf{v} = 2 \int dv \, \mathbf{v} \cdot \overline{T}_{l,i} \sin \alpha_i \nabla \zeta \tag{157}$$

(158)

## calculation of volume-integrated basis-function-weighted metric information

• The required geometric information is calculated in ma00aa().

#### **Parameters**

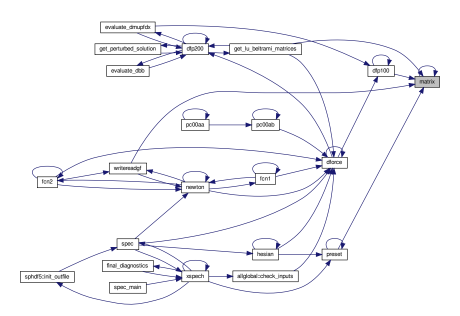
in	Ivol	
in	mn	
in	Irad	

References allglobal::ate, allglobal::ato, allglobal::aze, allglobal::aze, allglobal::azo, allglobal::dbdx, allglobal::ddttcc, allglobal::ddttcs, allglobal::ddttcs, allglobal::ddttcs, allglobal::ddttcc, allglobal::ddtzcc, allglobal::ddtzcc, allglobal::ddtzcc, allglobal::ddtzcc, allglobal::ddtzcc, allglobal::ddtzcc, allglobal::ddtzcc, allglobal::ddtzcc, allglobal::ddtzcc, allglobal::dma, allglobal::dmb, allglobal::dmd, allglobal::idnc, allglobal::ibnc, allglobal::ibnc, allglobal::ibnc, allglobal::imn, allglobal::imn, allglobal::lmc, allglobal::lmc, allglobal::lmc, allglobal::lmc, allglobal::lmc, allglobal::mpi\_comm\_spec, inputlist::mpol, allglobal::myid, allglobal::madof, allglobal::ncpu, allglobal::notstellsym, constants::one, fileunits::ounit, allglobal::rtm, allglobal::rtt, numerical::small, allglobal::tdstcc, allglobal::tdstcs, allglobal::tdstcc, allglobal::tdstcs, allglobal::tdstcc, allglobal::ttsscc, allglo



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Here is the caller graph for this function:



Constructs matrices that represent the Beltrami linear system, matrix-free.

### **Parameters**

lvol	
mn	
Irad	
resultA	
resultD	
idx	

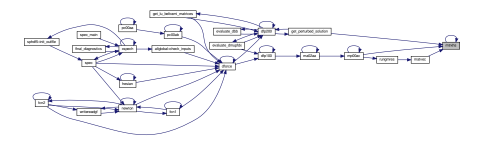
References allglobal::ate, allglobal::ate, allglobal::aze, allglobal::aze, allglobal::aze, allglobal::aze, allglobal::dtc, allglobal::dtc, allglobal::dtc, allglobal::dzc, allglobal::dzc, constants::half, allglobal::im, allglobal::in, allglobal::lcoordinatesingularity, allglobal::lma, allglobal::lmavalue, allglobal::lmbvalue, allglobal::lmc, allglobal::lmcvalue, allglobal::lmd, allglobal::lmf, allglobal::lmf, allglobal::lmf, allglobal::lmg, allglobal::lmg, allglobal::lmgvalue, allglobal::lmhvalue, allglobal::mpi\_comm\_spec, inputlist::mpol, mtrxhs(), allglobal::myid, allglobal::nadof, allglobal::ncpu, allglobal::notstellsym, constants::one, fileunits::ounit, allglobal::rtm, allglobal::rtt, numerical::small, allglobal::tsc, allglobal::tsc, allglobal::ttc, allglobal::ttc, allglobal::tts, constants::two, allglobal::tzc, allglobal::tzs, inputlist::wmacros, allglobal::yesstellsym, and constants::zero.

Referenced by get\_perturbed\_solution(), matvec(), mp00ac(), and mtrxhs().

Here is the call graph for this function:



Here is the caller graph for this function:



Constructs matrices for the precondtioner.

## Preconditioner

GMRES iteratively looks for  $\mathbf{a}_n$  that minimises the residual  $\epsilon_{\mathrm{GMRES}} = \|\hat{\mathcal{A}} \cdot \mathbf{a}_n - \mathbf{b}\|$ , where  $\|.\|$  is the Euclidean norm. Instead of solving the original problem which is usually ill-conditioned, a left preconditioner matrix  $\mathcal{M}$  is applied on both side of  $\mathcal{A} \cdot \mathbf{a} = \mathbf{b}$  so that the transformed problem is well conditioned. The convergence speed of (the preconditioned) GMRES depends highly on the quality of  $\mathcal{M}$ . A good preconditioner will require the matrix product  $\mathcal{M}^{-1}\hat{\mathcal{A}}$  to be as close as possible to an identity matrix. Also, inverting the preconditioner  $\mathcal{M}$  should be considerably cheaper than inverting  $\hat{\mathcal{A}}$  itself.

If the i-th and j-th unknowns in a correspond to  $A_{\theta,m_i,n_i,l_i}$  and  $A_{\theta,m_j,n_j,l_j}$ , respectively, then the matrix element  $\hat{\mathcal{A}}_{i,j}$  describes the coupling strength between harmonics  $(m_i,n_i)$  and  $(m_j,n_j)$ . Noting that if the Fourier series of the boundary  $R_{m,n}$  and  $Z_{m,n}$  have spectral convergence, then the coupling terms between  $A_{\theta,m_i,n_i,l_i}$  and  $A_{\theta,m_j,n_j,l_j}$ , formed by the  $(|m_i-m_j|,|n_i-n_j|)$  harmonics of the coordinate metrics, should also decay exponentially with  $|m_i-m_j|$  and  $|n_i-n_j|$  and are thus small compared to the 'diagonals"  $m_i=m_j$  and  $n_i=n_j$ . Therefore, we can construct  $\mathcal M$  from the elements of  $\hat{\mathcal A}$  by eliminating all the coupling terms with  $m_i\neq m_j$  or  $n_i\neq n_j$ , and keeping the rest ('diagonals" and terms related to Lagrange mulitpliers). Physically, the matrix  $\mathcal M$  is equivalent to the  $\hat{\mathcal A}$  matrix of a tokamak with similar major radius and minor radius to the stellarator we are solving for. The preconditioning matrix  $\mathcal M$  is sparse, with the number of nonzero elements  $\sim O(MNL^2)$ , while the total number of elements in  $\mathcal M$  is  $O(M^2N^2L^2)$ . After the construction of  $\mathcal M$ , the approximate inverse  $\mathcal M$  is computed by an incomplete LU factorisation.

This subroutine constructs such a preconditioner matrix  $\mathcal{M}$  and store it inside a sparse matrix. The matrix elements are the same as **matrix.f90**, however, only the aforementioned terms are kept. The sparse matrix uses the storage structure of **Compact Sparse Row (CSR)**.

7.11 Build matrices 55

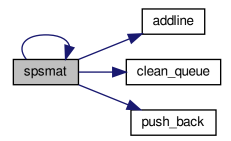
#### **Parameters**

Ivol	
mn	
Irad	

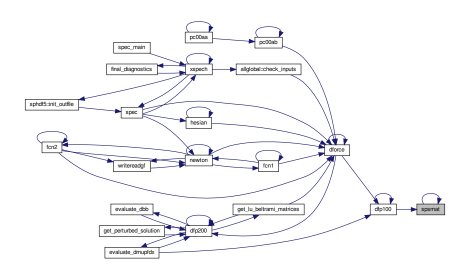
References addline(), allglobal::ate, allglobal::ate, allglobal::aze, allglobal::aze, clean\_queue(), allglobal::cpus, allglobal::ddttcc, allglobal::ddttccc, allglobal::ddtccc, allglobal::ddtccc, allglobal::ddtccc, allglobal::ddtccc, allglobal::ddtccc, allglobal::dtccc, allglobal::dtccc, allglobal::dtccc, allglobal::dtccc, allglobal::ibnc, allglobal::ibnc, allglobal::ibnc, allglobal::ibnc, allglobal::ibnc, allglobal::ilnc, allglobal::ilnc, allglobal::lmc, allglobal::lmc, allglobal::lmc, allglobal::lmc, allglobal::lmc, allglobal::lmc, allglobal::lmc, allglobal::mpi\_ccmm\_spec, inputlist::mpol, allglobal::myid, allglobal::nadof, allglobal::nccc, allglobal::ndmasmax, allglobal::ndmasmax, allglobal::notstellsym, constants::one, fileunits::ounit, push\_back(), allglobal::rtm, allglobal::rtt, numerical::small, spsmat(), allglobal::tdstcc, allglobal::tdstcc, allglobal::tdstcc, allglobal::tdstcc, allglobal::tdstcc, allglobal::tdstcc, allglobal::tdstcc, allglobal::tdstcc, allglobal::ttsscc, allglob

Referenced by dfp100(), and spsmat().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.12 Metric quantities

#### **Functions/Subroutines**

• subroutine metrix (Iquad, Ivol)

Calculates the metric quantities,  $\sqrt{g} g^{\mu\nu}$ , which are required for the energy and helicity integrals.

## 7.12.1 Detailed Description

### 7.12.2 Function/Subroutine Documentation

Calculates the metric quantities,  $\sqrt{g}\,g^{\mu\nu}$ , which are required for the energy and helicity integrals. **metrics** 

• The Jacobian,  $\sqrt{g}$ , and the "lower" metric elements,  $g_{\mu\nu}$ , are calculated by coords(), and are provided on a regular grid in "real-space", i.e.  $(\theta, \zeta)$ , at a given radial location, i.e. where s is input.

## plasma region

• In the plasma region, the required terms are  $\bar{g}_{\mu\nu} \equiv g_{\mu\nu}/\sqrt{g}$ .

$$\sqrt{g} g^{ss} = (g_{\theta\theta}g_{\zeta\zeta} - g_{\theta\zeta}g_{\theta\zeta})/\sqrt{g} 
\sqrt{g} g^{s\theta} = (g_{\theta\zeta}g_{s\zeta} - g_{s\theta}g_{\zeta\zeta})/\sqrt{g} 
\sqrt{g} g^{s\zeta} = (g_{s\theta}g_{\theta\zeta} - g_{\theta\theta}g_{s\zeta})/\sqrt{g} 
\sqrt{g} g^{\theta\theta} = (g_{\zeta\zeta}g_{ss} - g_{s\zeta}g_{s\zeta})/\sqrt{g} 
\sqrt{g} g^{\theta\zeta} = (g_{s\zeta}g_{s\theta} - g_{\theta\zeta}g_{ss})/\sqrt{g} 
\sqrt{g} g^{\zeta\zeta} = (g_{ss}g_{\theta\theta} - g_{s\theta}g_{s\theta})/\sqrt{g}$$
(159)

## **FFTs**

• After constructing the required quantities in real space, FFTs provided the required Fourier harmonics, which are returned through global.f90. (The "extended" Fourier resolution is used.)

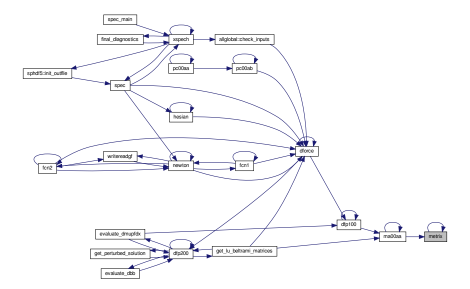
References allglobal::cfmn, allglobal::cpus, allglobal::dbdx, allglobal::efmn, allglobal::goomne, allglobal::goomne, allglobal::goomne, allglobal::gssmne, allglobal::gssmne, allglobal::gssmne, allglobal::gssmne, allglobal::gszmne, allglobal::gszmne, allglobal::gszmne, allglobal::gtzmne, allglobal::gtzmne, allglobal::gtzmne, allglobal::guvij, allglobal::guvijsave, allglobal::gyzuj, allglobal::gzzmne, allglobal::gzzmne, allglobal::im, allglobal::im, allglobal::im, allglobal::in, allglobal::ncpu, allglobal::ntz, allglobal::ntz, allglobal::ofmn, constants::one, fileunits::ounit, allglobal::sfmn, allglobal::sg, numerical::small, and constants::zero.

Referenced by ma00aa(), and metrix().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.13 Solver for Beltrami (linear) system

## **Functions/Subroutines**

subroutine mp00ac (Ndof, Xdof, Fdof, Ddof, Ldfjac, iflag)
 Solves Beltrami/vacuum (linear) system, given matrices.
 unpacking fluxes, helicity multiplier

## 7.13.1 Detailed Description

## 7.13.2 Function/Subroutine Documentation

Solves Beltrami/vacuum (linear) system, given matrices.

### unpacking fluxes, helicity multiplier

• The vector of "parameters",  $\mu$ , is unpacked. (Recall that  $\mu$  was "packed" in ma02aa() .) In the following,  $\psi \equiv (\Delta \psi_t, \Delta \psi_p)^T$ .

## construction of linear system

- The equation  $\nabla imes {f B} = \mu {f B}$  is cast as a matrix equation,

$$\mathcal{M} \cdot \mathbf{a} = \mathcal{R},\tag{160}$$

where  $\mathbf{a}$  represents the degrees-of-freedom in the magnetic vector potential,  $\mathbf{a} \equiv \{A_{\theta,e,i,l}, A_{\zeta,e,i,l}, \dots\}$ .

• The matrix  $\mathcal{M}$  is constructed from  $\mathcal{A} \equiv \text{dMA}$  and  $\mathcal{D} \equiv \text{dMD}$ , which were constructed in matrix(), according to

$$\mathcal{M} \equiv \mathcal{A} - \mu \mathcal{D}. \tag{161}$$

Note that in the vacuum region,  $\mu=0$ , so  $\mathcal{M}$  reduces to  $\mathcal{M}\equiv\mathcal{A}$ .

- The construction of the vector  $\mathcal{R}$  is as follows:
  - if Lcoordinatesingularity=T, then

$$\mathcal{R} \equiv -\left(\mathcal{B} - \mu \mathcal{E}\right) \cdot \boldsymbol{\psi} \tag{162}$$

- if Lcoordinatesingularity=F and Lplasmaregion=T, then

$$\mathcal{R} \equiv -\mathcal{B} \cdot \psi \tag{163}$$

- if Lcoordinatesingularity=F and Lvacuumregion=T, then

$$\mathcal{R} \equiv -\mathcal{G} - \mathcal{B} \cdot \psi \tag{164}$$

The quantities  $\mathcal{B}\equiv \mathtt{dMB},\,\mathcal{E}\equiv \mathtt{dME}$  and  $\mathcal{G}\equiv \mathtt{dMG}$  are constructed in matrix() .

## solving linear system

It is *not* assumed that the linear system is positive definite. The LAPACK routine DSYSVX is used to solve the linear system.

### unpacking, ...

- The magnetic degrees-of-freedom are unpacked by packab() .
- The error flag, ImagneticOK, is set that indicates if the Beltrami fields were successfully constructed.

## construction of "constraint" function

• The construction of the function  $f(\mu)$  is required so that iterative methods can be used to construct the Beltrami field consistent with the required constraints (e.g. on the enclosed fluxes, helicity, rotational-transform, ...).

#### See also

ma02aa() for additional details.

## plasma region

- For Lcoordinatesingularity=T, the returned function is:

$$\mathbf{f}(\mu,\Delta\psi_p) \equiv \begin{cases} & ( & 0 & , & 0 \ )^T, & \text{if Lconstraint} = -1 \\ & ( & 0 & , & 0 \ )^T, & \text{if Lconstraint} = 0 \\ & ( & \pm(+1)-\text{iota (lvol )} & , & 0 \ )^T, & \text{if Lconstraint} = 1 \\ & ( & ? & , & ? \ )^T, & \text{if Lconstraint} = 2 \end{cases}$$
 (165)

- For Lcoordinatesingularity=F, the returned function is:

$$\mathbf{f}(\mu,\Delta\psi_p) \equiv \left\{ \begin{array}{lll} (&0&,&0&\\ (&0&,&0&\\ (&\pm(-1)-\mathrm{oita}\,(\mathrm{lvol-1})&,&\pm(+1)-\mathrm{iota}\,(\mathrm{lvol})&)^T,&\mathrm{if}\,\,\mathrm{Lconstraint}&=&-1\\ (&?&,&?&\end{array} \right.$$

### vacuum region

- For the vacuum region, the returned function is:

$$\mathbf{f}(\Delta\psi_t,\Delta\psi_p) \equiv \begin{cases} ( & 0 & , & 0 & )^T, & \text{if Lconstraint} = -1\\ ( & I-\text{curtor} & , & G-\text{curpol} & )^T, & \text{if Lconstraint} = & 0\\ ( & \pounds(-1)-\text{oita(lvol-1)} & , & G-\text{curpol} & )^T, & \text{if Lconstraint} = & 1\\ ( & ? & , & ? & )^T, & \text{if Lconstraint} = & 2 \end{cases}$$
(167)

• The rotational-transform,  $\pm$ , is computed by tr00ab(); and the enclosed currents, I and G, are computed by curent().

## early termination

• If  $|\mathbf{f}| < \text{mupftol}$ , then early termination is enforced (i.e., iflag is set to a negative integer). (See ma02aa() for details of how mp00ac() is called iteratively.)

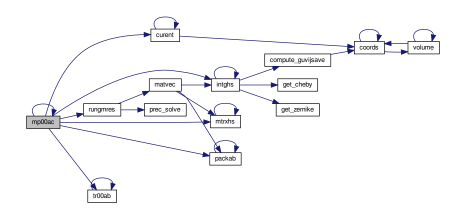
#### **Parameters**

in	Ndof	
in	Xdof	
	Fdof	
	Ddof	
in	Ldfjac	
	iflag	indicates whether (i) iflag=1: "function" values are required; or (ii) iflag=2: "derivative" values are required

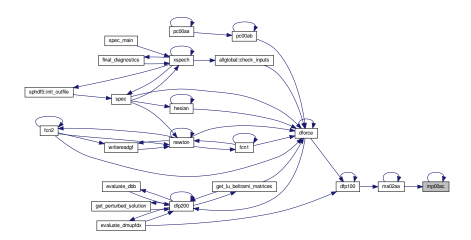
References allglobal::adotx, allglobal::ate, allglobal::ato, allglobal::aze, allglobal::azo, allglobal::cpus, curent(), inputlist::currol, inputlist::curror, allglobal::dotx, allglobal::ditotadxup, allglobal::ditgpdxtp, allglobal::dma, allglobal::dmas, allglobal::dmb, allglobal::dmd, allglobal::dmds, allglobal::dmg, allglobal::dpflux, allglobal::dtflux, inputlist::epsgmres, inputlist::epsilu, allglobal::gmreslastsolution, constants::half, inputlist::helicity, allglobal::idmas, allglobal::imagneticok, allglobal::in, intghs(), inputlist::iota, allglobal::iquad, allglobal::ivol, allglobal::jdmas, allglobal::labintegral, allglobal::lbbintegral, inputlist::lconstraint, allglobal::lcoordinatesingularity, inputlist::lgmresprec, allglobal::liluprecond, inputlist::lmatsolver, allglobal::plasmaregion, inputlist::lrad, allglobal::lvacuumregion, numerical::machprec, allglobal::mn, allglobal::mns, mp00ac(), allglobal::mpi\_comm\_spec, mtrxhs(), inputlist::mu, inputlist::mupftol, allglobal::myid, allglobal::nadof, allglobal::ncpu, allglobal::ndmas, allglobal::ndmasmax, inputlist::nitergmres, allglobal::notmatrixfree, allglobal::notstellsym, allglobal::nt, inputlist::ntor, allglobal::nz, inputlist::oita, constants::one, fileunits::ounit, packab(), rungmres(), numerical::small, allglobal::solution, tr00ab(), inputlist::wmacros, allglobal::xoffset, allglobal::yesstellsym, and constants::zero.

Referenced by ma02aa(), and mp00ac().

Here is the call graph for this function:



Here is the caller graph for this function:



## 7.14 Force-driver

#### **Functions/Subroutines**

- subroutine newton (NGdof, position, ihybrd) Employs Newton method to find  $\mathbf{F}(\mathbf{x}) = 0$ , where  $\mathbf{x} \equiv \{\text{geometry}\}\$ and  $\mathbf{F}$  is defined in dforce() .
- subroutine writereadgf (readorwrite, NGdof, ireadhessian)

read or write force-derivative matrix

- subroutine fcn1 (NGdof, xx, fvec, irevcm)
- subroutine fcn2 (NGdof, xx, fvec, fjac, Ldfjac, irevcm)

## 7.14.1 Detailed Description

## 7.14.2 Function/Subroutine Documentation

- The iterative, Newton search to find  $\mathbf{x} \equiv \{\text{geometry}\} \equiv \{R_{i,l}, Z_{i,l}\}$  such that  $\mathbf{F}(\mathbf{x}) = 0$ , where  $\mathbf{F}$  and its derivatives,  $\nabla_{\mathbf{x}}\mathbf{F}$ , are calculated by dforce(), is provided by either
  - C05NDF if Lfindzero=1, which only uses function values; or
  - C05PDF if Lfindzero=2, which uses user-provided derivatives.
- The iterative search will terminate when the solution is within c05xtol of the true solution (see NAG documentation).
- The input variable c05factor is provided to determine the initial step bound (see NAG documentation).

## logic, writing/reading from file

7.14 Force-driver 61

• Before proceeding with iterative search, dforce() is called to determine the magnitude of the initial force imbalance, and if this is less than forcetol then the iterative search will not be performed.

- As the iterations proceed, wrtend() will be called to save itermediate information (also see xspech()).
- If the derivative matrix,  $\nabla_{\mathbf{x}} \mathbf{F}$ , is required, i.e. if Lfindzero=2, and if LreadGF=T then the derivative matrix will initially be read from .ext.sp.DF, if it exists, or from .sp.DF.
- · As the iterations proceed, the derivative matrix will be written to .ext.sp.DF .

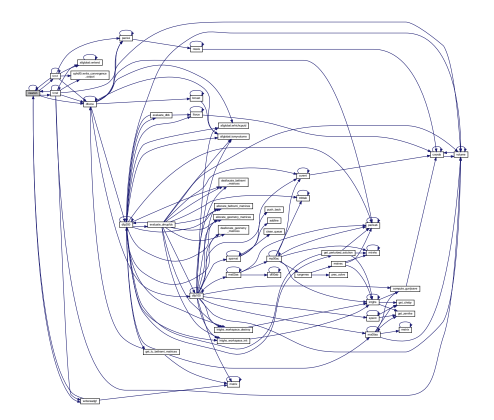
#### **Parameters**

in	NGdof	
in,out	position	
out	ihybrd	

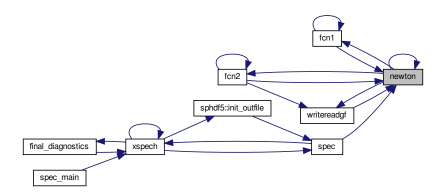
References allglobal::bbe, allglobal::bbo, inputlist::c05factor, inputlist::c05xmax, inputlist::c05xtol, allglobal::cpus, allglobal::dbbdmp, allglobal::dessian, allglobal::dffdrz, dforce(), allglobal::dmupfdx, allglobal::energy, fcn1(), fcn2(), allglobal::forceerr, inputlist::forcetol, allglobal::hessian, inputlist::igeometry, allglobal::iie, allglobal::iio, allglobal::im, allglobal::irbc, allglobal::irbs, allglobal::izbc, allglobal::izbs, newtontime::lastcpu, inputlist::lcheck, inputlist::lfindzero, allglobal::lgdof, allglobal::lhessianallocated, allglobal::localconstraint, inputlist::lreadgf, allglobal::mn, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ncpu, newtontime::ndcalls, newton(), newtontime::nfcalls, allglobal::nfreeboundaryiterations, allglobal::notstellsym, inputlist::nvol, constants::one, fileunits::ounit, numerical::sqrtmachprec, constants::ten, constants::two, inputlist::wmacros, writereadgf(), and constants::zero.

Referenced by fcn1(), fcn2(), newton(), spec(), and writereadgf().

Here is the call graph for this function:



Here is the caller graph for this function:



read or write force-derivative matrix

## **Parameters**

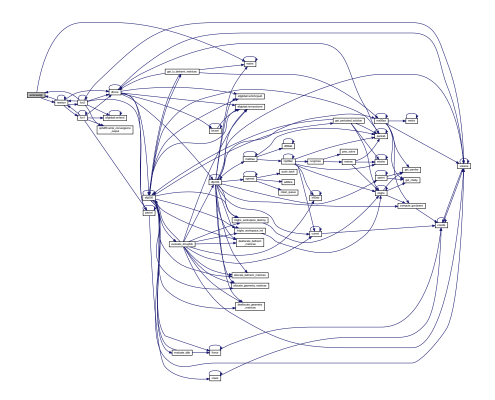
in	readorwrite	
in	NGdof	
out	ireadhessian	

References allglobal::cpus, fileunits::dunit, allglobal::hessian, inputlist::igeometry, allglobal::im, allglobal::in, inputlist::istellsym, inputlist::Ifreebound, allglobal::lhessianallocated, matrix(), allglobal::mn, allglobal::mpi\_comm\_spec, inputlist::mpol, allglobal::myid, newton(), inputlist::ntor, inputlist::nvol, fileunits::ounit, and constants::zero.

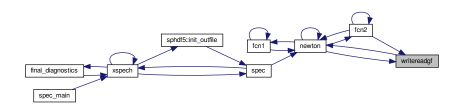
Referenced by fcn2(), and newton().

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Here is the call graph for this function:



Here is the caller graph for this function:



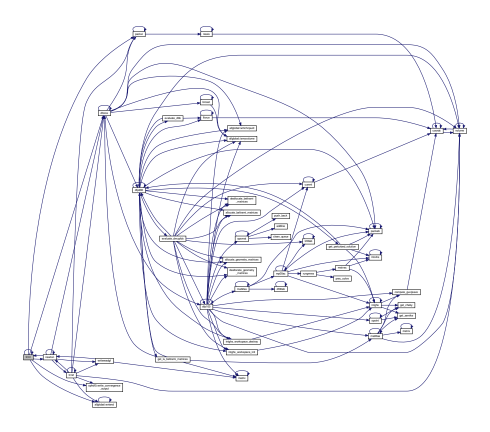
## **Parameters**

in	NGdof	
in	XX	
out	fvec	
in	irevcm	

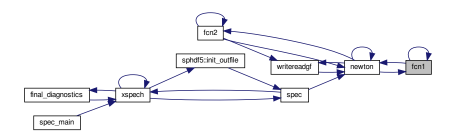
References allglobal::bbe, allglobal::bbo, inputlist::c05factor, inputlist::c05xmax, inputlist::c05xtol, allglobal::cpus,

allglobal::dbbdmp, allglobal::dessian, allglobal::dffdrz, dforce(), allglobal::dmupfdx, allglobal::energy, fcn1(), allglobal::forceerr, inputlist::forcetol, allglobal::hessian, inputlist::igeometry, allglobal::iie, allglobal::iio, allglobal::im, allglobal::irbc, allglobal::irbs, allglobal::izbc, allglobal::izbs, newtontime::lastcpu, inputlist::lcheck, inputlist::lfindzero, allglobal::lgdof, allglobal::lhessianallocated, inputlist::lreadgf, allglobal::mn, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ncpu, newtontime::ndcalls, newton(), newtontime::nfcalls, allglobal::nfreeboundaryiterations, allglobal::notstellsym, inputlist::nvol, constants::one, fileunits::ounit, packxi(), numerical::sqrtmachprec, constants::ten, constants::two, inputlist::wmacros, sphdf5::write\_convergence\_output(), allglobal::wrtend(), and constants::zero. Referenced by fcn1(), and newton().

Here is the call graph for this function:



Here is the caller graph for this function:



7.14 Force-driver 65

```
real, dimension(1:ldfjac,1:ngdof), intent(out) fjac,
integer, intent(in) Ldfjac,
integer, intent(in) irevcm )
```

fcn2

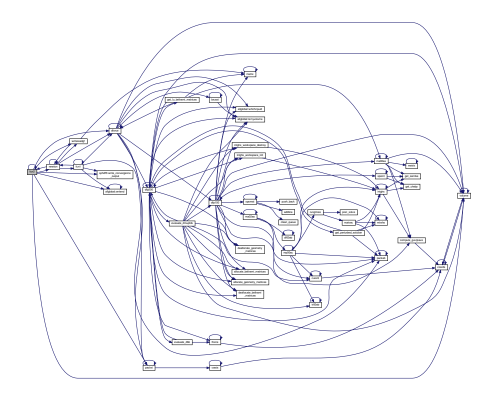
#### **Parameters**

in	NGdof	
in	XX	
out	fvec	
out	fjac	
in	Ldfjac	
in	irevcm	

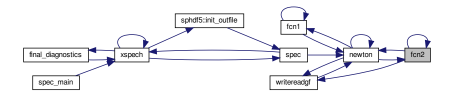
References allglobal::bbe, allglobal::bbo, inputlist::c05factor, inputlist::c05xmax, inputlist::c05xtol, allglobal::cpus, allglobal::dbbdmp, allglobal::dessian, allglobal::dffdrz, dforce(), allglobal::dmupfdx, allglobal::energy, fcn2(), allglobal::forceerr, inputlist::forcetol, allglobal::hessian, inputlist::igeometry, allglobal::iie, allglobal::iio, allglobal::im, allglobal::irbc, allglobal::irbs, allglobal::izbc, allglobal::izbs, newtontime::lastcpu, inputlist::lcheck, inputlist::lfindzero, allglobal::lgdof, allglobal::lhessianallocated, inputlist::lreadgf, allglobal::mn, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ncpu, newtontime::ndcalls, newton(), newtontime::nfcalls, allglobal::nfreeboundaryiterations, allglobal::notstellsym, inputlist::nvol, constants::one, fileunits::ounit, packxi(), numerical::sqrtmachprec, constants::ten, constants::two, volume(), inputlist::wmacros, sphdf5::write\_convergence\_output(), writereadgf(), allglobal::wrtend(), and constants::zero.

Referenced by fcn2(), and newton().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.15 "packing" of Beltrami field solution vector

# **Functions/Subroutines**

• subroutine packab (packorunpack, Ivol, NN, solution, ideriv)

Packs and unpacks Beltrami field solution vector.

 subroutine packxi (NGdof, position, Mvol, mn, iRbc, iZbs, iRbs, iZbc, packorunpack, LComputeDerivatives, LComputeAxis)

Packs, and unpacks, geometrical degrees of freedom; and sets coordinate axis.

# 7.15.1 Detailed Description

#### 7.15.2 Function/Subroutine Documentation

Packs and unpacks Beltrami field solution vector.

## construction of "vector" of independent degrees of freedom

- Numerical routines for solving linear equations typically require the unknown, independent degrees of freedom to be "packed" into a vector,  $\mathbf{x}$ .
- The magnetic field is defined by the independent degrees of freedom in the Chebyshev-Fourier representation of the vector potential,  $A_{\theta,e,i,l}$  and  $A_{\zeta,e,i,l}$ ; and the non-stellarator-symmetric terms if relevant,  $A_{\theta,o,i,l}$  and  $A_{\zeta,o,i,l}$ ; and the Lagrange multipliers,  $a_i,b_i,c_i,d_i,e_i$ , etc. as required to enforce the constraints:

$$\mathbf{x} \equiv \{ A_{\theta,e,i,l}, A_{\zeta,e,i,l}, A_{\theta,o,i,l}, A_{\zeta,o,i,l}, a_i, b_i, c_i, d_i, e_i, f_i, g_1, h_1 \}.$$
(168)

• The "packing" index is assigned in preset().

#### **Parameters**

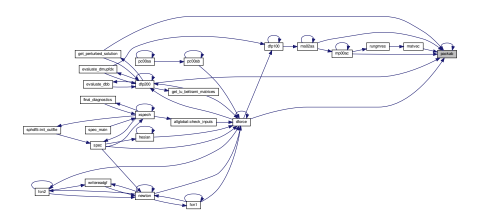
References allglobal::ate, allglobal::ate, allglobal::aze, allglobal::aze, allglobal::aze, allglobal::im, allglobal:im,

allglobal::lma, allglobal::lmavalue, allglobal::lmb, allglobal::lmbvalue, allglobal::lmc, allglobal::lmcvalue, allglobal::lmdvalue, allglobal::lmg, allglobal::myid, allglobal::notstellsym, fileunits::ounit, packab(), numerical::small, allglobal::tt, allglobal::yesstellsym, and constants::zero.

Referenced by dforce(), dfp200(), get\_perturbed\_solution(), ma02aa(), matvec(), mp00ac(), and packab(). Here is the call graph for this function:



Here is the caller graph for this function:



```
7.15.2.2 packxi() subroutine packxi (
    integer, intent(in) NGdof,
    real, dimension(0:ngdof) position,
    integer, intent(in) Mvol,
    integer, intent(in) mn,
    real, dimension(1:mn,0:mvol) iRbc,
    real, dimension(1:mn,0:mvol) iRbs,
    real, dimension(1:mn,0:mvol) iZbs,
    real, dimension(1:mn,0:mvol) iZbc,
    character packorunpack,
    logical, intent(in) LComputeDerivatives,
    logical, intent(in) LComputeAxis)
```

Packs, and unpacks, geometrical degrees of freedom; and sets coordinate axis. geometrical degrees of freedom

• The geometrical degrees-of-freedom, namely the  $R_{j,v}$  and  $Z_{j,v}$  where v labels the interface and j labels the Fourier harmonic, must be "packxi", and "unpackxi", into a single vector,  $\boldsymbol{\xi}$ , so that standard numerical routines can be called to find solutions to force-balance, i.e.  $\mathbf{F}[\boldsymbol{\xi}] = 0$ .

• A coordinate "pre-conditioning" factor is included:

$$\boldsymbol{\xi}_k \equiv \frac{R_{j,v}}{\Psi_{j,v}},\tag{169}$$

where  $\Psi_{j,v} \equiv \texttt{psifactor}(\texttt{j,v})$  , which is defined in global.f90 .

#### coordinate axis

- The coordinate axis is not an independent degree-of-freedom of the geometry. It is constructed by extrapolating the geometry of the innermost interface down to a line.
- Note that if the coordinate axis depends only on the geometry of the innermost interface then the block tridiagonal structure of the the force-derivative matrix is preserved.
- · Define the arc-length weighted averages,

$$R_0(\zeta) \equiv \frac{\int_0^{2\pi} R_1(\theta, \zeta) dl}{L(\zeta)}, \qquad Z_0(\zeta) \equiv \frac{\int_0^{2\pi} Z_1(\theta, \zeta) dl}{L(\zeta)}, \tag{170}$$

where  $L(\zeta) \equiv \int_0^{2\pi} dl$  and  $dl \equiv \sqrt{\partial_\theta R_1(\theta,\zeta)^2 + \partial_\theta Z_1(\theta,\zeta)^2} \, d\theta$ .

- Note that if dl does not depend on  $\theta$ , i.e. if  $\theta$  is the equal arc-length angle, then the expressions simplify.
- Note that the geometry of the coordinate axis thus constructed only depends on the geometry of the innermost interface, by which I mean that the geometry of the coordinate axis is independent of the angle parameterization.

#### some numerical comments

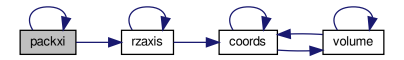
- First, the differential poloidal length,  $dl \equiv \sqrt{R_{\theta}^2 + Z_{\theta}^2}$ , is computed in real space using an inverse FFT from the Fourier harmonics of R and Z.
- Second, the Fourier harmonics of the dl are computed using an FFT. The integration over  $\theta$  to construct  $L \equiv \int dl$  is now trivial: just multiply the m=0 harmonics of dl by  $2\pi$ . The ajk (1:mn) variable is used.
- Next, the weighted  $R\ dl$  and  $Z\ dl$  are computed in real space, and the poloidal integral is similarly taken.
- · Lastly, the Fourier harmonics are constructed using an FFT after dividing in real space.

#### **Parameters**

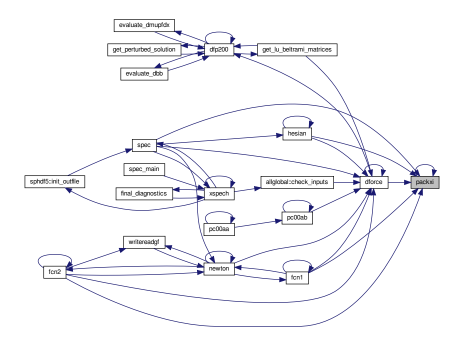
in	NGdof	
	position	
in	Mvol	
in	mn	
	iRbc	
	iZbs	
	iRbs	
	iZbc	
	packorunpack	
in	LComputeDerivatives	
in	LComputeAxis	

References allglobal::ajk, allglobal::cfmn, allglobal::comn, allglobal::cpus, allglobal::efmn, allglobal::evmn, inputlist::igeometry, allglobal::ijimag, allglobal::ijireal, allglobal::im, allglobal::irij, allglobal::irij, allglobal::irij, allglobal::irij, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ncpu, allglobal::notstellsym, allglobal::nt, inputlist::ntor, allglobal::ntz, inputlist::nvol, allglobal::nz, allglobal::odmn, allglobal::ofmn, fileunits::ounit, packxi(), allglobal::psifactor, allglobal::rscale, rzaxis(), allglobal::sfmn, allglobal::simn, allglobal::tzij, allglobal::yesstellsym, and constants::zero.

Referenced by dforce(), fcn1(), fcn2(), hesian(), packxi(), and spec(). Here is the call graph for this function:



Here is the caller graph for this function:



# 7.16 Conjugate-Gradient method

# Functions/Subroutines

- subroutine pc00aa (NGdof, position, Nvol, mn, ie04dgf)
   Use preconditioned conjugate gradient method to find minimum of energy functional.
- subroutine pc00ab (mode, NGdof, Position, Energy, Gradient, nstate, iuser, ruser)

  Returns the energy functional and it's derivatives with respect to geometry.

# 7.16.1 Detailed Description

## 7.16.2 Function/Subroutine Documentation

```
integer, intent(in) Nvol,
integer, intent(in) mn,
integer ie04dgf)
```

Use preconditioned conjugate gradient method to find minimum of energy functional.

# energy functional

The energy functional is described in pc00ab().

# relevant input variables

- The following input variables control the operation of E04DGF:
  - epsilon : weighting of "spectral energy"; see pc00ab()
  - maxstep: this is given to E04DGF for the Maximum Step Length
  - maxiter: upper limit on derivative calculations used in the conjugate gradient iterations
  - verify: if verify=1, then E04DGF will confirm user supplied gradients (provided by pc00ab()) are correct;
- Todo Unfortunately,  $\verb"E04DGF"$  seems to require approximately 3N function evaluations before proceeding to minimize the energy functional, where there are N degrees of freedom. I don't know how to turn this off!

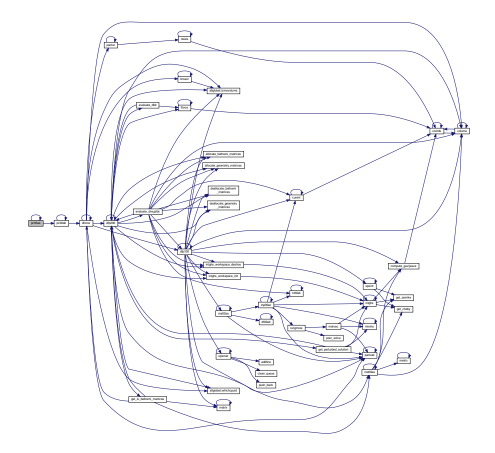
#### **Parameters**

in	NGdof
in,out	position
in	Nvol
in	mn
	ie04dgf

References allglobal::cpus, allglobal::energy, allglobal::forceerr, inputlist::forcetol, allglobal::myid, allglobal::ncpu, fileunits::ounit, pc00aa(), pc00ab(), constants::ten, and constants::zero.

Referenced by pc00aa().

Here is the call graph for this function:



Here is the caller graph for this function:



```
7.16.2.2 pc00ab() subroutine pc00ab (
    integer mode,
    integer NGdof,
    real, dimension(1:ngdof) Position,
    real Energy,
    real, dimension(1:ngdof) Gradient,
    integer nstate,
    integer, dimension(1:2) iuser,
    real, dimension(1:1) ruser )
```

Returns the energy functional and it's derivatives with respect to geometry.

# **Energy functional**

· The energy functional is

$$F \equiv \sum_{l=1}^{N} \int_{\mathcal{V}} \left( \frac{p}{\gamma - 1} + \frac{B^2}{2} \right) dv, \tag{171}$$

where  $N \equiv N \text{ vol}$  is the number of interfaces.

- Assuming that the toroidal and poloidal fluxes,  $\psi_t$  and  $\psi_p$ , the helicity,  $\mathcal{K}$ , the helicity multiplier,  $\mu$ , and/or the interface rotational-transforms,  $\pm$ , are appropriately constrained, the Beltrami fields in each volume depend only the geometry of the adjacent interfaces. So, the energy functional is assumed to be a function of "position", i.e.  $F = F(R_{l,j}, Z_{l,j})$ .
- Introducing a ficitious time, t, the position may be advanced according to

$$\frac{\partial R_{j}}{\partial t} \equiv -\frac{\partial}{\partial R_{j}} \sum_{l=1}^{N} \int \left(\frac{p}{\gamma - 1} + \frac{B^{2}}{2}\right) dv, 
\frac{\partial Z_{j}}{\partial t} \equiv -\frac{\partial}{\partial Z_{j}} \sum_{l=1}^{N} \int \left(\frac{p}{\gamma - 1} + \frac{B^{2}}{2}\right) dv.$$
(172)

· There remain degrees of freedom in the angle representation of the interfaces.

# Spectral energy minimization

· Consider variations which do not affect the geometry of the surfaces,

$$\delta R = R_{\theta} u, \tag{173}$$

$$\delta Z = Z_{\theta} u, \tag{174}$$

where u is a angle variation.

· The corresponding variation in each of the Fourier harmonics is

$$\delta R_j \equiv \oint \!\! \! \int \!\! d\theta d\zeta \ R_\theta \ u \ \cos \alpha_j, \tag{175}$$

$$\delta Z_j \equiv \oint \!\! \oint \! d\theta d\zeta \ Z_\theta \ u \ \sin \alpha_j, \tag{176}$$

· Following Hirshman et al., introducing the normalized spectral width

$$M \equiv \frac{\sum_{j} (m_{j}^{p} + n_{j}^{q}) (R_{l,j}^{2} + Z_{l,j}^{2})}{\sum_{j} (R_{l,j}^{2} + Z_{l,j}^{2})},$$
(177)

· Using the notation

$$N \equiv \sum_{j} \lambda_{j} (R_{l,j}^{2} + Z_{l,j}^{2}),$$
 (178)

$$D \equiv \sum_{i} (R_{l,j}^2 + Z_{l,j}^2), \tag{179}$$

where  $\lambda_j \equiv m_i^p + n_i^q$ , the variation in the normalized spectral width is

$$\delta M = (\delta N - M\delta D)/D. \tag{180}$$

· For tangential variations,

$$\delta N = 2 \oint \!\! \int \!\! d\theta d\zeta \ u \left( R_{\theta} \sum_{j} \lambda_{j} R_{j} \cos \alpha_{j} + Z_{\theta} \sum_{j} \lambda_{j} Z_{j} \sin \alpha_{j} \right), \tag{181}$$

$$\delta D = 2 \oint \!\! \int \!\! d\theta d\zeta \ u \left( R_{\theta} \sum_{j} R_{j} \cos \alpha_{j} + Z_{\theta} \sum_{j} Z_{j} \sin \alpha_{j} \right). \tag{182}$$

· The "tangential spectral-width descent direction" is thus

$$\frac{\partial u}{\partial t} = -\left[R_{\theta} \sum_{j} (\lambda_{j} - M) R_{j} \cos \alpha_{j} / D + Z_{\theta} \sum_{j} (\lambda_{j} - M) Z_{j} \sin \alpha_{j} / D\right]. \tag{183}$$

· This suggests that position should be advanced according to

$$\frac{\partial R_j}{\partial t} \equiv -\frac{\partial}{\partial R_j} \sum_{l=1}^N \int \left( \frac{p}{\gamma - 1} + \frac{B^2}{2} \right) dv - [R_\theta (R_\theta X + Z_\theta Y)]_j, \tag{184}$$

$$\frac{\partial Z_j}{\partial t} \equiv -\frac{\partial}{\partial Z_j} \sum_{l=1}^N \int \left(\frac{p}{\gamma - 1} + \frac{B^2}{2}\right) dv - [Z_{\theta}(R_{\theta}X + Z_{\theta}Y)]_j, \tag{185}$$

where  $X \equiv \sum_{j} (\lambda_j - M) R_j \cos \alpha_j / D$  and  $Y \equiv \sum_{j} (\lambda_j - M) Z_j \sin \alpha_j / D$ .

# numerical implementation

· The spectral condensation terms,

$$R_{ heta}(R_{ heta}X+Z_{ heta}Y) = \sum_{j,k,l} m_j m_k (\lambda_l - M) R_j (+R_k R_l \sin lpha_j \sin lpha_k \cos lpha_l - Z_k Z_l \sin lpha_j \cos lpha_k \sin lpha_l)$$
 (28) 
$$Z_{ heta}(R_{ heta}X+Z_{ heta}Y) = \sum_{j,k,l} m_j m_k (\lambda_l - M) Z_j (-R_k R_l \cos lpha_j \sin lpha_k \cos lpha_l + Z_k Z_l \cos lpha_j \cos lpha_k \sin lpha_l)$$

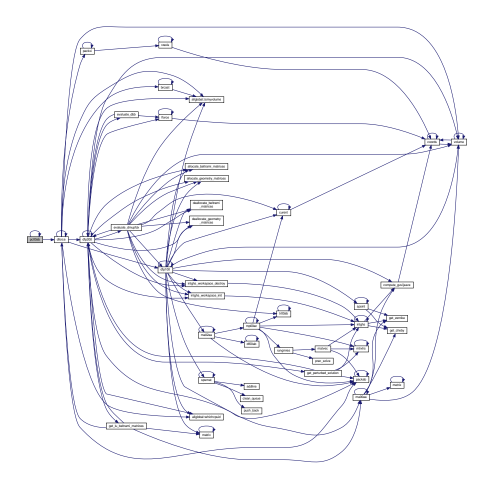
are calculated using triple angle expressions...

#### Todo IT IS VERY LIKELY THAT FFTs WOULD BE FASTER!!!

References allglobal::cpus, allglobal::dbbdrz, dforce(), allglobal::diidrz, inputlist::epsilon, allglobal::forceerr, inputlist::forcetol, constants::half, inputlist::igeometry, allglobal::lbbintegral, allglobal::mn, allglobal::myid, inputlist::nvol, constants::one, fileunits::ounit, pc00ab(), allglobal::yesstellsym, and constants::zero.

Referenced by pc00aa(), and pc00ab().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.17 Initialization of the code

# **Functions/Subroutines**

subroutine preset

Allocates and initializes internal arrays.

# 7.17.1 Detailed Description

# 7.17.2 Function/Subroutine Documentation

#### 7.17.2.1 preset() subroutine preset

Allocates and initializes internal arrays.

## LGdof and NGdof: number of geometrical degrees-of-freedom

- LGdof 

   the number of degrees-of-freedom in the geometry (i.e. Fourier harmonics) of each interface
- $NGdof \equiv total$  number of degrees-of-freedom in geometry, i.e. of all interfaces

#### iota and oita: rotational transform on interfaces

- The input variables iota and oita are the rotational transform on "inner-side" and on the "outer-side" of each interface.
- · These quantities are formally inputs.
- Note that if  $q_l + \gamma q_r \neq 0$ , then iota is given by

$$\pm \equiv \frac{p_l + \gamma p_r}{q_l + \gamma q_r},$$
(188)

where  $p_l \equiv \texttt{pl}$ ,  $q_l \equiv \texttt{ql}$  , etc.; and similarly for oita .

## dtflux(1:Mvol) and dpflux(1:Mvol): enclosed fluxes

- dtflux  $\equiv \Delta \psi_{tor}/2\pi$  and dpflux  $\equiv \Delta \psi_{pol}/2\pi$  in each volume.
- Note that the total toroidal flux enclosed by the plasma boundary is  $\Phi_{edge} \equiv \mathtt{phiedge}$  .
- $\psi_{tor} \equiv tflux$  and  $\psi_{pol} \equiv tflux$  are immediately normalized (in readin() ) according to  $\psi_{tor,i} \to \psi_{tor,i}/\psi_0$ and  $\psi_{pol,i} \to \psi_{pol,i}/\psi_0$ , where  $\psi_0 \equiv \psi_{tor,N}$  on input.

## sweight(1:Mvol): star-like angle constraint weight

• the "star-like" poloidal angle constraint weights (only required for toroidal geometry, i.e. Igeometry=3) are given by

$$sweight_v \equiv upsilon \times (l_v/N_{vol})^w, \tag{189}$$

where  $l_v$  is the volume number, and  $w \equiv wpoloidal$ .

# TT(0:Mrad,0:1,0:1): Chebyshev polynomials at inner/outer interface

- TT (0:Lrad, 0:1, 0:1) gives the Chebyshev polynomials, and their first derivative, evaluated at s=-1and s = +1.
- Precisely, TT (1, i, d)  $\equiv T_i^{(d)}(s_i)$  for  $s_0 = -1$  and  $s_1 = +1$ .
- Note that  $T_l^{(0)}(s) = s^l$  and  $T_l^{(1)}(s) = s^{l+1}l^2$  for  $s = \pm 1$ .
- · Note that

$$T_{l}(-1) = \begin{cases} +1, & \text{if } l \text{ is even,} \\ -1, & \text{if } l \text{ is odd;} \end{cases} \qquad T_{l}(+1) = \begin{cases} +1, & \text{if } l \text{ is even,} \\ +1, & \text{if } l \text{ is odd;} \end{cases}$$
(190)  
$$T'_{l}(-1) = \begin{cases} -l^{2}, & \text{if } l \text{ is even,} \\ +l^{2}, & \text{if } l \text{ is odd.} \end{cases} \qquad T'_{l}(+1) = \begin{cases} +l^{2}, & \text{if } l \text{ is even,} \\ +l^{2}, & \text{if } l \text{ is odd.} \end{cases}$$
(191)

$$T'_l(-1) = \begin{cases} -l^2, & \text{if } l \text{ is even,} \\ +l^2, & \text{if } l \text{ is odd;} \end{cases} \qquad T'_l(+1) = \begin{cases} +l^2, & \text{if } l \text{ is even,} \\ +l^2, & \text{if } l \text{ is odd.} \end{cases}$$
(191)

- TT (0:Mrad, 0:1, 0:1) is used in routines that explicity require interface information, such as
  - the interface force-balance routine, Iforce()
  - the virtual casing routine, casing()
  - computing the rotational-transform on the interfaces, tr00ab()
  - computing the covariant components of the interface magnetic field, sc00aa()
  - enforcing the constraints on the Beltrami fields, matrix() and

computing the enclosed currents of the vacuum field, curent().

## ImagneticOK(1:Mvol): Beltrami/vacuum error flag

- · error flags that indicate if the magnetic field in each volume has been successfully constructed
- ImagneticOK is initialized to .false. in dforce() before the Beltrami solver routines are called. If the construction of the Beltrami field is successful (in either ma02aa() or mp00ac() ) then ImagneticOK is set to .true. .

#### Lhessianallocated

• The internal logical variable, Lhessianallocated, indicates whether the `'Hessian' matrix of second-partial derivatives (really, the first derivatives of the force-vector) has been allocated, or not!

## ki(1:mn,0:1): Fourier identification

· Consider the "abbreviated" representation for a double Fourier series,

$$\sum_{i} f_{i} \cos(m_{i}\theta - n_{i}\zeta) \equiv \sum_{n=0}^{N_{0}} f_{0,n} \cos(-n\zeta) + \sum_{m=1}^{M_{0}} \sum_{n=-N_{0}}^{N_{0}} f_{m,n} \cos(m\theta - n\zeta),$$
(192)

and the same representation but with enhanced resolution,

$$\sum_{k} \bar{f}_{k} \cos(\bar{m}_{k}\theta - \bar{n}_{k}\zeta) \equiv \sum_{n=0}^{N_{1}} f_{0,n} \cos(-n\zeta) + \sum_{m=1}^{M_{1}} \sum_{n=-N_{1}}^{N_{1}} f_{m,n} \cos(m\theta - n\zeta), \tag{193}$$

with  $M_1 \geq M_0$  and  $N_1 \geq N_0$ ; then  $k_i \equiv \text{ki}$  (i, 0) is defined such that  $\bar{m}_{k_i} = m_i$  and  $\bar{n}_{k_i} = n_i$ .

## kija(1:mn,1:mn,0:1), kijs(1:mn,1:mn,0:1): Fourier identification

• Consider the following quantities, which are computed in ma00aa(), where  $\bar{g}^{\mu\nu} = \sum_k \bar{g}_k^{\mu\nu} \cos \alpha_k$  for  $\alpha_k \equiv m_k \theta - n_k \zeta$ ,

$$\oint \!\! \int \!\! d\theta d\zeta \ \bar{g}^{\mu\nu} \cos \alpha_i \ \cos \alpha_j = \frac{1}{2} \oint \!\! \int \!\! d\theta d\zeta \ \bar{g}^{\mu\nu} (+\cos \alpha_{k_{ij+}} + \cos \alpha_{k_{ij-}}), \tag{194}$$

$$\oint \!\! \oint d\theta d\zeta \ \bar{g}^{\mu\nu} \sin \alpha_i \cos \alpha_j = \frac{1}{2} \oint \!\! \oint d\theta d\zeta \ \bar{g}^{\mu\nu} (+\sin \alpha_{k_{ij+}} + \sin \alpha_{k_{ij-}}), \tag{196}$$

$$\oint \!\! \int \!\! d\theta d\zeta \ \bar{g}^{\mu\nu} \sin \alpha_i \ \sin \alpha_j = \frac{1}{2} \oint \!\! \int \!\! d\theta d\zeta \ \bar{g}^{\mu\nu} (-\cos \alpha_{k_{ij+}} + \cos \alpha_{k_{ij-}}), \tag{197}$$

where  $(m_{k_{ij+}},n_{k_{ij+}})=(m_i+m_j,n_i+n_j)$  and  $(m_{k_{ij-}},n_{k_{ij-}})=(m_i-m_j,n_i-n_j)$ ; then kija (i, j, 0)  $\equiv k_{ij+}$  and kijs (i, j, 0)  $\equiv k_{ij-}$ .

• Note that Eqn. (193) does not include m < 0; so, if  $m_i - m_j < 0$  then  $k_{ij-}$  is re-defined such that  $(m_{k_{ij-}}, n_{k_{ij-}}) = (m_j - m_i, n_j - n_i)$ ; and similarly for the case m = 0 and n < 0. Also, take care that the sign of the sine harmonics in the above expressions will change for these cases.

# djkp iotakki

# cheby(0:Lrad,0:2): Chebyshev polynomial workspace

- cheby (0:Lrad, 0:2) is global workspace for computing the Chebyshev polynomials, and their derivatives, using the recurrence relations  $T_0(s) = 1$ ,  $T_1(s) = s$  and  $T_l(s) = 2sT_{l-1}(s) T_{l-2}(s)$ .
- These are computed as required, i.e. for arbitrary s, in bfield(), jo00aa() and ma00aa().
- Note that the quantities required for ma00aa() are for fixed s, and so these quantities should be precomputed.

# Iquad, gaussianweight, gaussianabscissae: Gauss-Legendre quadrature

- The volume integrals are computed using a "Fourier" integration over the angles and by Gauss-Legendre quadrature over the radial, i.e.  $\int \! f(s) ds = \sum_k \omega_k f(s_k)$ .
- The quadrature resolution in each volume is give by Iquad (1:Mvol) which is determined as follows:
  - if Nquad.gt.0, then Iquad(vvol)=Nquad
  - if Nquad.le.0 and .not.Lcoordinatesingularity, then Iquad(vvol)=2\*Lrad(vvol)-Nquad
  - if Nquad.le.0 and Lcoordinatesingularity , then Iquad(vvol)=2\*Lrad(vvol)-Nquad+Mpol
- The Gaussian weights and abscissae are given by gaussianweight (1:maxIquad, 1:Mvol) and gaussianabscissae(1:maxIquad, 1:Mvol), which are computed using modified Numerical Recipes routine gauleg().
- Iquad v is passed through to ma00aa() to compute the volume integrals of the metric elements; also see jo00aa(), where Iquad v is used to compute the volume integrals of  $||\nabla \times \mathbf{B} \mu \mathbf{B}||$ .

# LBsequad, LBnewton and LBlinear

• LBsequad, LBnewton and LBlinear depend simply on LBeltrami, which is described in global.f90

# BBweight(1:mn): weighting of force-imbalance harmonics

· weight on force-imbalance harmonics;

$$BBweight_i \equiv opsilon \times exp \left[ -escale \times (m_i^2 + n_i^2) \right]$$
 (198)

• this is only used in dforce() in constructing the force-imbalance vector

## mmpp(1:mn): spectral condensation weight factors

· spectral condensation weight factors;

$$mmpp(i) \equiv m_i^p, \tag{199}$$

where  $p \equiv pcondense$ .

#### NAdof, Ate, Aze, Ato and Azo: degrees-of-freedom in magnetic vector potential

- NAdof (1:Mvol) = total number of degrees-of-freedom in magnetic vector potential, including Lagrange multipliers, in each volume. This can de deduced from matrix().
- The components of the vector potential,  $\mathbf{A}=A_{\theta}\nabla+A_{\zeta}\nabla\zeta$ , are

$$A_{\theta}(s,\theta,\zeta) = \sum_{i,l} A_{\theta,e,i,l} \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} A_{\theta,o,i,l} \overline{T}_{l,i}(s) \sin \alpha_i, \tag{200}$$

$$A_{\zeta}(s,\theta,\zeta) = \sum_{i,l} A_{\zeta,e,i,l} \, \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} A_{\zeta,o,i,l} \, \overline{T}_{l,i}(s) \sin \alpha_i, \tag{201}$$

where  $\overline{T}_{l,i}(s) \equiv \overline{s}^{m_i/2} T_l(s)$ ,  $T_l(s)$  is the Chebyshev polynomial, and  $\alpha_j \equiv m_j \theta - n_j \zeta$ . The regularity factor,  $\overline{s}^{m_i/2}$ , where  $\overline{s} \equiv (1+s)/2$ , is only included if there is a coordinate singularity in the domain (i.e. only in the innermost volume, and only in cylindrical and toroidal geometry.)

• The Chebyshev-Fourier harmonics of the covariant components of the magnetic vector potential are kept in

$$A_{\theta,e,i,l} \equiv \text{Ate}(v,0,j) \%s(1), \qquad (202)$$

$$A_{\zeta,e,i,l} \equiv \operatorname{Aze}(v,0,j) \%s(1),$$
 (203)

$$A_{\theta,o,i,l} \equiv \text{Ato}(v,0,j) \%s(1), \text{and}$$
 (204)

$$A_{\zeta,o,i,l} \equiv \text{Azo}(v,0,j)\%s(1); \tag{205}$$

where v=1, Mvol labels volume, j=1, mn labels Fourier harmonic, and  $l=0, \texttt{Lrad}\ (v)$  labels Chebyshev polynomial. (These arrays also contains derivative information.)

- If Linitguess=1, a guess for the initial state for the Beltrami fields is constructed. An initial state is required for iterative solvers of the Beltrami fields, see LBeltrami.
- If Linitguess=2, the initial state for the Beltrami fields is read from file (see ra00aa()). An initial state is required for iterative solvers of the Beltrami fields, see LBeltrami.

#### workspace

goomne, goomno: metric information These are defined in metrix(), and used in ma00aa(). gssmne, gssmno: metric information These are defined in metrix(), and used in ma00aa(). gstmne, gstmno: metric information These are defined in metrix(), and used in ma00aa(). gszmne, gszmno: metric information These are defined in metrix(), and used in ma00aa(). gttmne, gttmno: metric information These are defined in metrix(), and used in ma00aa(). gtzmne, gtzmno: metric information These are defined in metrix(), and used in ma00aa(). gzzmne, gzzmno: metric information These are defined in metrix(), and used in ma00aa(). cosi(1:Ntz,1:mn) and sini(1:Ntz,1:mn)

· Trigonometric factors used in various Fast Fourier transforms, where

$$\cos i_{j,i} = \cos(m_i \theta_j - n_i \zeta_j), \tag{206}$$

$$\sin_{j,i} = \sin(m_i \theta_j - n_i \zeta_j). \tag{207}$$

# psifactor(1:mn,1:Mvol): coordinate "pre-conditioning" factor

· In toroidal geometry, the coordinate "pre-conditioning" factor is

$$f_{j,v} \equiv \begin{cases} \psi_{t,v}^0 &, & \text{for } m_j = 0, \\ \psi_{t,v}^{m_j/2} &, & \text{otherwise.} \end{cases}$$
 (208)

where  $\psi_{t,v} \equiv exttt{tflux}$  is the (normalized?) toroidal flux enclosed by the v-th interface.

- psifactor is used in packxi(), dforce() and hesian().
- inifactor is similarly constructed, with

$$f_{j,v} \equiv \begin{cases} \psi_{t,v}^{1/2} &, & \text{for } m_j = 0, \\ \psi_{t,v}^{m_j/2} &, & \text{otherwise.} \end{cases}$$
 (209)

and used only for the initialization of the surfaces taking into account axis information if provided.

### **Bsupumn and Bsupvmn**

# diotadxup and glambda: transformation to straight fieldline angle

- Given the Beltrami fields in any volume, the rotational-transform on the adjacent interfaces may be determined (in tr00ab()) by constructing the straight fieldline angle on the interfaces.
- The rotational transform on the inner or outer interface of a given volume depends on the magnetic field in that volume, i.e.  $\pm \pm = \pm (\mathbf{B}_{\pm})$ , so that

$$\delta \boldsymbol{\tau}_{\pm} = \frac{\partial \boldsymbol{\tau}_{\pm}}{\partial \mathbf{B}_{\pm}} \cdot \delta \mathbf{B}_{\pm}. \tag{210}$$

• The magnetic field depends on the Fourier harmonics of both the inner and outer interface geometry (represented here as  $x_j$ ), the helicity multiplier, and the enclosed poloidal flux, i.e.  $\mathbf{B}_{\pm} = \mathbf{B}_{\pm}(x_j, \mu, \Delta \psi_p)$ , so that

$$\delta \mathbf{B}_{\pm} = \frac{\partial \mathbf{B}_{\pm}}{\partial x_{i}} \delta x_{j} + \frac{\partial \mathbf{B}_{\pm}}{\partial \mu} \delta \mu + \frac{\partial \mathbf{B}_{\pm}}{\partial \Delta \psi_{p}} \delta \Delta \psi_{p}. \tag{211}$$

- The rotational-transforms, thus, can be considered to be functions of the geometry, the helicity-multiplier and the enclosed poloidal flux,  $\psi_{\pm} = \psi_{\pm}(x_i, \mu, \Delta \psi_p)$ .
- The rotational-transform, and its derivatives, on the inner and outer interfaces of each volume is stored in diotadxup(0:1,-1:2,1:Mvol) . The indices label:

- the first index labels the inner or outer interface,
- the the second one labels derivative, with
  - \* -1: indicating the derivative with respect to the interface geometry, i.e.  $\frac{\partial + \pm}{\partial x_j}$ ,
  - \* 0 : the rotational-transform itself,
  - \* 1,2 : the derivatives with respec to  $\mu$  and  $\Delta\psi_p$ , i.e.  $\frac{\partial\,\epsilon_{\,\pm}}{\partial\mu}$  and  $\frac{\partial\,\epsilon_{\,\pm}}{\partial\Delta\psi_p}$ ;
- The third index labels volume.
- The values of diotadxup are assigned in mp00aa() after calling tr00ab().

## vvolume, IBBintegral and IABintegral

· volume integrals

vvolume(i) = 
$$\int_{\mathcal{V}_i} dv$$
 (212)

lBBintegral(i) = 
$$\int_{\mathcal{V}_i} \mathbf{B} \cdot \mathbf{B} \, dv$$
 (213)

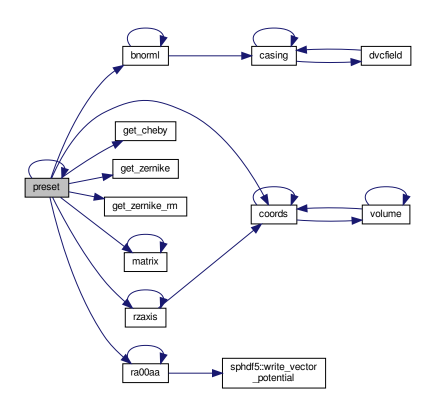
lABintegral(i) = 
$$\int_{\mathcal{V}_i} \mathbf{A} \cdot \mathbf{B} \, dv$$
 (214)

References allglobal::ajk, allglobal::ate, allglobal::ato, allglobal::aze, allglobal::azo, allglobal::bbe, allglobal::bbe, allglobal::bbweight, allglobal::beltramierror, allglobal::bemn, allglobal::bloweremn, allglobal::bloweromn, inputlist::bnc, bnorml(), inputlist::bns, allglobal::bomn, allglobal::bsupumn, allglobal::btemn, allglobal::btemn, allglobal::btemn, allglobal::bzemn, allglobal::bzemn, allglobal::cfmn, allglobal::cheby, allglobal::comn, coords(), allglobal::cosi, fftw interface::cplxin, fftw interface::cplxout, allglobal::diotadxup, allglobal::ditgpdxtp, allglobal::djkm, allglobal::djkp, allglobal::dpflux, allglobal::dradr, allglobal::dradz, allglobal::drbc, allglobal::drbs, allglobal::drij, allglobal::drodr, allglobal::drodz, allglobal::dtflux, allglobal::dxyz, allglobal::dzadr, allglobal::dzadz, allglobal::dzbc, allglobal::dzbs, allglobal::dzij, allglobal::dzodr, allglobal::dzodz, allglobal::efmn, inputlist::escale, allglobal::evmn, inputlist::forcetol, allglobal::fse, allglobal::fso, allglobal::gaussianabscissae, allglobal::gaussianweight, get cheby(), get zernike(), get zernike rm(), allglobal::glambda, allglobal::gmreslastsolution, constants::goldenmean, allglobal::goomne, allglobal::goomno, allglobal::gssmne, allglobal::gssmne, allglobal::gstmne, allglobal::gstmne, allglobal::gszmne, allglobal::gszmno, allglobal::gteta, allglobal::gttmne, allglobal::gttmne, allglobal::gtzmne, allglobal::gtzmno, allglobal::gvuij, allglobal::gvuij, allglobal::gzeta, allglobal::gzzmne, allglobal::gzzmno, constants::half, allglobal::halfmm, inputlist::helicity, allglobal::hnt, allglobal::hnz, allglobal::ibnc, allglobal::ibns, allglobal::iemn, inputlist::igeometry, allglobal::iie, allglobal::iiio, allglobal::ijimag, allglobal::ijreal, allglobal::im, allglobal::imagneticok, allglobal::ime, inputlist::impol, allglobal::ims, allglobal::in, allglobal::ine, allglobal::inifactor, allglobal::ins, inputlist::intor, allglobal::iomn, inputlist::iota, allglobal::iotakadd, allglobal::iotakkii, allglobal::iotaksgn, allglobal::iotaksub, allglobal::ipdtdpf, allglobal::iquad, allglobal::irbc, allglobal::irbs, allglobal::irij, inputlist::istellsym, allglobal::ivnc, allglobal::ivns, inputlist::ivolume, allglobal::izbc, allglobal::izbs, allglobal::izij, allglobal::jiimag, allglobal::jireal, allglobal::jkimag, allglobal::jkreal, allglobal::jxyz, allglobal::ki, allglobal::kija, allglobal::kijs, allglobal::kijmag, allglobal::kjreal, allglobal::labintegral, allglobal::lbbintegral, inputlist::lbeltrami, allglobal::lblinear, allglobal::lbnewton, allglobal::lbsequad, inputlist::lcheck, inputlist::lconstraint, allglobal::lcoordinatesingularity, inputlist::lfindzero, inputlist::lfreebound, allglobal::lgdof, inputlist::lgmresprec, allglobal::lhessianallocated, inputlist::lhevalues, inputlist::lhevectors, inputlist::lhmatrix, allglobal::liluprecond, inputlist::linitgues, inputlist::linitialize, allglobal::lma, inputlist::lmatsolver, allglobal::lmavalue, allglobal::lmb, allglobal::lmbvalue, allglobal::lmc, allglobal::lm allglobal::lmd, allglobal::lmdvalue, allglobal::lme, allglobal::lmevalue, allglobal::lmf, allglobal::lmfvalue, allglobal::lmg, allglobal::lmgvalue, allglobal::lmh, allglobal::lmhvalue, allglobal::lmns, allglobal::lmpol, allglobal::lntor, allglobal::localconstraint, inputlist::lp, inputlist::lperturbed, inputlist::lq, inputlist::lrad, inputlist::lreflect, matrix(), inputlist::maxrndgues, allglobal::mmpp, allglobal::mn, allglobal::mne, allglobal::mns, inputlist::mpol, inputlist::mregular, inputlist::mu, constants::mu0, allglobal::myid, allglobal::nadof, inputlist::ndiscrete, allglobal::ndmas, allglobal::ndmasmax, allglobal::nfielddof, inputlist::nfp, allglobal::ngdof, allglobal::notmatrixfree, allglobal::notstellsym, inputlist::nppts, inputlist::nquad, allglobal::nt, inputlist::ntor, allglobal::ntz, inputlist::nvol, allglobal::nxyz, allglobal::nz, allglobal:: allglobal::ofmn, inputlist::oita, constants::one, inputlist::opsilon, fileunits::ounit, inputlist::pcondense, allglobal::pemn, inputlist::pflux, inputlist::phiedge, constants::pi2, inputlist::pl, fftw interface::planb, inputlist::pr, preset(), allglobal::psifactor, inputlist::ql, inputlist::qr, constants::quart, ra00aa(), inputlist::rac, inputlist::ras, inputlist::rbc, inputlist::rbs, allglobal::regumm, allglobal::rij, inputlist::rp, inputlist::rq, allglobal::rscale,

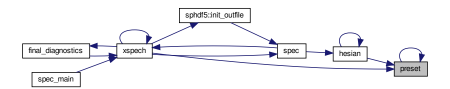
allglobal::rtm, allglobal::rtm, inputlist::rwc, inputlist::rws, rzaxis(), allglobal::semn, allglobal::sfmn, allglobal::sg, allglobal::simn, allglobal::simn, allglobal::simn, allglobal::sontz, numerical::sqrtmachprec, allglobal::sweight, inputlist::tflux, allglobal::trij, allglobal::tt, allglobal::tzij, inputlist::upsilon, inputlist::vnc, inputlist::vns, numerical::vsmall, allglobal::vvolume, inputlist::wpoloidal, allglobal::yesstellsym, inputlist::zac, inputlist::zac, inputlist::zbc, inputlist::zbs, allglobal::zernike, constants::zero, allglobal::zij, inputlist::zwc, and inputlist::zws.

Referenced by hesian(), preset(), and xspech().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.18 Output file(s)

## **Modules**

module sphdf5
 writing the HDF5 output file

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#### Functions/Subroutines

• subroutine ra00aa (writeorread)

Writes vector potential to .ext.sp.A .

• subroutine sphdf5::init\_outfile

Initialize the interface to the HDF5 library and open the output file.

subroutine sphdf5::mirror input to outfile

Mirror input variables into output file.

· subroutine sphdf5::init\_convergence\_output

Prepare convergence evolution output.

• subroutine sphdf5::write\_convergence\_output (nDcalls, ForceErr)

Write convergence output (evolution of interface geometry, force, etc).

subroutine sphdf5::write grid

Write the magnetic field on a grid.

subroutine sphdf5::init flt output (numTrajTotal)

Initialize field line tracing output group and create array datasets.

subroutine sphdf5::write\_poincare (offset, data, success)

Write a hyperslab of Poincare data corresponding to the output of one parallel worker.

• subroutine sphdf5::write transform (offset, length, lvol, diotadxup, fiota)

Write the rotational transform output from field line following.

subroutine sphdf5::finalize flt output

Finalize Poincare output.

• subroutine sphdf5::write\_vector\_potential (sumLrad, allAte, allAze, allAto, allAzo)

Write the magnetic vector potential Fourier harmonics to the output file group /vector\_potential.

subroutine sphdf5::hdfint

Write the final state of the equilibrium to the output file.

· subroutine sphdf5::finish\_outfile

Close all open HDF5 objects (we know of) and list any remaining still-open objects.

#### 7.18.1 Detailed Description

# 7.18.2 Function/Subroutine Documentation

Writes vector potential to .ext.sp.A .

# representation of vector potential

• The components of the vector potential,  $\mathbf{A} = A_{\theta} \nabla + A_{\zeta} \nabla \zeta$ , are

$$A_{\theta}(s,\theta,\zeta) = \sum_{i,l} A_{\theta,e,i,l} \, \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} A_{\theta,o,i,l} \, \overline{T}_{l,i}(s) \sin \alpha_i, \tag{215}$$

$$A_{\zeta}(s,\theta,\zeta) = \sum_{i,l} A_{\zeta,e,i,l} \, \overline{T}_{l,i}(s) \cos \alpha_i + \sum_{i,l} A_{\zeta,o,i,l} \, \overline{T}_{l,i}(s) \sin \alpha_i, \tag{216}$$

where  $\overline{T}_{l,i}(s) \equiv \overline{s}^{m_i/2} T_l(s)$ ,  $T_l(s)$  is the Chebyshev polynomial, and  $\alpha_j \equiv m_j \theta - n_j \zeta$ . The regularity factor,  $\overline{s}^{m_i/2}$ , where  $\overline{s} \equiv (1+s)/2$ , is only included if there is a coordinate singularity in the domain (i.e. only in the innermost volume, and only in cylindrical and toroidal geometry.)

# file format

• The format of the files containing the vector potential is as follows:

```
open(aunit, file="."//trim(ext)//".sp.A", status="replace", form="unformatted" )
write(aunit) mvol, mpol, ntor, mn, nfp ! integers;
write(aunit) im(1:mn) ! integers; poloidal modes;
write(aunit) in(1:mn) ! integers; toroidal modes;
do vvol = 1, mvol ! integers; loop over volumes;
write(aunit) lrad(vvol) ! integers; the radial resolution in each volume may be different;
do ii = 1, mn
write(aunit) ate(vvol,ii)%s(0:lrad(vvol)) ! reals;
write(aunit) aze(vvol,ii)%s(0:lrad(vvol)) ! reals;
write(aunit) aze(vvol,ii)%s(0:lrad(vvol)) ! reals;
write(aunit) azo(vvol,ii)%s(0:lrad(vvol)) ! reals;
enddo ! end of do ii;
enddo ! end of do vvol;
close(aunit)
```

### **Parameters**

in	writeorread	'W' to write the vector potential; 'R' to read it
----	-------------	---

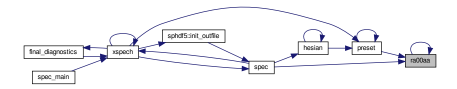
References allglobal::ate, allglobal::ato, fileunits::aunit, allglobal::aze, allglobal::azo, allglobal::cpus, allglobal::im, allglobal::im, inputlist::lrad, allglobal::mn, allglobal::mpi\_comm\_spec, inputlist::mpol, allglobal::myid, allglobal::ncpu, inputlist::nfp, inputlist::ntor, fileunits::ounit, ra00aa(), inputlist::wmacros, sphdf5::write\_vector\_potential(), and constants::zero.

Referenced by preset(), ra00aa(), and spec().

Here is the call graph for this function:



Here is the caller graph for this function:



# **7.18.2.2 mirror\_input\_to\_outfile()** subroutine sphdf5::mirror\_input\_to\_outfile Mirror input variables into output file.

The goal of this routine is to have an exact copy of the input file contents that were used to parameterize a given SPEC run. This also serves to check after the run if SPEC correctly understood the text-based input file.

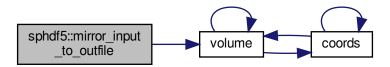
References inputlist::absacc, inputlist::absreq, inputlist::adiabatic, inputlist::bnc, inputlist::bns, inputlist::bnsblend, inputlist::c05factor, inputlist::c05xmax, inputlist::c05xtol, inputlist::curpol, inputlist::curpol

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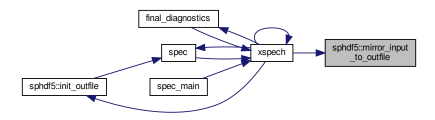
inputlist::igeometry, inputlist::imethod, inputlist::impol, allglobal::in, inputlist::iorder, inputlist::iorder, inputlist::iotatol, inputlist::iprecon, inputlist::istellsym, inputlist::isurf, inputlist::ivolume, inputlist::ladiabatic, inputlist::lbeltrami, inputlist::lcheck, inputlist::lconstraint, inputlist::lerrortype, inputlist::lextrap, inputlist::lfindzero, inputlist::lfreebound, inputlist::lgmresprec, inputlist::lhevalues, inputlist::lhevectors, inputlist::lhmatrix, inputlist::linitialize, inputlist::lmatsolver, inputlist::lperturbed, inputlist::lposdef, inputlist::lq, inputlist::lrad, inputlist::lreadgf, inputlist::lreflect, inputlist::lrzaxis, inputlist::lsparse, inputlist::lsvdiota, inputlist::ltiming, inputlist::lzerovac, inputlist::maxrndgues, inputlist::maxrndgues, inputlist::maxrndgues, inputlist::mupflo, allglobal::myid, inputlist::mpol, inputlist::mpol, inputlist::mregular, inputlist::mup, inputlist::mupfits, inputlist::mupflo, allglobal::myid, inputlist::ndiscrete, inputlist::nfp, inputlist::ngrid, inputlist::npts, inputlist::npts, inputlist::npts, inputlist::pondense, inputlist::pflux, inputlist::ntoraxis, inputlist::nvol, inputlist::ppts, inputlist::pr, inputlist::pscale, inputlist::ql, inputlist::rac, inputlist::rac, inputlist::rbc, inputlist::rbs, inputlist::rereq, inputlist::rpol, inputlist::rpol, inputlist::ror, inputlist::rwc, inputlist::rws, inputlist::rbs, inputlist::ror, i

Referenced by xspech().

Here is the call graph for this function:



Here is the caller graph for this function:



# **7.18.2.3** init\_convergence\_output() subroutine sphdf5::init\_convergence\_output Prepare convergence evolution output.

• The group iterations is created in the output file. This group contains the interface geometry at each iteration, which is useful for constructing movies illustrating the convergence. The data structure in use is an unlimited array of the following compound datatype:

```
DATATYPE H5T_COMPOUND {

H5T_NATIVE_INTEGER "nDcalls";

H5T_NATIVE_DOUBLE "Energy";

H5T_NATIVE_DOUBLE "ForceErr";

H5T_ARRAY { [Mvol+1] [mn] H5T_NATIVE_DOUBLE } "iRbc";

H5T_ARRAY { [Mvol+1] [mn] H5T_NATIVE_DOUBLE } "iZbs";

H5T_ARRAY { [Mvol+1] [mn] H5T_NATIVE_DOUBLE } "iRbs";

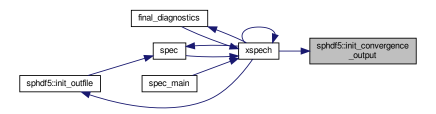
H5T_ARRAY { [Mvol+1] [mn] H5T_NATIVE_DOUBLE } "iZbc";

}
```

References sphdf5::dt\_energy\_id, sphdf5::dt\_forceerr\_id, sphdf5::dt\_irbc\_id, sphdf5::dt\_irbs\_id, sphdf5::dt\_izbc\_id, sphdf5::dt\_izbc\_id, sphdf5::dt\_izbs\_id, sphdf5::dt\_izbs\_id, sphdf5::memspace, allglobal::mn, allglobal::myid, and sphdf5::plist\_id.

Referenced by xspech().

Here is the caller graph for this function:



# 7.18.2.4 write\_grid() subroutine sphdf5::write\_grid

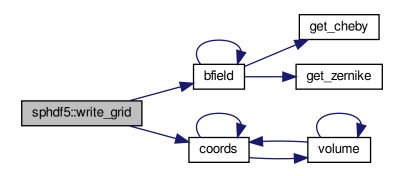
Write the magnetic field on a grid.

The magnetic field is evaluated on a regular grid in  $(s, \theta, \zeta)$  and the corresponding cylindrical coordinates (R, Z) as well as the cylindrical components of the magnetic field  $(B^R, B^\varphi, B^Z)$  are written out.

References bfield(), coords(), sphdf5::file\_id, allglobal::gbzeta, inputlist::igeometry, allglobal::ijimag, allglobal::ijreal, allglobal::ijreal, allglobal::lcoordinatesingularity, allglobal::lplasmaregion, inputlist::lrad, allglobal::lvacuumregion, allglobal::mn, allglobal::myid, inputlist::ngrid, allglobal::node, allglobal::nt, allglobal::ntz, inputlist::nvol, allglobal::nz, constants::one, constants::pi2, allglobal::rij, inputlist::rpol, inputlist::rtor, allglobal::sg, constants::two, constants::zero, and allglobal::zij.

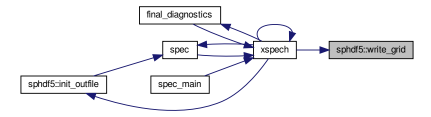
Referenced by xspech().

Here is the call graph for this function:



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Here is the caller graph for this function:



Initialize field line tracing output group and create array datasets.

The field-line tracing diagnostic is parallelized over volumes, where all threads/ranks produce individual output. This is gathered in the output file, stacked over the radial dimension. The success flag signals if the integrator was successful in following the fieldline for the derired number of toroidal periods.

#### **Parameters**

|--|

References sphdf5::dset\_id\_diotadxup, sphdf5::dset\_id\_fiota, sphdf5::dset\_id\_r, sphdf5::dset\_id\_s, sphdf5::dset\_id\_s, sphdf5::dset\_id\_s, sphdf5::dset\_id\_s, sphdf5::dset\_id\_s, sphdf5::dset\_id\_t, sphdf5::filespace\_fiota, sphdf5::filespace\_r, sphdf5::filespace\_s, sphdf5::filespace\_s, sphdf5::filespace\_t, sphdf5::filespace\_t, sphdf5::filespace\_t, sphdf5::grptransform, sphdf5::hdfier, sphdf5::memspace\_diotadxup, sphdf5::memspace\_r, sphdf5::memspace\_s, sphdf5::memspace\_s, sphdf5::memspace\_s, sphdf5::memspace\_s, sphdf5::memspace\_s, sphdf5::memspace\_s, sphdf5::memspace\_s, sphdf5::rankt, sphdf

Referenced by pp00aa().

Here is the caller graph for this function:



Write a hyperslab of Poincare data corresponding to the output of one parallel worker.

#### **Parameters**

offset	radial offset at which the data belongs
data	output from field-line tracing

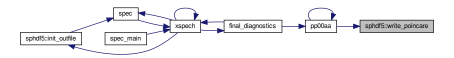
#### **Parameters**

success	flags to indicate if integrator was successful	]
---------	--	---

References sphdf5::dset\_id\_r, sphdf5::dset\_id\_s, sphdf5::dset\_id\_success, sphdf5::dset\_id\_t, sphdf5::dset\_id\_z, sphdf5::filespace\_r, sphdf5::filespace\_s, sphdf5::filespace\_s, sphdf5::filespace\_t, sphdf5::filespace\_z, sphdf5::memspace\_r, sphdf5::memspace\_s, sphdf5::memspace\_s, sphdf5::memspace\_t, sphdf5::memspace\_t, sphdf5::memspace\_t, sphdf5::memspace\_t, sphdf5::memspace\_z, allglobal::myid, inputlist::nppts, and allglobal::nz.

Referenced by pp00aa().

Here is the caller graph for this function:



# $\textbf{7.18.2.7} \quad \textbf{write\_transform()} \quad \texttt{subroutine sphdf5::write\_transform ()}$

```
integer, intent(in) offset,
integer, intent(in) length,
integer, intent(in) lvol,
real, dimension(:), intent(in) diotadxup,
real, dimension(:,:), intent(in) fiota)
```

Write the rotational transform output from field line following.

#### **Parameters**

offset	radial offset at which the data belongs
length	length of dataset to write
Ivol	nested volume index
diotadxup	derivative of rotational transform (?)
fiota	rotational transform

References sphdf5::dset\_id\_diotadxup, sphdf5::dset\_id\_fiota, sphdf5::filespace\_diotadxup, sphdf5::filespace\_fiota, sphdf5::hdfier, sphdf5::memspace\_diotadxup, sphdf5::memspace\_fiota, and sphdf5::rankt.

Referenced by pp00aa().

Here is the caller graph for this function:



# 7.18.2.8 finalize\_flt\_output() subroutine sphdf5::finalize\_flt\_output

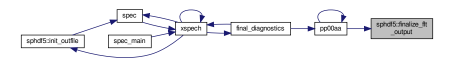
Finalize Poincare output.

This closes the still-open datasets related to field-line tracing, which had to be kept open during the tracing to be able to write the outputs directly when a given worker thread is finished.

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References sphdf5::dset\_id\_diotadxup, sphdf5::dset\_id\_fiota, sphdf5::dset\_id\_r, sphdf5::dset\_id\_s, sphdf5::dset\_id\_s, sphdf5::dset\_id\_s, sphdf5::dset\_id\_s, sphdf5::dset\_id\_s, sphdf5::dset\_id\_z, sphdf5::filespace\_diotadxup, sphdf5::filespace\_fiota, sphdf5::filespace\_r, sphdf5::filespace\_s, sphdf5::filespace\_s, sphdf5::filespace\_z, sphdf5::grppoincare, sphdf5::grptransform, sphdf5::hdfier, sphdf5::memspace\_diotadxup, sphdf5::memspace\_r, sphdf5::memspace\_s, sphdf5:

Here is the caller graph for this function:



```
7.18.2.9 write_vector_potential() subroutine sphdf5::write_vector_potential (
    integer, intent(in) sumLrad,
    real, dimension(:,:), intent(in) allAte,
    real, dimension(:,:), intent(in) allAze,
    real, dimension(:,:), intent(in) allAto,
    real, dimension(:,:), intent(in) allAzo)
```

Write the magnetic vector potential Fourier harmonics to the output file group /vector\_potential.

The data is stacked in the radial direction over  $\mbox{\tt Lrad}$ , since  $\mbox{\tt Lrad}$  can be different in each volume, but HDF5 only supports rectangular arrays. So, one needs to split the  $\mbox{\tt sumLrad}$  dimension into chunks given by the input  $\mbox{\tt Lrad}$  array.

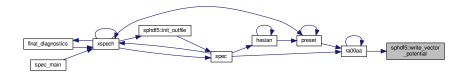
# **Parameters**

sumLrad	total sum over Lrad in all nested volumes
allAte	$A_{\mathrm{even}}^{ heta}$ for all nested volumes
allAze	$A_{ m even}^{\zeta}$ for all nested volumes
allAto	$A_{\mathrm{odd}}^{ heta}$ for all nested volumes
allAzo	$A_{ m odd}^{\zeta}$ for all nested volumes

References allglobal::ate, allglobal::ate, allglobal::aze, allglobal::aze, sphdf5::file\_id, allglobal::mn, and allglobal::myid.

Referenced by ra00aa().

Here is the caller graph for this function:



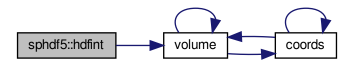
# 7.18.2.10 hdfint() subroutine sphdf5::hdfint

Write the final state of the equilibrium to the output file.

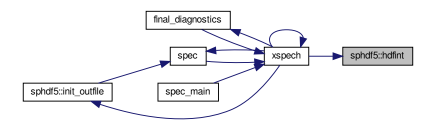
- In addition to the input variables, which are described in global(), the following quantities are written to ext. ←
   sp.h5:
- All quantities marked as real should be treated as double precision.

References inputlist::adiabatic, allglobal::beltramierror, inputlist::bnc, inputlist::bns, allglobal::bsupumn, allglobal::bsupumn, allglobal::bsupumn, allglobal::btomn, allglobal::bzemn, allglobal::bzemn, allglobal::bzemn, allglobal::bzemn, allglobal::cpus, allglobal::drbc, allglobal::drbc, allglobal::drbc, allglobal::drbc, allglobal::ibnc, allglobal::ibnc, allglobal::ibnc, allglobal::ibns, allglobal::in, allglobal::irbc, allglobal::irbs, allglobal::ivnc, allglobal::ivns, inputlist::ivolume, allglobal::izbc, allglobal::izbs, inputlist::lcheck, allglobal::lmns, inputlist::lperturbed, inputlist::lrad, allglobal::mn, inputlist::mu, allglobal::myid, allglobal::ncpu, inputlist::nvol, fileunits::ounit, inputlist::pflux, inputlist::rbc, inputlist::rbs, inputlist::tflux, allglobal::tt, inputlist::vnc, inputlist::vns, volume(), allglobal::vvolume, inputlist::zbc, and inputlist::zbs. Referenced by xspech().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.19 Coordinate axis

## **Functions/Subroutines**

• subroutine rzaxis (Mvol, mn, inRbc, inZbs, inRbs, inZbc, ivol, LcomputeDerivatives)

The coordinate axis is assigned via a poloidal average over an arbitrary surface.

# 7.19.1 Detailed Description

#### 7.19.2 Function/Subroutine Documentation

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```
real, dimension(1:mn,0:mvol) inZbs,
real, dimension(1:mn,0:mvol) inRbs,
real, dimension(1:mn,0:mvol) inZbc,
integer, intent(in) ivol,
logical, intent(in) LcomputeDerivatives )
```

The coordinate axis is assigned via a poloidal average over an arbitrary surface.

Specifies position of coordinate axis;  $\mathbf{x}_a(\zeta) \equiv \int \mathbf{x}_1(\theta, \zeta) dl / \int dl$ .

#### coordinate axis

- The coordinate axis is *not* an independent degree-of-freedom of the geometry. It is constructed by extrapolating the geometry of a given interface, as determined by  $i \equiv ivol$  which is given on input, down to a line.
- If the coordinate axis depends only on the *geometry* of the interface and not the angle parameterization, then the block tri-diagonal structure of the the force-derivative matrix is preserved.
- · Define the arc-length-weighted averages,

$$R_0(\zeta) \equiv \frac{\int_0^{2\pi} R_i(\theta, \zeta) \, dl}{\int_0^{2\pi} dl}, \qquad Z_0(\zeta) \equiv \frac{\int_0^{2\pi} Z_i(\theta, \zeta) \, dl}{\int_0^{2\pi} dl}, \tag{217}$$

where 
$$dl \equiv \dot{l} d\theta = \sqrt{\partial_{\theta} R_i(\theta, \zeta)^2 + \partial_{\theta} Z_i(\theta, \zeta)^2} d\theta$$
.

- (Note that if  $\dot{l}$  does not depend on  $\theta$ , i.e. if  $\theta$  is the equal arc-length angle, then the expressions simplify. This constraint is not enforced.)
- The geometry of the coordinate axis thus constructed only depends on the geometry of the interface, i.e. the angular parameterization of the interface is irrelevant.

## coordinate axis: derivatives

 The derivatives of the coordinate axis with respect to the Fourier harmonics of the given interface are given by

$$\frac{\partial R_0}{\partial R_{i,j}^c} = \int \left(\cos \alpha_j \ \dot{l} - \Delta R_i R_{i,\theta} \ m_j \sin \alpha_j / \ \dot{l}\right) d\theta / L \tag{218}$$

$$\frac{\partial R_0}{\partial R_{i,j}^s} = \int \left( \sin \alpha_j \ \dot{l} + \Delta R_i R_{i,\theta} \ m_j \cos \alpha_j / \ \dot{l} \right) d\theta / L \tag{219}$$

$$\frac{\partial R_0}{\partial Z_{i,j}^c} = \int \left( -\Delta R_i Z_{i,\theta} \, m_j \sin \alpha_j / \, \dot{l} \right) d\theta / L \tag{220}$$

$$\frac{\partial R_0}{\partial Z_{i,j}^s} = \int \left( +\Delta R_i Z_{i,\theta} \, m_j \cos \alpha_j / \, \dot{l} \right) d\theta / L \tag{221}$$

$$\frac{\partial Z_0}{\partial R_{i,j}^c} = \int \left( -\Delta Z_i R_{i,\theta} \, m_j \sin \alpha_j / \, \dot{l} \right) d\theta / L \tag{222}$$

$$\frac{\partial Z_0}{\partial R_{i,j}^s} = \int \left( +\Delta Z_i R_{i,\theta} \, m_j \cos \alpha_j / \, \dot{l} \right) d\theta / L \tag{223}$$

$$\frac{\partial Z_0}{\partial Z_{i,j}^c} = \int \left(\cos \alpha_j \ \dot{l} - \Delta Z_i Z_{i,\theta} \ m_j \sin \alpha_j / \ \dot{l}\right) d\theta / L \tag{224}$$

$$\frac{\partial Z_0}{\partial Z_{i,j}^s} = \int \left( \sin \alpha_j \ \dot{l} + \Delta Z_i Z_{i,\theta} \ m_j \cos \alpha_j / \ \dot{l} \right) d\theta / L \tag{225}$$

where 
$$L(\zeta) \equiv \int_0^{2\pi} \!\! dl.$$

#### some numerical comments

- First, the differential poloidal length,  $\dot{l} \equiv \sqrt{R_{\theta}^2 + Z_{\theta}^2}$ , is computed in real space using an inverse FFT from the Fourier harmonics of R and Z.
- Second, the Fourier harmonics of dl are computed using an FFT. The integration over  $\theta$  to construct  $L \equiv \int dl$  is now trivial: just multiply the m=0 harmonics of dl by  $2\pi$ . The ajk (1:mn) variable is used, and this is assigned in readin().
- ullet Next, the weighted  $R\,dl$  and  $Z\,dl$  are computed in real space, and the poloidal integral is similarly taken.
- · Last, the Fourier harmonics are constructed using an FFT after dividing in real space.

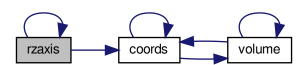
# **Parameters**

in	Mvol	
in	mn	
	inRbc	
	inZbs	
	inRbs	
	inZbc	
in	ivol	
	LcomputeDerivatives	

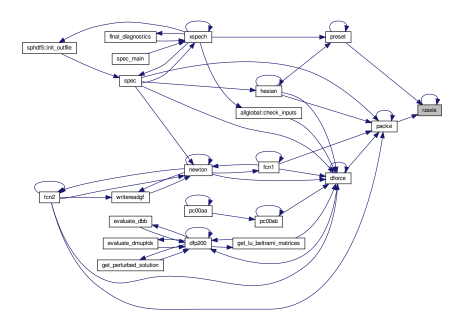
References allglobal::ajk, allglobal::cfmn, allglobal::comn, coords(), allglobal::cosi, allglobal::cpus, allglobal::dbdx, allglobal::dradz, allglobal::dradz, allglobal::dradz, allglobal::dzadz, allglobal::dzadz, allglobal::dzadz, allglobal::dzadz, allglobal::dzadz, allglobal::dzadz, allglobal::dzadz, allglobal::dzadz, allglobal::dzadz, allglobal::ijimag, allglobal::ijimag, allglobal::ijimag, allglobal::ijimag, allglobal::ijimag, allglobal::jimag, allglobal::

Referenced by packxi(), preset(), and rzaxis().

Here is the call graph for this function:



Here is the caller graph for this function:



## 7.20 Rotational Transform

#### **Functions/Subroutines**

subroutine tr00ab (Ivol, mn, NN, Nt, Nz, iflag, Idiota)
 Calculates rotational transform given an arbitrary tangential field.

## 7.20.1 Detailed Description

# 7.20.2 Function/Subroutine Documentation

```
7.20.2.1 tr00ab() subroutine tr00ab (
    integer, intent(in) lvol,
    integer, intent(in) mn,
    integer, intent(in) NN,
    integer, intent(in) Nt,
    integer, intent(in) Nz,
    integer, intent(in) iflag,
    real, dimension(0:1,-1:2), intent(inout) ldiota)
```

Calculates rotational transform given an arbitrary tangential field.

Calculates transform,  $\iota = \dot{\theta}(1 + \lambda_{\theta}) + \lambda_{\zeta}$ , given  $\mathbf{B}|_{\mathcal{I}}$ .

## constructing straight field line angle on interfaces

• The algorithm stems from introducing a straight field line angle  $\theta_s=\theta+\lambda(\theta,\zeta)$ , where

$$\lambda = \sum_{j} \lambda_{o,j} \sin(m_j \theta - n_j \zeta) + \sum_{j} \lambda_{e,j} \cos(m_j \theta - n_j \zeta)$$
 (226)

and insisting that

$$\frac{\mathbf{B} \cdot \nabla \theta_s}{\mathbf{B} \cdot \nabla \zeta} = \dot{\theta} (1 + \lambda_{\theta}) + \lambda_{\zeta} = t, \tag{227}$$

where + is a constant that is to be determined.

• Writing  $\dot{\theta} = -\partial_s A_{\mathcal{L}}/\partial_s A_{\theta}$ , we have

$$\partial_s A_\theta + \partial_s A_\zeta \lambda_\theta - \partial_s A_\theta \lambda_\zeta = -\partial_s A_\zeta \tag{228}$$

· Expanding this equation we obtain

$$(A'_{\theta,e,k}\cos\alpha_k + A'_{\theta,o,k}\sin\alpha_k) +$$

$$+ (A'_{\zeta,e,k}\cos\alpha_k + A'_{\zeta,o,k}\sin\alpha_k) (+m_j\lambda_{o,j}\cos\alpha_j - m_j\lambda_{e,j}\sin\alpha_j)$$

$$- (A'_{\theta,e,k}\cos\alpha_k + A'_{\theta,o,k}\sin\alpha_k) (-n_j\lambda_{o,j}\cos\alpha_j + n_j\lambda_{e,j}\sin\alpha_j)$$

$$= - (A'_{\zeta,e,k}\cos\alpha_k + A'_{\zeta,o,k}\sin\alpha_k),$$
(229)

where summation over  $k=1, \min j=2, \min s$  is implied

· After applying double angle formulae,

$$(A'_{\theta,e,k}\cos\alpha_k + A'_{\theta,o,k}\sin\alpha_k) +$$

$$+ \lambda_{o,j} \left( + m_j A'_{\zeta,e,k} + n_j A'_{\theta,e,k} \right) \left[ + \cos(\alpha_k + \alpha_j) + \cos(\alpha_k - \alpha_j) \right] / 2$$

$$+ \lambda_{e,j} \left( - m_j A'_{\zeta,e,k} - n_j A'_{\theta,e,k} \right) \left[ + \sin(\alpha_k + \alpha_j) - \sin(\alpha_k - \alpha_j) \right] / 2$$

$$+ \lambda_{o,j} \left( + m_j A'_{\zeta,o,k} + n_j A'_{\theta,o,k} \right) \left[ + \sin(\alpha_k + \alpha_j) + \sin(\alpha_k - \alpha_j) \right] / 2$$

$$+ \lambda_{e,j} \left( - m_j A'_{\zeta,o,k} - n_j A'_{\theta,o,k} \right) \left[ - \cos(\alpha_k + \alpha_j) + \cos(\alpha_k - \alpha_j) \right] / 2$$

$$= - \left( A'_{\zeta,e,k}\cos\alpha_k + A'_{\zeta,o,k}\sin\alpha_k \right), \tag{230}$$

and equating coefficients, an equation of the form  $\mathbf{A} \cdot \mathbf{x} = \mathbf{b}$  is obtained, where

$$\mathbf{x} = (\underbrace{t}_{\mathbf{x}[1]}, \underbrace{\lambda_{o,2}, \lambda_{o,3}, \dots}_{\mathbf{x}[2:N]}, \underbrace{\lambda_{e,2}, \lambda_{e,3}, \dots}_{\mathbf{x}[N+1:2N-1]})^{T}.$$
(231)

# alternative iterative method

• Consider the equation  $\dot{\theta}(1+\lambda_{\theta})+\lambda_{\zeta}=\pm$ , where  $\lambda=\sum_{j}\lambda_{j}\sin\alpha_{j}$ , given on a grid

$$\dot{\theta}_i + \dot{\theta}_i \sum_j m_j \cos \alpha_{i,j} \lambda_j - \sum_j n_j \cos \alpha_{i,j} \lambda_j = \pm,$$
(232)

where i labels the grid point.

· This is a matrix equation...

# Parameters

Ivol	
mn	
NN	
Nt	
Nz	
iflag	
Idiota	

References allglobal::ate, allglobal::ato, allglobal::aze, allglobal::azo, allglobal::cpus, allglobal::glambda, constants::goldenmean, constants::half, allglobal::hnt, allglobal::hnz, allglobal::im, inputlist::imethod, allglobal::ims, allglobal::in, allglobal::iotaksgn, allglobal::

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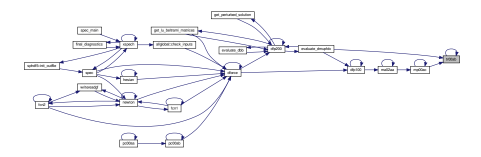
constants::third, tr00ab(), allglobal::tt, constants::two, numerical::vsmall, inputlist::wmacros, allglobal::yesstellsym, and constants::zero.

Referenced by evaluate\_dmupfdx(), mp00ac(), and tr00ab().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.21 Plasma volume

# Functions/Subroutines

subroutine volume (Ivol, vflag)

Computes volume of each region; and, if required, the derivatives of the volume with respect to the interface geometry.

# 7.21.1 Detailed Description

# 7.21.2 Function/Subroutine Documentation

Computes volume of each region; and, if required, the derivatives of the volume with respect to the interface geometry.

Calculates volume of each region;  $V_i \equiv \int dv$ .

# volume integral

- The volume enclosed by the v-th interface is given by the integral

$$V = \int_{\mathcal{V}} dv = \frac{1}{3} \int_{\mathcal{V}} \nabla \cdot \mathbf{x} \, dv = \frac{1}{3} \int_{\mathcal{S}} \mathbf{x} \cdot d\mathbf{s} = \frac{1}{3} \int_{0}^{2\pi} d\theta \int_{0}^{2\pi/N} d\zeta \quad \mathbf{x} \cdot \mathbf{x}_{\theta} \times \mathbf{x}_{\zeta}|^{s}$$
(233)

where we have used  $\nabla \cdot \mathbf{x} = 3$ , and have assumed that the domain is periodic in the angles.

# representation of surfaces

· The coordinate functions are

$$R(\theta,\zeta) = \sum_{i} R_{e,i} \cos \alpha_i + \sum_{i} R_{o,i} \sin \alpha_i$$
 (234)

$$Z(\theta,\zeta) = \sum_{i} Z_{e,i} \cos \alpha_i + \sum_{i} Z_{o,i} \sin \alpha_i, \qquad (235)$$

where  $\alpha_i \equiv m_i \theta - n_i \zeta$ .

## geometry

- The geometry is controlled by the input parameter Igeometry as follows:
- Igeometry.eq.1: Cartesian:  $\sqrt{g}=R_s$

$$V = \int_0^{2\pi} d\theta \int_0^{2\pi/N} d\zeta R$$
$$= 2\pi \frac{2\pi}{N} R_{e,1}$$
(236)

• Igeometry.eq.2: cylindrical:  $\sqrt{g}=RR_s=\frac{1}{2}\partial_s(R^2)$ 

$$V = \frac{1}{2} \int_{0}^{2\pi} d\theta \int_{0}^{2\pi/N} d\zeta R^{2}$$

$$= \frac{1}{2} 2\pi \frac{2\pi}{N} \frac{1}{2} \sum_{i} \sum_{j} R_{e,i} R_{e,j} \left[ \cos(\alpha_{i} - \alpha_{j}) + \cos(\alpha_{i} + \alpha_{j}) \right]$$

$$+ \frac{1}{2} 2\pi \frac{2\pi}{N} \frac{1}{2} \sum_{i} \sum_{j} R_{o,i} R_{o,j} \left[ \cos(\alpha_{i} - \alpha_{j}) - \cos(\alpha_{i} + \alpha_{j}) \right]$$
(237)

• Igeometry.eq.3: toroidal:  $\mathbf{x} \cdot \mathbf{e}_{\theta} \times \mathbf{e}_{\zeta} = R(ZR_{\theta} - RZ_{\theta})$  This is computed by fast Fourier transform:

$$V = \frac{1}{3} \int_{0}^{2\pi} d\theta \int_{0}^{2\pi/N} d\zeta R(ZR_{\theta} - RZ_{\theta})$$

$$= \frac{1}{3} \sum_{i} \sum_{j} \sum_{k} R_{e,i} (Z_{e,j}R_{o,k} - R_{e,j}Z_{o,k}) (+m_{k}) \iint d\theta d\zeta \cos \alpha_{i} \cos \alpha_{j} \cos \alpha_{k}$$

$$+ \frac{1}{3} \sum_{i} \sum_{j} \sum_{k} R_{e,i} (Z_{o,j}R_{e,k} - R_{o,j}Z_{e,k}) (-m_{k}) \iint d\theta d\zeta \cos \alpha_{i} \sin \alpha_{j} \sin \alpha_{k}$$

$$+ \frac{1}{3} \sum_{i} \sum_{j} \sum_{k} R_{o,i} (Z_{e,j}R_{e,k} - R_{e,j}Z_{e,k}) (-m_{k}) \iint d\theta d\zeta \sin \alpha_{i} \cos \alpha_{j} \sin \alpha_{k}$$

$$+ \frac{1}{3} \sum_{i} \sum_{j} \sum_{k} R_{o,i} (Z_{o,j}R_{o,k} - R_{o,j}Z_{o,k}) (+m_{k}) \iint d\theta d\zeta \sin \alpha_{i} \sin \alpha_{j} \cos \alpha_{k}$$

$$(238)$$

- (Recall that the integral over an odd function is zero, so various terms in the above expansion have been ignored.)
- · The trigonometric terms are

```
4 \cos \alpha_i \cos \alpha_j \cos \alpha_k = + \cos(\alpha_i + \alpha_j + \alpha_k) + \cos(\alpha_i + \alpha_j - \alpha_k) + \cos(\alpha_i - \alpha_j + \alpha_k) + \cos(\alpha_i - \alpha_j - \alpha_k)
4 \cos \alpha_i \sin \alpha_j \sin \alpha_k = - \cos(\alpha_i + \alpha_j + \alpha_k) + \cos(\alpha_i + \alpha_j - \alpha_k) + \cos(\alpha_i - \alpha_j + \alpha_k) - \cos(\alpha_i - \alpha_j - \alpha_k)
4 \sin \alpha_i \cos \alpha_j \sin \alpha_k = - \cos(\alpha_i + \alpha_j + \alpha_k) + \cos(\alpha_i + \alpha_j - \alpha_k) - \cos(\alpha_i - \alpha_j + \alpha_k) + \cos(\alpha_i - \alpha_j - \alpha_k)
4 \sin \alpha_i \sin \alpha_j \cos \alpha_k = - \cos(\alpha_i + \alpha_j + \alpha_k) - \cos(\alpha_i + \alpha_j - \alpha_k) + \cos(\alpha_i - \alpha_j + \alpha_k) + \cos(\alpha_i - \alpha_j - \alpha_k)
```

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· The required derivatives are

$$3\frac{\partial V}{\partial R_{e,i}} = (+Z_{e,j}R_{o,k}m_k - R_{e,j}Z_{o,k}m_k - R_{e,j}Z_{o,k}m_k) \quad \iint d\theta d\zeta \cos \alpha_i \cos \alpha_j \cos \alpha_k$$

$$+ (-Z_{o,j}R_{e,k}m_k + R_{o,j}Z_{e,k}m_k + R_{o,j}Z_{e,k}m_k) \quad \iint d\theta d\zeta \cos \alpha_i \sin \alpha_j \sin \alpha_k$$

$$+ (-R_{o,k}Z_{e,j}m_i) \quad \iint d\theta d\zeta \sin \alpha_i \cos \alpha_j \sin \alpha_k$$

$$+ (-R_{e,k}Z_{o,j}m_i) \quad \iint d\theta d\zeta \sin \alpha_i \sin \alpha_j \cos \alpha_k$$

$$(240)$$

$$3\frac{\partial V}{\partial Z_{o,i}} = (-R_{e,k}R_{e,j}m_i) \quad \iint d\theta d\zeta \cos \alpha_i \cos \alpha_j \cos \alpha_k$$

$$+ (-R_{o,k}R_{o,j}m_i) \quad \iint d\theta d\zeta \cos \alpha_i \sin \alpha_j \sin \alpha_k$$

$$+ (-R_{e,j}R_{e,k}m_k) \quad \iint d\theta d\zeta \sin \alpha_i \cos \alpha_j \sin \alpha_k$$

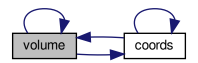
$$+ (+R_{o,j}R_{o,k}m_k) \quad \iint d\theta d\zeta \sin \alpha_i \sin \alpha_j \cos \alpha_k$$

$$(241)$$

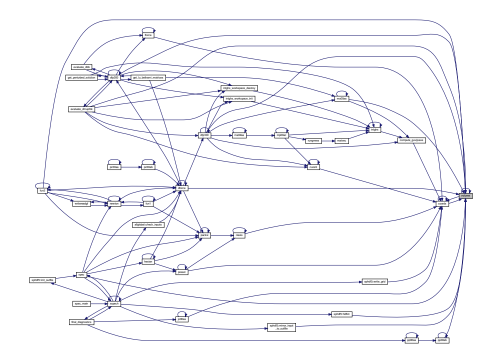
References coords(), allglobal::cosi, allglobal::cpus, allglobal::dbdx, allglobal::djkm, allglobal::djkp, allglobal::dvolume, constants::four, constants::half, inputlist::igeometry, allglobal::im, allglobal::in, allglobal::irbc, allglobal::irbc, allglobal::irbc, allglobal::irbc, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ntz, inputlist::nvol, constants::one, fileunits::ounit, constants::pi2, inputlist::pscale, constants::quart, allglobal::rij, allglobal::sini, numerical::small, constants::third, constants::two, volume(), numerical::vsmall, allglobal::vvolume, allglobal::yesstellsym, constants::zero, and allglobal::zij.

Referenced by coords(), dforce(), dfp100(), dfp200(), evaluate\_dmupfdx(), fcn2(), sphdf5::hdfint(), ma00aa(), sphdf5::mirror input to outfile(), pp00ab(), spec(), and volume().

Here is the call graph for this function:



Here is the caller graph for this function:



# 7.22 Smooth boundary

## **Functions/Subroutines**

• subroutine wa00aa (iwa00aa)

Constructs smooth approximation to wall.

· subroutine vacuumphi (Nconstraints, rho, fvec, iflag)

Compute vacuum magnetic scalar potential (?)

# 7.22.1 Detailed Description

#### 7.22.2 Function/Subroutine Documentation

# **7.22.2.1 wa00aa()** subroutine wa00aa ( integer *iwa00aa* )

Constructs smooth approximation to wall.

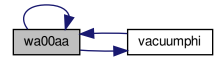
# solution of Laplace's equation in two-dimensions

- · The wall is given by a discrete set of points.
- · The points must go anti-clockwise.

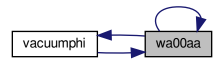
References laplaces::alpha, laplaces::cc, allglobal::cpus, laplaces::dorm, laplaces::exterior, fileunits::gunit, constants::half, laplaces::iangle, laplaces::ic, laplaces::icint, allglobal::im, allglobal::in, allglobal::irbc, allglobal::irbs, allglobal::izbc, allglobal::izbs, allglobal::lcoordinatesingularity, allglobal::mn, inputlist::mpol, allglobal::myid, allglobal::ncpu, laplaces::nintervals, laplaces::niterations, laplaces::np1, laplaces::np4, laplaces::nsegments, allglobal::nt, inputlist::ntor, allglobal::ntz, inputlist::nvol, allglobal::nz, inputlist::odetol, constants::one, laplaces::originalalpha, fileunits::ounit, laplaces::phi, laplaces::phid, constants::pi2, allglobal::rij, laplaces::rmid, laplaces::stage1, constants::ten, vacuumphi(), numerical::vsmall, wa00aa(), inputlist::wmacros, laplaces::xpoly, allglobal::yesstellsym, laplaces::ypoly, constants::zero, and allglobal::zij.

Referenced by vacuumphi(), and wa00aa().

Here is the call graph for this function:



Here is the caller graph for this function:



Compute vacuum magnetic scalar potential (?)

# **Parameters**

Nconstraints	
rho	
fvec	
iflag	

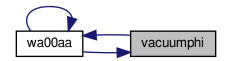
References laplaces::alpha, laplaces::cc, allglobal::cpus, laplaces::dorm, laplaces::exterior, constants::half, laplaces::iangle, laplaces::ic, laplaces::icint, allglobal::myid, allglobal::ncpu, laplaces::nintervals, laplaces::niterations, laplaces::np1, laplaces::np4, laplaces::nsegments, allglobal::ntz, constants::one, laplaces::originalalpha, fileunits::ounit, laplaces::phi, laplaces::phid, constants::pi2, allglobal::rij, laplaces::rmid, laplaces::stage1, wa00aa(), inputlist::wmacros, laplaces::xpoly, laplaces::ypoly, constants::zero, and allglobal::zij.

Referenced by wa00aa().

Here is the call graph for this function:



Here is the caller graph for this function:



## 7.23 Enhanced resolution for metric elements

Enhanced resolution is required for the metric elements,  $g_{ij}/\sqrt{g}$ , which is given by mne, ime, and ine. The Fourier resolution here is determined by 1 Mpol = 2 Mpol and 1 Ntor = 2 Ntor.

# **Variables**

- integer allglobal::mne
  - enhanced resolution for metric elements
- integer, dimension(:), allocatable allglobal::ime
   enhanced poloidal mode numbers for metric elements
- integer, dimension(:), allocatable **allglobal::ine**enhanced toroidal mode numbers for metric elements

# 7.23.1 Detailed Description

Enhanced resolution is required for the metric elements,  $g_{ij}/\sqrt{g}$ , which is given by mne, ime, and ine. The Fourier resolution here is determined by 1Mpol=2\*Mpol and 1Ntor=2\*Ntor.

# 7.24 Enhanced resolution for transformation to straight-field line angle

Enhanced resolution is required for the transformation to straight-field line angle on the interfaces, which is given by mns, ims and ins. The Fourier resolution here is determined by iMpol and iNtor.

# Variables

- integer allglobal::mns
  - enhanced resolution for straight field line transformation
- integer, dimension(:), allocatable allglobal::ims
   enhanced poloidal mode numbers for straight field line transformation

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• integer, dimension(:), allocatable **allglobal::ins**enhanced toroidal mode numbers for straight field line transformation

# 7.24.1 Detailed Description

Enhanced resolution is required for the transformation to straight-field line angle on the interfaces, which is given by mns, ims and ins. The Fourier resolution here is determined by iMpol and iNtor.

## 7.25 Internal Variables

Collaboration diagram for Internal Variables:



## **Modules**

- · Fourier representation
- · Interface geometry: iRbc, iZbs etc.

The Fourier harmonics of the interfaces are contained in iRbc(1:mn, 0:Mvol) and iZbs(1:mn, 0:Mvol), where iRbc(1, j), iZbs(1, j) contains the Fourier harmonics,  $R_j$ ,  $Z_j$ , of the l-th interface.

· Fourier Transforms

The coordinate geometry and fields are mapped to/from Fourier space and real space using FFTW3. The resolution of the real space grid is given by Nt=Ndiscrete\*4\*Mpol and Nz=Ndiscrete\*4\*Ntor.

· Volume-integrated Chebyshev-metrics

These are allocated in dforce(), defined in ma00aa(), and are used in matrix() to construct the matrices.

- · Vector potential and the Beltrami linear system
- · Field matrices: dMA, dMB, dMC, dMD, dME, dMF
- · Construction of "force"

The force vector is comprised of Bomn and Iomn.

· Covariant field on interfaces: Btemn, Bzemn, Btomn, Bzomn

The covariant field.

- · covariant field for Hessian computation: Bloweremn, Bloweromn
- Geometrical degrees-of-freedom: LGdof, NGdof

The geometrical degrees-of-freedom.

- · Parallel construction of derivative matrix
- · Derivatives of multiplier and poloidal flux with respect to geometry: dmupfdx
- Trigonometric factors
- · Volume integrals: IBBintegral, IABintegral
- · Internal global variables

internal global variables; internal logical variables; default values are provided here; these may be changed according to input values

Miscellaneous

The following are miscellaneous flags required for the virtual casing field, external (vacuum) field integration, ...

# **Variables**

type(derivative) allglobal::dbdx
 dB/dX (?)

# 7.25.1 Detailed Description

# 7.26 Fourier representation

Collaboration diagram for Fourier representation:



#### **Variables**

· integer allglobal::mn

total number of Fourier harmonics for coordinates/fields; calculated from Mpol, Ntor in readin()

• integer, dimension(:), allocatable allglobal::im

poloidal mode numbers for Fourier representation

• integer, dimension(:), allocatable allglobal::in

toroidal mode numbers for Fourier representation

• real, dimension(:), allocatable allglobal::halfmm

I saw this already somewhere...

• real, dimension(:), allocatable allglobal::regumm

I saw this already somewhere...

· real aligiobal::rscale

no idea

• real, dimension(:,:), allocatable allglobal::psifactor

no idea

• real, dimension(:,:), allocatable allglobal::inifactor

no idea

• real, dimension(:), allocatable allglobal::bbweight

weight on force-imbalance harmonics; used in dforce()

real, dimension(:), allocatable allglobal::mmpp

spectral condensation factors

### 7.26.1 Detailed Description

# 7.27 Interface geometry: iRbc, iZbs etc.

The Fourier harmonics of the interfaces are contained in iRbc(1:mn, 0:Mvol) and iZbs(1:mn, 0:Mvol), where iRbc(1,j), iZbs(1,j) contains the Fourier harmonics,  $R_j$ ,  $Z_j$ , of the l-th interface. Collaboration diagram for Interface geometry: iRbc, iZbs etc.:



# **Variables**

real, dimension(:,:), allocatable allglobal::irbc

cosine R harmonics of interface surface geometry; stellarator symmetric

• real, dimension(:,:), allocatable allglobal::izbs

sine Z harmonics of interface surface geometry; stellarator symmetric

• real, dimension(:,:), allocatable allglobal::irbs

sine R harmonics of interface surface geometry; non-stellarator symmetric

• real, dimension(:,:), allocatable allglobal::izbc

cosine Z harmonics of interface surface geometry; non-stellarator symmetric

real, dimension(:,:), allocatable allglobal::drbc

cosine R harmonics of interface surface geometry; stellarator symmetric; linear deformation

real, dimension(:,:), allocatable allglobal::dzbs

sine Z harmonics of interface surface geometry; stellarator symmetric; linear deformation

real, dimension(:,:), allocatable allglobal::drbs

sine R harmonics of interface surface geometry; non-stellarator symmetric; linear deformation

real, dimension(:,:), allocatable allglobal::dzbc

cosine Z harmonics of interface surface geometry; non-stellarator symmetric; linear deformation

real, dimension(:,:), allocatable allglobal::irij

interface surface geometry; real space

real, dimension(:,:), allocatable allglobal::izij

interface surface geometry; real space

• real, dimension(:,:), allocatable allglobal::drij

interface surface geometry; real space

• real, dimension(:,:), allocatable allglobal::dzij

interface surface geometry; real space

real, dimension(:,:), allocatable allglobal::trij

interface surface geometry; real space

• real, dimension(:,:), allocatable allglobal::tzij

interface surface geometry; real space

real, dimension(:), allocatable allglobal::ivns

sine harmonics of vacuum normal magnetic field on interfaces; stellarator symmetric

• real, dimension(:), allocatable allglobal::ibns

sine harmonics of plasma normal magnetic field on interfaces; stellarator symmetric

real, dimension(:), allocatable allglobal::ivnc

cosine harmonics of vacuum normal magnetic field on interfaces; non-stellarator symmetric

real, dimension(:), allocatable allglobal::ibnc

cosine harmonics of plasma normal magnetic field on interfaces; non-stellarator symmetric

real, dimension(:), allocatable allglobal::lrbc

local workspace

• real, dimension(:), allocatable allglobal::lzbs

local workspace

• real, dimension(:), allocatable allglobal::Irbs

local workspace

• real, dimension(:), allocatable allglobal::lzbc

local workspace

- integer allglobal::num\_modes
- integer, dimension(:), allocatable allglobal::mmrzrz
- integer, dimension(:), allocatable allglobal::nnrzrz
- real, dimension(:,:,:), allocatable allglobal::allrzrz

## 7.27.1 Detailed Description

The Fourier harmonics of the interfaces are contained in iRbc(1:mn, 0:Mvol) and iZbs(1:mn, 0:Mvol), where iRbc(1, j), iZbs(1, j) contains the Fourier harmonics,  $R_j$ ,  $Z_j$ , of the l-th interface.

## 7.28 Fourier Transforms

The coordinate geometry and fields are mapped to/from Fourier space and real space using FFTW3. The resolution of the real space grid is given by Nt=Ndiscrete\*4\*Mpol and Nz=Ndiscrete\*4\*Ntor.

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Collaboration diagram for Fourier Transforms:



#### **Variables**

• integer allglobal::nt

discrete resolution along  $\theta$  of grid in real space

• integer allglobal::nz

discrete resolution along  $\zeta$  of grid in real space

· integer allglobal::ntz

discrete resolution; Ntz=Nt\*Nz shorthand

· integer allglobal::hnt

discrete resolution; Ntz=Nt\*Nz shorthand

• integer allglobal::hnz

discrete resolution; Ntz=Nt\*Nz shorthand

· real allglobal::sontz

one / sqrt (one\*Ntz); shorthand

real, dimension(:,:,:), allocatable allglobal::rij

real-space grid; R

• real, dimension(:,:,:), allocatable allglobal::zij

real-space grid; Z

• real, dimension(:,:,:), allocatable allglobal::xij

what is this?

• real, dimension(:,:,:), allocatable allglobal::yij

what is this?

• real, dimension(:,:), allocatable allglobal::sg

real-space grid; jacobian and its derivatives

real, dimension(:,:,:,:), allocatable allglobal::guvij

real-space grid; metric elements

real, dimension(:,:,:), allocatable allglobal::gvuij

real-space grid; metric elements (?); 10 Dec 15;

• real, dimension(:,:,:,:), allocatable allglobal::guvijsave

what is this?

• integer, dimension(:,:), allocatable allglobal::ki

identification of Fourier modes

integer, dimension(:,:,:), allocatable allglobal::kijs

identification of Fourier modes

• integer, dimension(:,:,:), allocatable allglobal::kija

identification of Fourier modes

• integer, dimension(:), allocatable allglobal::iotakkii

identification of Fourier modes

• integer, dimension(:,:), allocatable allglobal::iotaksub

identification of Fourier modes

- integer, dimension(:,:), allocatable allglobal::iotakadd identification of Fourier modes
- integer, dimension(:,:), allocatable allglobal::iotaksgn identification of Fourier modes

1 12 2 13 14 15 15 15

• real, dimension(:), allocatable allglobal::efmn

Fourier harmonics; dummy workspace.

real, dimension(:), allocatable allglobal::ofmn
 Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::cfmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::sfmn

Fourier harmonics; dummy workspace.

real, dimension(:), allocatable allglobal::evmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::odmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::comn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::simn

Fourier harmonics; dummy workspace.

- real, dimension(:), allocatable allglobal::ijreal what is this?
- real, dimension(:), allocatable allglobal::ijimag
   what is this?
- real, dimension(:), allocatable allglobal::jireal what is this?
- real, dimension(:), allocatable allglobal::jiimag
   what is this?
- real, dimension(:), allocatable allglobal::jkreal what is this?
- real, dimension(:), allocatable allglobal::jkimag what is this?
- real, dimension(:), allocatable allglobal::kjreal
   what is this ?
- real, dimension(:), allocatable allglobal::kjimag what is this?
- real, dimension(:,:,:), allocatable allglobal::bsupumn

tangential field on interfaces;  $\theta$ -component; required for virtual casing construction of field; 11 Oct 12

real, dimension(:,:,:), allocatable allglobal::bsupvmn

tangential field on interfaces;  $\zeta$  -component; required for virtual casing construction of field; 11 Oct 12

 real, dimension(:,:), allocatable allglobal::goomne described in preset()

• real, dimension(:,:), allocatable allglobal::goomno

described in preset()

• real, dimension(:,:), allocatable allglobal::gssmne

described in preset()

real, dimension(:,:), allocatable allglobal::gssmno

described in preset()

real, dimension(:,:), allocatable allglobal::gstmne

described in preset()

real, dimension(:,:), allocatable allglobal::gstmno

described in preset()

- real, dimension(:,:), allocatable allglobal::gszmne described in preset()
- real, dimension(:,:), allocatable allglobal::gszmno described in preset()
- real, dimension(:,:), allocatable allglobal::gttmne described in preset()
- real, dimension(:,:), allocatable allglobal::gttmno described in preset()
- real, dimension(:,:), allocatable allglobal::gtzmne described in preset()
- real, dimension(:,:), allocatable allglobal::gtzmno described in preset()
- real, dimension(:,:), allocatable allglobal::gzzmne described in preset()
- real, dimension(:,:), allocatable allglobal::gzzmno described in preset()

#### 7.28.1 Detailed Description

The coordinate geometry and fields are mapped to/from Fourier space and real space using FFTW3. The resolution of the real space grid is given by Nt=Ndiscrete\*4\*Mpol and Nz=Ndiscrete\*4\*Mtor. Various workspace arrays are allocated. These include Rij(1:Ntz,0:3,0:3) and Zij(1:Ntz,0:3,0:3), which contain the coordinates in real space and their derivatives; sg(0:3,Ntz), which contains the Jacobian and its derivatives; and guv(0:6,0:3,1:Ntz), which contains the metric elements and their derivatives.

## 7.29 Volume-integrated Chebyshev-metrics

These are allocated in dforce(), defined in ma00aa(), and are used in matrix() to construct the matrices. Collaboration diagram for Volume-integrated Chebyshev-metrics:



#### **Variables**

- real, dimension(:,:,:,:), allocatable allglobal::dtoocc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,;;;:), allocatable allglobal::dtoocs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::dtoosc
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::dtooss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ttsscc volume-integrated Chebychev-metrics; see matrix()

- real, dimension(:,:,:,:), allocatable allglobal::ttsscs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ttsssc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ttssss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdstcc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdstcs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdstsc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdstss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdszcc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdszcs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdszsc
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:), allocatable allglobal::tdszss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddttcc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddttcs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddttsc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddttss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddtzcc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddtzcs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddtzsc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddtzss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddzzcc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddzzcs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddzzsc
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddzzss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:), allocatable allglobal::tsc what is this?
- real, dimension(:,:), allocatable allglobal::tss what is this?
- · real, dimension(:,:), allocatable allglobal::dtc

what is this?

• real, dimension(:,:), allocatable allglobal::dts

what is this?

real, dimension(:,:), allocatable allglobal::dzc what is this?

real, dimension(:,:), allocatable allglobal::dzs what is this?

real, dimension(:,:), allocatable allglobal::ttc what is this?

real, dimension(:,:), allocatable allglobal::tzc what is this?

real, dimension(:,:), allocatable allglobal::tts what is this?

real, dimension(:,:), allocatable allglobal::tzs what is this?

• real, dimension(:), allocatable allglobal::dtflux

 $\delta \psi_{toroidal}$  in each annulus

• real, dimension(:), allocatable allglobal::dpflux

 $\delta\psi_{poloidal}$  in each annulus

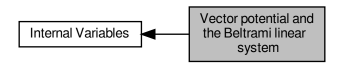
real, dimension(:), allocatable allglobal::sweight
 minimum poloidal length constraint weight

#### 7.29.1 Detailed Description

These are allocated in dforce(), defined in ma00aa(), and are used in matrix() to construct the matrices.

# 7.30 Vector potential and the Beltrami linear system

Collaboration diagram for Vector potential and the Beltrami linear system:



## **Variables**

- integer, dimension(:), allocatable allglobal::nadof degrees of freedom in Beltrami fields in each annulus
- integer, dimension(:), allocatable allglobal::nfielddof
   degrees of freedom in Beltrami fields in each annulus, field only, no Lagrange multipliers
- type(subgrid), dimension(:,:,:), allocatable allglobal::ate
   magnetic vector potential cosine Fourier harmonics; stellarator-symmetric
- type(subgrid), dimension(:,:,:), allocatable allglobal::aze
   magnetic vector potential cosine Fourier harmonics; stellarator-symmetric
- type(subgrid), dimension(:,:,:), allocatable allglobal::ato
   magnetic vector potential sine Fourier harmonics; non-stellarator-symmetric

• type(subgrid), dimension(:,:,:), allocatable allglobal::azo magnetic vector potential sine Fourier harmonics; non-stellarator-symmetric • integer, dimension(:,:), allocatable allglobal::Ima Lagrange multipliers (?) integer, dimension(:,:), allocatable allglobal::lmb Lagrange multipliers (?) • integer, dimension(:,:), allocatable allglobal::Imc Lagrange multipliers (?) integer, dimension(:,:), allocatable allglobal::Imd Lagrange multipliers (?) • integer, dimension(:,:), allocatable allglobal::Ime Lagrange multipliers (?) • integer, dimension(:,:), allocatable allglobal::Imf Lagrange multipliers (?) • integer, dimension(:,:), allocatable allglobal::Img Lagrange multipliers (?) • integer, dimension(:,:), allocatable allglobal::Imh Lagrange multipliers (?) • real, dimension(:,:), allocatable allglobal::Imavalue what is this? real, dimension(:,:), allocatable allglobal::Imbvalue what is this? • real, dimension(:,:), allocatable allglobal::Imcvalue what is this? • real, dimension(:,:), allocatable allglobal::Imdvalue what is this? real, dimension(:,:), allocatable allglobal::Imevalue what is this? real, dimension(:,:), allocatable allglobal::Imfvalue what is this? • real, dimension(:,:), allocatable allglobal::Imgvalue what is this? • real, dimension(:,:), allocatable allglobal::Imhvalue what is this? integer, dimension(:,:), allocatable allglobal::fso what is this? integer, dimension(:,:), allocatable allglobal::fse what is this? logical allglobal::lcoordinatesingularity set by LREGION macro; true if inside the innermost volume · logical allglobal::lplasmaregion set by LREGION macro; true if inside the plasma region · logical allglobal::lvacuumregion set by LREGION macro; true if inside the vacuum region

 logical allglobal::lsavedguvij flag used in matrix free · logical allglobal::localconstraint what is this?

## 7.30.1 Detailed Description

- In each volume, the total degrees of freedom in the Beltrami linear system is NAdof(1:Nvol). This depends on Mpol, Ntor and Lrad(vvol).
- The covariant components of the vector potential are written as

$$A_{\theta} = \sum_{i} \sum_{l=0}^{L} A_{\theta,e,i,l} T_{l}(s) \cos \alpha_{i} + \sum_{i} \sum_{l=0}^{L} A_{\theta,o,i,l} T_{l}(s) \sin \alpha_{i}$$
 (242)

$$A_{\zeta} = \sum_{i} \sum_{l=0}^{L} A_{\zeta,e,i,l} T_{l}(s) \cos \alpha_{i} + \sum_{i} \sum_{l=0}^{L} A_{\zeta,o,i,l} T_{l}(s) \sin \alpha_{i},$$
 (243)

where  $T_l(s)$  are the Chebyshev polynomials and  $\alpha_i \equiv m_i \theta - n_i \zeta$ .

• The following internal arrays are declared in preset():

dAte (0, i) %s(I) 
$$\equiv A_{\theta,e,i,l}$$
 dAze (0, i) %s(I)  $\equiv A_{\zeta,e,i,l}$  dAto (0, i) %s(I)  $\equiv A_{\theta,o,i,l}$  dAzo (0, i) %s(I)  $\equiv A_{\zeta,o,i,l}$ 

# 7.31 Field matrices: dMA, dMB, dMC, dMD, dME, dMF

Collaboration diagram for Field matrices: dMA, dMB, dMC, dMD, dME, dMF:



## **Variables**

- real, dimension(:,:), allocatable allglobal::dma energy and helicity matrices; quadratic forms
- real, dimension(:,:), allocatable allglobal::dmb
   energy and helicity matrices; quadratic forms
- real, dimension(:,:), allocatable allglobal::dmd
   energy and helicity matrices; quadratic forms
- real, dimension(:), allocatable allglobal::dmas sparse version of dMA, data
- real, dimension(:), allocatable allglobal::dmds sparse version of dMD, data
- integer, dimension(:), allocatable allglobal::idmas
   sparse version of dMA and dMD, indices
- integer, dimension(:), allocatable allglobal::jdmas sparse version of dMA and dMD, indices
- integer, dimension(:), allocatable allglobal::ndmasmax number of elements for sparse matrices
- integer, dimension(:), allocatable allglobal::ndmas

number of elements for sparse matrices

· real, dimension(:), allocatable allglobal::dmg

what is this?

• real, dimension(:), allocatable allglobal::adotx

the matrix-vector product

• real, dimension(:), allocatable allglobal::ddotx

the matrix-vector product

real, dimension(:,:), allocatable allglobal::solution

this is allocated in dforce; used in mp00ac and ma02aa; and is passed to packab

• real, dimension(:,:,:), allocatable allglobal::gmreslastsolution

used to store the last solution for restarting GMRES

· real, dimension(:), allocatable allglobal::mbpsi

matrix vector products

· logical allglobal::liluprecond

whether to use ILU preconditioner for GMRES

real, dimension(:,:), allocatable allglobal::beltramiinverse

Beltrami inverse matrix.

real, dimension(:,:,:), allocatable allglobal::diotadxup

measured rotational transform on inner/outer interfaces for each volume; d(transform)/dx; (see dforce)

real, dimension(:,:,:), allocatable allglobal::ditgpdxtp

measured toroidal and poloidal current on inner/outer interfaces for each volume; d(ltor,Gpol)/dx; (see dforce)

real, dimension(:,:,:,:), allocatable allglobal::glambda

save initial guesses for iterative calculation of rotational-transform

integer allglobal::lmns

what is this?

## 7.31.1 Detailed Description

• The energy,  $W \equiv \int dv \mathbf{B} \cdot \mathbf{B}$ , and helicity,  $K \equiv \int dv \mathbf{A} \cdot \mathbf{B}$ , functionals may be written

$$W = \frac{1}{2} a_i A_{i,j} a_j + a_i B_{i,j} \psi_j + \frac{1}{2} \psi_i C_{i,j} \psi_j$$

$$K = \frac{1}{2} a_i D_{i,j} a_j + a_i E_{i,j} \psi_j + \frac{1}{2} \psi_i F_{i,j} \psi_j$$
(244)

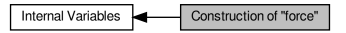
$$K = \frac{1}{2} a_i D_{i,j} a_j + a_i E_{i,j} \psi_j + \frac{1}{2} \psi_i F_{i,j} \psi_j$$
 (245)

where  $\mathbf{a} \equiv \{A_{\theta,e,i,l}, A_{\zeta,e,i,l}, A_{\theta,o,i,l}, A_{\zeta,o,i,l}, f_{e,i}, f_{o,i}\}$  contains the independent degrees of freedom and  $\psi \equiv \{\Delta\psi_t, \Delta\psi_p\}$ .

• These are allocated and deallocated in dforce(), assigned in matrix(), and used in mp00ac() and (?) df00aa().

# 7.32 Construction of "force"

The force vector is comprised of Bomn and Iomn. Collaboration diagram for Construction of "force":



#### **Variables**

- real, dimension(:,:,:), allocatable allglobal::bemn force vector; stellarator-symmetric (?)
- real, dimension(:,:), allocatable allglobal::iomn force vector; stellarator-symmetric (?)
- real, dimension(:,;,:), allocatable allglobal::somn force vector; non-stellarator-symmetric (?)
- real, dimension(:,:,:), allocatable allglobal::pomn force vector; non-stellarator-symmetric (?)
- real, dimension(:,:,:), allocatable allglobal::bomn force vector; stellarator-symmetric (?)
- real, dimension(:,:), allocatable allglobal::iemn force vector; stellarator-symmetric (?)
- real, dimension(:,:,:), allocatable allglobal::semn force vector; non-stellarator-symmetric (?)
- real, dimension(:,:,:), allocatable allglobal::pemn force vector; non-stellarator-symmetric (?)
- real, dimension(:), allocatable allglobal::bbe force vector (?); stellarator-symmetric (?)
- real, dimension(:), allocatable allglobal::iio force vector (?); stellarator-symmetric (?)
- real, dimension(:), allocatable allglobal::bbo force vector (?); non-stellarator-symmetric (?)
- real, dimension(:), allocatable allglobal::iie
   force vector (?); non-stellarator-symmetric (?)

# 7.32.1 Detailed Description

The force vector is comprised of Bomn and Iomn.

#### 7.33 Covariant field on interfaces: Btemn, Bzemn, Btomn, Bzomn

The covariant field.

Collaboration diagram for Covariant field on interfaces: Btemn, Bzemn, Btomn, Bzomn:



### **Variables**

- real, dimension(:,:,:), allocatable allglobal::btemn
   covariant θ cosine component of the tangential field on interfaces; stellarator-symmetric
- real, dimension(:,:,:), allocatable allglobal::bzemn
   covariant ζ cosine component of the tangential field on interfaces; stellarator-symmetric
- real, dimension(:,:,:), allocatable allglobal::btomn

covariant  $\theta$  sine component of the tangential field on interfaces; non-stellarator-symmetric

• real, dimension(:,:,:), allocatable allglobal::bzomn

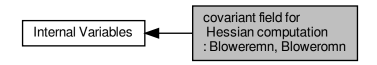
covariant  $\zeta$  sine component of the tangential field on interfaces; non-stellarator-symmetric

#### 7.33.1 Detailed Description

The covariant field.

# 7.34 covariant field for Hessian computation: Bloweremn, Bloweromn

Collaboration diagram for covariant field for Hessian computation: Bloweremn, Bloweremn:



#### **Variables**

- real, dimension(:,:), allocatable allglobal::bloweremn covariant field for Hessian computation
- real, dimension(:,:), allocatable allglobal::bloweromn covariant field for Hessian computation

## 7.34.1 Detailed Description

# 7.35 Geometrical degrees-of-freedom: LGdof, NGdof

The geometrical degrees-of-freedom.

Collaboration diagram for Geometrical degrees-of-freedom: LGdof, NGdof:



#### **Variables**

• integer allglobal::lgdof

geometrical degrees of freedom associated with each interface

• integer allglobal::ngdof

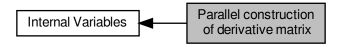
total geometrical degrees of freedom

## 7.35.1 Detailed Description

The geometrical degrees-of-freedom.

## 7.36 Parallel construction of derivative matrix

Collaboration diagram for Parallel construction of derivative matrix:



#### **Variables**

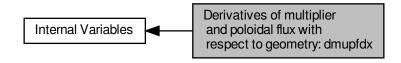
- real, dimension(:,:,:), allocatable allglobal::dbbdrz
   derivative of magnetic field w.r.t. geometry (?)
- real, dimension(:,:), allocatable allglobal::diidrz
   derivative of spectral constraints w.r.t. geometry (?)
- real, dimension(:,:,:,:), allocatable allglobal::dffdrz
   derivatives of B<sup>2</sup> at the interfaces wrt geometry
- real, dimension(:,:,:), allocatable allglobal::dbbdmp derivatives of B<sup>2</sup> at the interfaces wrt mu and dpflux

#### 7.36.1 Detailed Description

- The derivatives of force-balance,  $[[p+B^2/2]]$ , and the spectral constraints (see sw03aa()), with respect to the interface geometry is constructed in parallel by dforce().
- force-balance across the *l*-th interface depends on the fields in the adjacent interfaces.

# 7.37 Derivatives of multiplier and poloidal flux with respect to geometry: dmupfdx

Collaboration diagram for Derivatives of multiplier and poloidal flux with respect to geometry: dmupfdx:



#### **Variables**

- real, dimension(:,:,:,:), allocatable allglobal::dmupfdx
   derivatives of mu and dpflux wrt geometry at constant interface transform
- logical allglobal::lhessianallocated

flag to indicate that force gradient matrix is allocated (?)

• real, dimension(:,:), allocatable allglobal::hessian

force gradient matrix (?)

• real, dimension(:,:), allocatable allglobal::dessian

derivative of force gradient matrix (?)

#### 7.37.1 Detailed Description

- The information in dmupfdx describes how the helicity multiplier,  $\mu$ , and the enclosed poloidal flux,  $\Delta \psi_p$ , must vary as the geometry is varied in order to satisfy the interface transform constraint.
- The internal variable dmupfdx (1:Mvol, 1:2, 1:LGdof, 0:1) is allocated/deallocated in newton(), and hesian() if selected.
- The magnetic field depends on the Fourier harmonics of both the inner and outer interface geometry (represented here as  $x_j$ ), the helicity multiplier, and the enclosed poloidal flux, i.e.  $\mathbf{B}_{\pm} = \mathbf{B}_{\pm}(x_j, \mu, \Delta \psi_p)$ , so that

$$\delta \mathbf{B}_{\pm} = \frac{\partial \mathbf{B}_{\pm}}{\partial x_{j}} \delta x_{j} + \frac{\partial \mathbf{B}_{\pm}}{\partial \mu} \delta \mu + \frac{\partial \mathbf{B}_{\pm}}{\partial \Delta \psi_{p}} \delta \Delta \psi_{p}. \tag{246}$$

• This information is used to adjust the calculation of how force-balance, i.e.  $B^2$  at the interfaces, varies with geometry at fixed interface rotational transform. Given

$$B_{\pm}^{2} = B_{\pm}^{2}(x_{j}, \mu, \Delta \psi_{p}), \tag{247}$$

we may derive

$$\frac{\partial B_{\pm}^{2}}{\partial x_{i}} = \frac{\partial B_{\pm}^{2}}{\partial x_{i}} + \frac{\partial B_{\pm}^{2}}{\partial \mu} \frac{\partial \mu}{\partial x_{i}} + \frac{\partial B_{\pm}^{2}}{\partial \Delta \psi_{p}} \frac{\partial \Delta \psi_{p}}{\partial x_{i}}$$
(248)

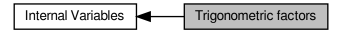
• The constraint to be enforced is that  $\mu$  and  $\Delta \psi_p$  must generally vary as the geometry is varied if the value of the rotational-transform constraint on the inner/outer interface is to be preserved, i.e.

$$\begin{pmatrix}
\frac{\partial \boldsymbol{t}_{-}}{\partial \mathbf{B}_{-}} \cdot \frac{\partial \mathbf{B}_{-}}{\partial \mu} & , & \frac{\partial \boldsymbol{t}_{-}}{\partial \mathbf{B}_{-}} \cdot \frac{\partial \mathbf{B}_{-}}{\partial \Delta \psi_{p}} \\
\frac{\partial \boldsymbol{t}_{+}}{\partial \mathbf{B}_{+}} \cdot \frac{\partial \mathbf{B}_{+}}{\partial \mu} & , & \frac{\partial \boldsymbol{t}_{+}}{\partial \mathbf{B}_{+}} \cdot \frac{\partial \mathbf{B}_{+}}{\partial \Delta \psi_{p}}
\end{pmatrix}
\begin{pmatrix}
\frac{\partial \mu}{\partial x_{j}} \\
\frac{\partial \Delta \psi_{p}}{\partial x_{j}}
\end{pmatrix} = - \begin{pmatrix}
\frac{\partial \boldsymbol{t}_{-}}{\partial \mathbf{B}_{-}} \cdot \frac{\partial \mathbf{B}_{-}}{\partial x_{j}} \\
\frac{\partial \boldsymbol{t}_{+}}{\partial \mathbf{B}_{+}} \cdot \frac{\partial \mathbf{B}_{+}}{\partial x_{j}}
\end{pmatrix}.$$
(249)

- This  $2 \times 2$  linear equation is solved in dforce() and the derivatives of the rotational-transform are given in diotadxup, see preset.f90.
- A finite-difference estimate is computed if Lcheck==4.

## 7.38 Trigonometric factors

Collaboration diagram for Trigonometric factors:



#### **Variables**

- real, dimension(:,:), allocatable allglobal::cosi some precomputed cosines
- real, dimension(:,:), allocatable allglobal::sini some precomputed sines
- real, dimension(:), allocatable **allglobal::gteta** something related to  $\sqrt{g}$  and  $\theta$ ?
- real, dimension(:), allocatable allglobal::gzeta something related to  $\sqrt{g}$  and  $\zeta$  ?
- real, dimension(:), allocatable **allglobal::ajk**definition of coordinate axis
- real, dimension(:,:,:,:), allocatable allglobal::dradr derivatives of coordinate axis
- real, dimension(:,:,:,:), allocatable allglobal::dradz derivatives of coordinate axis
- real, dimension(:,:,:,:), allocatable allglobal::dzadr derivatives of coordinate axis
- real, dimension(:,:,:,:), allocatable allglobal::dzadz derivatives of coordinate axis
- real, dimension(:,:,:), allocatable allglobal::drodr derivatives of coordinate axis
- real, dimension(:,:,:), allocatable allglobal::drodz derivatives of coordinate axis
- real, dimension(:,:,:), allocatable allglobal::dzodr derivatives of coordinate axis
- real, dimension(:,:,:), allocatable allglobal::dzodz derivatives of coordinate axis
- integer, dimension(:,:), allocatable allglobal::djkp for calculating cylindrical volume
- integer, dimension(:,:), allocatable allglobal::djkm for calculating cylindrical volume

#### 7.38.1 Detailed Description

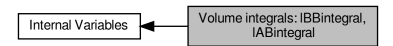
- To facilitate construction of the metric integrals, various trigonometric identities are exploited.
- The following are used for volume integrals (see volume() ):

$$a_{i,j,k} = 4 m_k \oint d\theta d\zeta \cos(\alpha_i) \cos(\alpha_j) \cos(\alpha_k) / (2\pi)^2,$$
 (250)

$$b_{i,j,k} = 4 m_j \oint d\theta d\zeta \cos(\alpha_i) \sin(\alpha_j) \sin(\alpha_k) / (2\pi)^2,$$
 (251)

# 7.39 Volume integrals: IBBintegral, IABintegral

Collaboration diagram for Volume integrals: IBBintegral, IABintegral:



#### **Variables**

- real, dimension(:), allocatable allglobal::lbbintegral
   B.B integral.
- real, dimension(:), allocatable allglobal::labintegral
   A.B integral.
- real, dimension(:), allocatable **allglobal::vvolume** volume integral of  $\sqrt{g}$ ; computed in volume
- real aligiobal::dvolume

derivative of volume w.r.t. interface geometry

#### 7.39.1 Detailed Description

• The energy functional,  $F \equiv \sum_{l} F_{l}$ , where

$$F_{l} \equiv \left( \int_{\mathcal{V}_{l}} \frac{p_{l}}{\gamma - 1} + \frac{B_{l}^{2}}{2} dv \right) = \frac{P_{l}}{\gamma - 1} V_{l}^{1 - \gamma} + \int_{\mathcal{V}_{l}} \frac{B_{l}^{2}}{2} dv, \tag{252}$$

where the second expression is derived using  $p_l V_l^{\gamma} = P_l$ , where  $P_l$  is the adiabatic-constant. In Eqn. (252), it is implicit that  ${\bf B}$  satisfies (i) the toroidal and poloidal flux constraints; (ii) the interface constraint,  ${\bf B} \cdot \nabla s = 0$ ; and (iii) the helicity constraint (or the transform constraint).

• The derivatives of  $F_l$  with respect to the inner and outer adjacent interface geometry are stored in  $dFF(1 \leftarrow :Nvol,0:1,0:mn+mn-1)$ , where

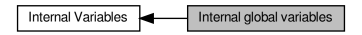
$$\begin{split} F_l & \equiv \text{dFF} \, (\text{1,0,0}) \\ \partial F_l / \partial R_{l-1,j} & \equiv \text{dFF} \, (\text{11,0,j}) \\ \partial F_l / \partial Z_{l-1,j} & \equiv \text{dFF} \, (\text{11,0,mn j}) \\ \partial F_l / \partial R_{l,j} & \equiv \text{dFF} \, (\text{11,1,j}) \\ \partial F_l / \partial Z_{l,j} & \equiv \text{dFF} \, (\text{11,1,mn j}) \end{split}$$

• The volume integrals  $\int dv$ ,  $\int B^2 dv$  and  $\int \mathbf{A} \cdot \mathbf{B} dv$  in each volume are computed and saved in volume (0  $\leftarrow$  :2,1:Nvol).

# 7.40 Internal global variables

internal global variables; internal logical variables; default values are provided here; these may be changed according to input values

Collaboration diagram for Internal global variables:



## Variables

integer allglobal::ivol

labels volume; some subroutines (called by NAG) are fixed argument list but require the volume label

real aliglobal::gbzeta

toroidal (contravariant) field; calculated in bfield; required to convert  $\dot{\theta}$  to  $B^{\theta}$ ,  $\dot{s}$  to  $B^{s}$ 

· integer, dimension(:), allocatable allglobal::iquad

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internal copy of Nquad

• real, dimension(:,:), allocatable allglobal::gaussianweight

weights for Gaussian quadrature

• real, dimension(:,:), allocatable allglobal::gaussianabscissae

abscissae for Gaussian quadrature

· logical allglobal::lblinear

controls selection of Beltrami field solver; depends on LBeltrami

· logical allglobal::Ibnewton

controls selection of Beltrami field solver; depends on LBeltrami

· logical allglobal::lbsequad

controls selection of Beltrami field solver; depends on LBeltrami

real, dimension(1:3) allglobal::orzp

used in mg00aa() to determine  $(s,\theta,\zeta)$  given  $(R,Z,\varphi)$ 

## 7.40.1 Detailed Description

internal global variables; internal logical variables; default values are provided here; these may be changed according to input values

#### 7.41 Miscellaneous

The following are miscellaneous flags required for the virtual casing field, external (vacuum) field integration, ... Collaboration diagram for Miscellaneous:



## Variables

· integer allglobal::globaljk

labels position

real, dimension(:,:), allocatable allglobal::dxyz

computational boundary; position

real, dimension(:,:), allocatable allglobal::nxyz

computational boundary; normal

real, dimension(:,:), allocatable allglobal::jxyz

plasma boundary; surface current

real, dimension(1:2) allglobal::tetazeta

what is this?

• real allglobal::virtualcasingfactor = -one / (four\*pi)

this agrees with diagno

integer allglobal::iberror

for computing error in magnetic field

· integer allglobal::nfreeboundaryiterations

number of free-boundary iterations already performed

• integer, parameter allglobal::node = 2

best to make this global for consistency between calling and called routines

• logical allglobal::first\_free\_bound = .false.

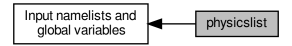
flag to indicate that this is the first free-boundary iteration

## 7.41.1 Detailed Description

The following are miscellaneous flags required for the virtual casing field, external (vacuum) field integration, ...

# 7.42 physicslist

The namelist physicslist controls the geometry, profiles, and numerical resolution. Collaboration diagram for physicslist:



#### **Variables**

```
• integer inputlist::igeometry = 3
```

selects Cartesian, cylindrical or toroidal geometry;

• integer inputlist::istellsym = 1

 $stellarator\ symmetry\ is\ enforced\ if\ Istellsym==1$ 

• integer inputlist::lfreebound = 0

compute vacuum field surrounding plasma

• real inputlist::phiedge = 1.0

total enclosed toroidal magnetic flux;

• real inputlist::curtor = 0.0

total enclosed (toroidal) plasma current;

• real inputlist::curpol = 0.0

total enclosed (poloidal) linking current;

• real inputlist::gamma = 0.0

adiabatic index; cannot set  $|\gamma|=1$ 

• integer inputlist::nfp = 1

field periodicity

integer inputlist::nvol = 1

number of volumes

• integer inputlist::mpol = 0

number of poloidal Fourier harmonics

• integer inputlist::ntor = 0

number of toroidal Fourier harmonics

• integer, dimension(1:mnvol+1) inputlist::lrad = 4

Chebyshev resolution in each volume.

• integer inputlist::lconstraint = -1

selects constraints; primarily used in ma02aa() and mp00ac().

• real, dimension(1:mnvol+1) inputlist::tflux = 0.0

toroidal flux,  $\psi_t$ , enclosed by each interface

real, dimension(1:mnvol+1) inputlist::pflux = 0.0

poloidal flux,  $\psi_p$ , enclosed by each interface

real, dimension(1:mnvol) inputlist::helicity = 0.0

helicity, K, in each volume,  $V_i$ 

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```
real inputlist::pscale = 0.0
      pressure scale factor

    real, dimension(1:mnvol+1) inputlist::pressure = 0.0

      pressure in each volume
• integer inputlist::ladiabatic = 0
      logical flag

    real, dimension(1:mnvol+1) inputlist::adiabatic = 0.0

      adiabatic constants in each volume
real, dimension(1:mnvol+1) inputlist::mu = 0.0
      helicity-multiplier, \mu, in each volume
real, dimension(1:mnvol+1) inputlist::ivolume = 0.0
       Toroidal current constraint normalized by \mu_0 ( I_{volume} = \mu_0 \cdot [A]), in each volume. This is a cumulative quantity:
       I_{\mathcal{V},i} = \int_0^{\psi_{t,i}} \mathbf{J} \cdot \mathbf{dS}. Physically, it represents the sum of all non-pressure driven currents.
real, dimension(1:mnvol) inputlist::isurf = 0.0
       Toroidal current normalized by \mu_0 at each interface (cumulative). This is the sum of all pressure driven currents.

    integer, dimension(0:mnvol) inputlist::pl = 0

       "inside" interface rotational-transform is \iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.

    integer, dimension(0:mnvol) inputlist::ql = 0

       "inside" interface rotational-transform is \iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.
• integer, dimension(0:mnvol) inputlist::pr = 0
       "inside" interface rotational-transform is \iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.
• integer, dimension(0:mnvol) inputlist::qr = 0
       "inside" interface rotational-transform is \iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.

    real, dimension(0:mnvol) inputlist::iota = 0.0

      rotational-transform, ε, on inner side of each interface
• integer, dimension(0:mnvol) inputlist::lp = 0
       "outer" interface rotational-transform is \,\iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.

    integer, dimension(0:mnvol) inputlist::lq = 0

       "outer" interface rotational-transform is \,\iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.
• integer, dimension(0:mnvol) inputlist::rp = 0
       "outer" interface rotational-transform is \,\iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.

    integer, dimension(0:mnvol) inputlist::rq = 0

       "outer" interface rotational-transform is \,\iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.
• real, dimension(0:mnvol) inputlist::oita = 0.0
      rotational-transform, ε, on outer side of each interface

 real inputlist::mupftol = 1.0e-14

      accuracy to which \mu and \Delta\psi_p are required
• integer inputlist::mupfits = 8
      an upper limit on the transform/helicity constraint iterations;
real inputlist::rpol = 1.0
      poloidal extent of slab (effective radius)
real inputlist::rtor = 1.0
      toroidal extent of slab (effective radius)
• integer inputlist::lreflect = 0
      =1 reflect the upper and lower bound in slab, =0 do not reflect
• real, dimension(0:mntor) inputlist::rac = 0.0
      stellarator symmetric coordinate axis;
• real, dimension( 0:mntor) inputlist::zas = 0.0
      stellarator symmetric coordinate axis;
  real, dimension(0:mntor) inputlist::ras = 0.0
```

non-stellarator symmetric coordinate axis;

- real, dimension( 0:mntor) inputlist::zac = 0.0
   non-stellarator symmetric coordinate axis;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::rbc = 0.0 stellarator symmetric boundary components:
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::zbs = 0.0 stellarator symmetric boundary components;
- real, dimension(-mntor:mntor,-mmpol:mmpol) **inputlist::rbs** = 0.0 non-stellarator symmetric boundary components;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::zbc = 0.0 non-stellarator symmetric boundary components;
- real, dimension(-mntor:mntor,-mmpol:mmpol) **inputlist::rwc** = 0.0 stellarator symmetric boundary components of wall;
- real, dimension(-mntor:mntor,-mmpol:mmpol) **inputlist::zws** = 0.0 stellarator symmetric boundary components of wall;
- real, dimension(-mntor:mntor,-mmpol:mmpol) **inputlist::rws** = 0.0 non-stellarator symmetric boundary components of wall;
- real, dimension(-mntor:mntor,-mmpol:mmpol) **inputlist::zwc** = 0.0 non-stellarator symmetric boundary components of wall;
- real, dimension(-mntor:mntor,-mmpol:mmpol) **inputlist::vns** = 0.0 stellarator symmetric normal field at boundary; vacuum component;
- real, dimension(-mntor:mntor,-mmpol:mmpol) **inputlist::bns** = 0.0 stellarator symmetric normal field at boundary; plasma component;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::vnc = 0.0
   non-stellarator symmetric normal field at boundary; vacuum component;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::bnc = 0.0
   non-stellarator symmetric normal field at boundary; plasma component;

## 7.42.1 Detailed Description

The namelist physicslist controls the geometry, profiles, and numerical resolution.

#### 7.42.2 Variable Documentation

# **7.42.2.1 igeometry** integer inputlist::igeometry = 3 selects Cartesian, cylindrical or toroidal geometry;

- Igeometry=1 : Cartesian; geometry determined by R;
- Igeometry=2 : cylindrical; geometry determined by R;
- Igeometry=3: toroidal; geometry determined by R and Z;

Referenced by bnorml(), allglobal::broadcast\_inputs(), allglobal::check\_inputs(), coords(), dfpred(), dfpred(), dfpred(), dfpred(), fcnred(), fcnr

# **7.42.2.2 nfp** integer inputlist::nfp = 1 field periodicity

- all Fourier representations are of the form  $\cos(m\theta nN\zeta)$ ,  $\sin(m\theta nN\zeta)$ , where  $N \equiv \text{Nfp}$
- constraint: Nfp >= 1

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), jo00aa(), sphdf5::mirror\_input\_to\_outfile(), preset(), ra00aa(), spec(), and allglobal::wrtend().

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**7.42.2.3 nvol** integer inputlist::nvol = 1 number of volumes

- each volume  $\mathcal{V}_l$  is bounded by the  $\mathcal{I}_{l-1}$  and  $\mathcal{I}_l$  interfaces
- note that in cylindrical or toroidal geometry,  $\mathcal{I}_0$  is the degenerate coordinate axis
- constraint: Nvol<=MNvol

Referenced by brcast(), allglobal::broadcast\_inputs(), allglobal::check\_inputs(), df00ab(), dforce(), dfp100(), dfp200(), dvcfield(), evaluate\_dbb(), evaluate\_dmupfdx(), fcn1(), fcn2(), final\_diagnostics(), sphdf5::hdfint(), hesian(), jo00aa(), lforce(), sphdf5::mirror\_input\_to\_outfile(), newton(), packxi(), pc00ab(), pp00aa(), pp00ab(), preset(), spec(), stzxyz(), tr00ab(), volume(), wa00aa(), sphdf5::write\_grid(), writereadgf(), and allglobal::wrtend().

**7.42.2.4 mpol** integer inputlist::mpol = 0 number of poloidal Fourier harmonics

· all Fourier representations of doubly-periodic functions are of the form

$$f(\theta,\zeta) = \sum_{n=0}^{\text{Ntor}} f_{0,n} \cos(-n \operatorname{Nfp} \zeta) + \sum_{m=1}^{\text{Mpol}} \sum_{n=-\text{Ntor}}^{\text{Ntor}} f_{m,n} \cos(m\theta - n \operatorname{Nfp} \zeta),$$
 (253)

Internally these "double" summations are written as a "single" summation, e.g.  $f(\theta, \zeta) = \sum_j f_j \cos(m_j \theta - n_j \zeta)$ .

Referenced by allocate\_geometry\_matrices(), bfield(), bfield\_tangent(), allglobal::broadcast\_inputs(), allglobal::check\_inputs(), dfp200(), intghs(), intghs\_workspace\_init(), jo00aa(), ma00aa(), matrix(), sphdf5::mirror\_input\_to\_outfile(), mtrxhs(), preset(), ra00aa(), spsint(), spsmat(), tr00ab(), wa00aa(), writereadgf(), and allglobal::wrtend().

**7.42.2.5 ntor** integer inputlist::ntor = 0 number of toroidal Fourier harmonics

• all Fourier representations of doubly-periodic functions are of the form

$$f(\theta,\zeta) = \sum_{n=0}^{\text{Ntor}} f_{0,n} \cos(-n \operatorname{Nfp} \zeta) + \sum_{m=1}^{\text{Mpol}} \sum_{n=-\text{Ntor}}^{\text{Ntor}} f_{m,n} \cos(m\theta - n \operatorname{Nfp} \zeta), \tag{254}$$

Internally these "double" summations are written as a "single" summation, e.g.  $f(\theta, \zeta) = \sum_j f_j \cos(m_j \theta - n_j \zeta)$ .

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), coords(), dforce(), dfp200(), evaluate\_dbb(), sphdf5::mirror\_input\_to\_outfile(), mp00ac(), packxi(), preset(), ra00aa(), rzaxis(), stzxyz(), tr00ab(), wa00aa(), writereadgf(), and allglobal::wrtend().

**7.42.2.6 Irad** integer, dimension(1:mnvol+1) inputlist::lrad = 4 Chebyshev resolution in each volume.

• constraint : Lrad(1:Mvol) >= 2

Referenced by allocate\_geometry\_matrices(), bfield(), bfield\_tangent(), bnorml(), brcast(), allglobal::broadcast\_inputs(), allglobal::check\_inputs(), curent(), dforce(), dfp100(), dfp200(), dvcfield(), evaluate\_dbb(), evaluate\_dmupfdx(), get\_lu\_beltrami\_matrices(), get\_perturbed\_solution(), sphdf5::hdfint(), intghs\_workspace\_init(), jo00aa(), lforce(), ma02aa(), matvec(), sphdf5::mirror\_input\_to\_outfile(), mp00ac(), packab(), pp00aa(), preset(), ra00aa(), spec(), tr00ab(), sphdf5::write\_grid(), and allglobal::wrtend().

**7.42.2.7 Iconstraint** integer inputlist::lconstraint = -1 selects constraints; primarily used in ma02aa() and mp00ac().

- if Lconstraint==-1, then in the plasma regions  $\Delta\psi_t$ ,  $\mu$  and  $\Delta\psi_p$  are not varied and in the vacuum region (only for free-boundary)  $\Delta\psi_t$  and  $\Delta\psi_p$  are not varied, and  $\mu=0$ .
- if Lconstraint==0, then in the plasma regions  $\Delta\psi_t$ ,  $\mu$  and  $\Delta\psi_p$  are not varied and in the vacuum region (only for free-boundary)  $\Delta\psi_t$  and  $\Delta\psi_p$  are varied to match the prescribed plasma current, current, and the "linking" current, curpol, and  $\mu=0$
- if Lconstraint==1, then in the plasma regions  $\mu$  and  $\Delta\psi_p$  are adjusted in order to satisfy the inner and outer interface transform constraints (except in the simple torus, where the enclosed poloidal flux is irrelevant, and only  $\mu$  is varied to satisfy the outer interface transform constraint); and in the vacuum region  $\Delta\psi_t$  and  $\Delta\psi_p$  are varied to match the transform constraint on the boundary and to obtain the prescribed linking current, curpol, and  $\mu=0$ .
- Todo if Lconstraint==2, under reconstruction.
- if Lconstraint.eq.3, then the  $\mu$  and  $\psi_p$  variables are adjusted in order to satisfy the volume and surface toroidal current computed with lbpol() (excepted in the inner most volume, where the volume current is irrelevant). Not implemented yet in free boundary.

Referenced by brcast(), allglobal::broadcast\_inputs(), allglobal::check\_inputs(), dfproe(), dfproe(), dfproe(), dfproe(), evaluate\_dbb(), evaluate\_dmupfdx(), get\_lu\_beltrami\_matrices(), get\_perturbed\_solution(), ma02aa(), sphdf5::mirror\_input\_to\_outfile mp00ac(), pp00aa(), preset(), spec(), and allglobal::wrtend().

**7.42.2.8 tflux** real, dimension(1:mnvol+1) inputlist::tflux = 0.0 toroidal flux,  $\psi_t$ , enclosed by each interface

- For each of the plasma volumes, this is a constraint: tflux is not varied
- For the vacuum region (only if Lfreebound==1), tflux may be allowed to vary to match constraints
- Note that tflux will be normalized so that tflux (Nvol) = 1.0, so that tflux is arbitrary up to a scale factor

See also

phiedge

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), dfp200(), sphdf5::hdfint(), sphdf5::mirror\_input\_to\_outfile(), preset(), spec(), and allglobal::wrtend().

```
7.42.2.9 helicity real, dimension(1:mnvol) inputlist::helicity = 0.0 helicity, \mathcal{K}, in each volume, \mathcal{V}_i
```

• on exit, helicity is set to the computed values of  $\mathcal{K} \equiv \int \mathbf{A} \cdot \mathbf{B} \ dv$ 

Referenced by brcast(), allglobal::broadcast\_inputs(), allglobal::check\_inputs(), df00ab(), sphdf5::hdfint(), hesian(), ma02aa(), sphdf5::mirror\_input\_to\_outfile(), mp00ac(), preset(), spec(), and allglobal::wrtend().

```
7.42.2.10 pscale real inputlist::pscale = 0.0 pressure scale factor
```

• the initial pressure profile is given by pscale \* pressure

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), dfp200(), evaluate\_dbb(), lforce(), sphdf5::mirror\_input\_to\_outfile(), spec(), volume(), and allglobal::wrtend().

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**7.42.2.11 pressure** real, dimension(1:mnvol+1) inputlist::pressure = 0.0 pressure in each volume

- The pressure is *not* held constant, but  $p_l V_l^{\gamma} = P_l$  is held constant, where  $P_l$  is determined by the initial pressures and the initial volumes,  $V_l$ .
- Note that if gamma==0.0, then  $p_l \equiv P_l$ .
- On output, the pressure is given by  $p_l = P_l/V_l^{\gamma}$ , where  $V_l$  is the final volume.
- pressure is only used in calculation of interface force-balance.

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), spec(), and allglobal::wrtend().

# **7.42.2.12 ladiabatic** integer inputlist::ladiabatic = 0 logical flag

- If Ladiabatic==0, the adiabatic constants are determined by the initial pressure and volume.
- If Ladiabatic==1, the adiabatic constants are determined by the given input adiabatic.

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), spec(), and allglobal::wrtend().

**7.42.2.13 adiabatic** real, dimension(1:mnvol+1) inputlist::adiabatic = 0.0 adiabatic constants in each volume

- The pressure is *not* held constant, but  $p_l V_l^{\gamma} = P_l \equiv \texttt{adiabatic}$  is constant.
- Note that if gamma==0.0, then pressure==adiabatic.
- pressure is only used in calculation of interface force-balance.

Referenced by allglobal::broadcast\_inputs(), dfp200(), evaluate\_dbb(), sphdf5::hdfint(), lforce(), sphdf5::mirror\_input\_to\_outfile(), spec(), and allglobal::wrtend().

```
7.42.2.14 pl integer, dimension(0:mnvol) inputlist::pl = 0 "inside" interface rotational-transform is \iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2. If both q_l=0 and q_r=0, then the (inside) interface rotational-transform is defined by iota . Referenced by allglobal::broadcast_inputs(), sphdf5::mirror_input_to_outfile(), preset(), and allglobal::wrtend().
```

```
7.42.2.15 ql integer, dimension(0:mnvol) inputlist::ql = 0 "inside" interface rotational-transform is t = (p_l + \gamma p_r)/(q_l + \gamma q_r), where \gamma is the golden mean, \gamma = (1 + \sqrt{5})/2. If both q_l = 0 and q_r = 0, then the (inside) interface rotational-transform is defined by iota . Referenced by allglobal::broadcast_inputs(), sphdf5::mirror_input_to_outfile(), preset(), and allglobal::wrtend().
```

```
7.42.2.16 pr integer, dimension(0:mnvol) inputlist::pr = 0 "inside" interface rotational-transform is \iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2. If both q_l=0 and q_r=0, then the (inside) interface rotational-transform is defined by iota . Referenced by allglobal::broadcast_inputs(), sphdf5::mirror_input_to_outfile(), preset(), and allglobal::wrtend().
```

7.42.2.17 qr integer, dimension(0:mnvol) inputlist::qr = 0

"inside" interface rotational-transform is  $t=(p_l+\gamma p_r)/(q_l+\gamma q_r)$ , where  $\gamma$  is the golden mean,  $\gamma=(1+\sqrt{5})/2$ . If both  $q_l=0$  and  $q_r=0$ , then the (inside) interface rotational-transform is defined by iota. Referenced by allglobal::broadcast inputs(), sphdf5::mirror input to outfile(), preset(), and allglobal::wrtend().

**7.42.2.18 iota** real, dimension(0:mnvol) inputlist::iota = 0.0 rotational-transform, t, on inner side of each interface

only relevant if illogical input for ql and qr are provided

Referenced by allglobal::broadcast\_inputs(), sphdf5::mirror\_input\_to\_outfile(), mp00ac(), pp00aa(), preset(), and allglobal::wrtend().

**7.42.2.19 Ip** integer, dimension(0:mnvol) inputlist::lp = 0

"outer" interface rotational-transform is  $t=(p_l+\gamma p_r)/(q_l+\gamma q_r)$ , where  $\gamma$  is the golden mean,  $\gamma=(1+\sqrt{5})/2$ . If both  $q_l=0$  and  $q_r=0$ , then the (outer) interface rotational-transform is defined by oita. Referenced by allglobal::broadcast\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

**7.42.2.20** Iq integer, dimension(0:mnvol) inputlist::lq = 0

"outer" interface rotational-transform is  $t = (p_l + \gamma p_r)/(q_l + \gamma q_r)$ , where  $\gamma$  is the golden mean,  $\gamma = (1 + \sqrt{5})/2$ . If both  $q_l = 0$  and  $q_r = 0$ , then the (outer) interface rotational-transform is defined by oita. Referenced by allglobal::broadcast\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

**7.42.2.21 rp** integer, dimension(0:mnvol) inputlist::rp = 0

"outer" interface rotational-transform is  $t=(p_l+\gamma p_r)/(q_l+\gamma q_r)$ , where  $\gamma$  is the golden mean,  $\gamma=(1+\sqrt{5})/2$ . If both  $q_l=0$  and  $q_r=0$ , then the (outer) interface rotational-transform is defined by oita. Referenced by allglobal::broadcast inputs(), sphdf5::mirror input to outfile(), preset(), and allglobal::wrtend().

**7.42.2.22 rq** integer, dimension(0:mnvol) inputlist::rq = 0

"outer" interface rotational-transform is  $t=(p_l+\gamma p_r)/(q_l+\gamma q_r)$ , where  $\gamma$  is the golden mean,  $\gamma=(1+\sqrt{5})/2$ . If both  $q_l=0$  and  $q_r=0$ , then the (outer) interface rotational-transform is defined by oita. Referenced by allglobal::broadcast inputs(), sphdf5::mirror input to outfile(), preset(), and allglobal::wrtend().

**7.42.2.23 oita** real, dimension(0:mnvol) inputlist::oita = 0.0 rotational-transform, t, on outer side of each interface

• only relevant if illogical input for ql and qr are provided

Referenced by allglobal::broadcast\_inputs(), sphdf5::mirror\_input\_to\_outfile(), mp00ac(), pp00aa(), preset(), and allglobal::wrtend().

7.42.2.24 mupftol real inputlist::mupftol = 1.0e-14 accuracy to which  $\mu$  and  $\Delta\psi_p$  are required

• only relevant if constraints on transform, enclosed currents etc. are to be satisfied iteratively, see Lconstraint

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), dforce(), evaluate\_dmupfdx(), ma02aa(), sphdf5::mirror\_input\_to\_outfile(), mp00ac(), and allglobal::wrtend().

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**7.42.2.25 mupfits** integer inputlist::mupfits = 8

an upper limit on the transform/helicity constraint iterations;

only relevant if constraints on transform, enclosed currents etc. are to be satisfied iteratively, see
 Lconstraint

• constraint: mupfits > 0

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), ma02aa(), sphdf5::mirror\_input\_to\_outfile(), and allglobal::wrtend().

**7.42.2.26 rpol** real inputlist::rpol = 1.0 poloidal extent of slab (effective radius)

- only relevant if Igeometry==1
- poloidal size is  $L=2\pi*{\tt rpol}$

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), coords(), sphdf5::mirror\_input\_to\_outfile(), sphdf5::write\_grid(), and allglobal::wrtend().

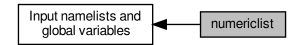
**7.42.2.27 rtor** real inputlist::rtor = 1.0 toroidal extent of slab (effective radius)

- only relevant if Igeometry==1
- toroidal size is  $L=2\pi*{\tt rtor}$

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), coords(), sphdf5::mirror\_input\_to\_outfile(), sphdf5::write\_grid(), and allglobal::wrtend().

## 7.43 numericlist

The namelist numericlist controls internal resolution parameters that the user rarely needs to consider. Collaboration diagram for numericlist:



#### **Variables**

• integer inputlist::linitialize = 0

Used to initialize geometry using a regularization / extrapolation method.

• integer inputlist::lautoinitbn = 1

Used to initialize  $B_{ns}$  using an initial fixed-boundary calculation.

integer inputlist::lzerovac = 0

Used to adjust vacuum field to cancel plasma field on computational boundary.

• integer inputlist::ndiscrete = 2

resolution of the real space grid on which fast Fourier transforms are performed is given by Ndiscrete\*Mpol\*4

• integer inputlist::nquad = -1

Resolution of the Gaussian quadrature.

• integer inputlist::impol = -4

Fourier resolution of straight-fieldline angle on interfaces.

integer inputlist::intor = -4

Fourier resolution of straight-fieldline angle on interfaces;.

• integer inputlist::lsparse = 0

controls method used to solve for rotational-transform on interfaces

• integer inputlist::lsvdiota = 0

controls method used to solve for rotational-transform on interfaces; only relevant if Lsparse = 0

• integer inputlist::imethod = 3

controls iterative solution to sparse matrix arising in real-space transformation to the straight-fieldline angle; only relevant if Lsparse.eq.2;

• integer inputlist::iorder = 2

controls real-space grid resolution for constructing the straight-fieldline angle; only relevant if Lsparse>0

• integer inputlist::iprecon = 0

controls iterative solution to sparse matrix arising in real-space transformation to the straight-fieldline angle; only relevant if Lsparse.eq.2;

• real inputlist::iotatol = -1.0

tolerance required for iterative construction of straight-fieldline angle; only relevant if Lsparse.ge.2

• integer inputlist::lextrap = 0

geometry of innermost interface is defined by extrapolation

integer inputlist::mregular = -1

maximum regularization factor

integer inputlist::lrzaxis = 1

controls the guess of geometry axis in the innermost volume or initialization of interfaces

• integer inputlist::ntoraxis = 3

the number of n harmonics used in the Jacobian m=1 harmonic elimination method; only relevant if  $Lrzaxis. \leftarrow qe.1$ .

## 7.43.1 Detailed Description

The namelist numericlist controls internal resolution parameters that the user rarely needs to consider.

#### 7.43.2 Variable Documentation

# **7.43.2.1 linitialize** integer inputlist::linitialize = 0

Used to initialize geometry using a regularization / extrapolation method.

- if  $\mbox{Linitialize} = -I$ , where I is a positive integer, the geometry of the  $i=1,N_V-I$  surfaces constructed by an extrapolation
- if Linitialize = 0, the geometry of the interior surfaces is provided after the namelists in the input file
- if Linitialize = 1, the interior surfaces will be intialized as  $R_{l,m,n} = R_{N,m,n} \psi_{t,l}^{m/2}$ , where  $R_{N,m,n}$  is the plasma boundary and  $\psi_{t,l}$  is the given toroidal flux enclosed by the l-th interface, normalized to the total enclosed toroidal flux; a similar extrapolation is used for  $Z_{l,m,n}$
- Note that the Fourier harmonics of the boundary is always given by the Rbc and Zbs given in physicslist.
- if Linitialize = 2, the interior surfaces and the plasma boundary will be intialized as  $R_{l,m,n}=R_{W,m,n}\psi_{t,l}^{m/2}$ , where  $R_{W,m,n}$  is the computational boundary and  $\psi_{t,l}$  is the given toroidal flux enclosed by the l-th interface, normalized to the total enclosed toroidal flux; a similar extrapolation is used for  $Z_{l,m,n}$

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• Note that, for free-boundary calculations, the Fourier harmonics of the computational boundary are *always* given by the Rwc and Zws given in physicslist.

• if Linitialize = 1, 2, it is not required to provide the geometry of the interfaces after the namelists

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), rzaxis(), and allglobal::wrtend().

### **7.43.2.2 lautoinitbn** integer inputlist::lautoinitbn = 1

Used to initialize  $B_{ns}$  using an initial fixed-boundary calculation.

- only relevant if Lfreebound = 1
- user-supplied Bns will only be considered if LautoinitBn = 0

Referenced by allglobal::broadcast inputs(), allglobal::check inputs(), spec(), and allglobal::wrtend().

#### **7.43.2.3 Izerovac** integer inputlist::lzerovac = 0

Used to adjust vacuum field to cancel plasma field on computational boundary.

• only relevant if Lfreebound = 1

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), spec(), and allglobal::wrtend().

#### **7.43.2.4 ndiscrete** integer inputlist::ndiscrete = 2

resolution of the real space grid on which fast Fourier transforms are performed is given by Ndiscrete\*Mpol\*4

• constraint Ndiscrete>0

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

#### **7.43.2.5 nquad** integer inputlist::nquad = -1

Resolution of the Gaussian quadrature.

- The resolution of the Gaussian quadrature,  $\int\!\!f(s)ds=\sum_k\omega_kf(s_k)$ , in each volume is given by  $\operatorname{Iquad}_v$ ,
- Iquad v is set in preset()

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

#### **7.43.2.6** impol integer inputlist::impol = -4

Fourier resolution of straight-fieldline angle on interfaces.

- the rotational-transform on the interfaces is determined by a transformation to the straight-fieldline angle, with poloidal resolution given by iMpol
- if iMpol <= 0, then iMpol = Mpol iMpol

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

#### **7.43.2.7 intor** integer inputlist::intor = -4

Fourier resolution of straight-fieldline angle on interfaces;.

- the rotational-transform on the interfaces is determined by a transformation to the straight-fieldline angle, with toroidal resolution given by iNtor
- if iNtor<=0 then iNtor = Ntor iNtor
- if Ntor==0, then the toroidal resolution of the angle transformation is set lNtor = 0

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

#### **7.43.2.8 Isparse** integer inputlist::lsparse = 0

controls method used to solve for rotational-transform on interfaces

- if Lsparse = 0, the transformation to the straight-fieldline angle is computed in Fourier space using a dense matrix solver, F04AAF
- if Lsparse = 1, the transformation to the straight-fieldline angle is computed in real space using a dense matrix solver, F04ATF
- if Lsparse = 2, the transformation to the straight-fieldline angle is computed in real space using a sparse matrix solver, F11DEF
- if Lsparse = 3, the different methods for constructing the straight-fieldline angle are compared

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), tr00ab(), and allglobal::wrtend().

# **7.43.2.9 Isvdiota** integer inputlist::lsvdiota = 0

controls method used to solve for rotational-transform on interfaces; only relevant if Lsparse = 0

- if Lsvdiota = 0, use standard linear solver to construct straight fieldline angle transformation
- if Lsvdiota = 1, use SVD method to compute rotational-transform

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), tr00ab(), and allglobal::wrtend().

## **7.43.2.10 imethod** integer inputlist::imethod = 3

controls iterative solution to sparse matrix arising in real-space transformation to the straight-fieldline angle; only relevant if Lsparse.eq.2;

See also

# tr00ab() for details

- if imethod = 1, the method is RGMRES
- if imethod = 2, the method is CGS
- if imethod = 3, the method is BICGSTAB

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), tr00ab(), and allglobal::wrtend().

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#### **7.43.2.11 iorder** integer inputlist::iorder = 2

controls real-space grid resolution for constructing the straight-fieldline angle; only relevant if Lsparse>0 determines order of finite-difference approximation to the derivatives

- if iorder = 2.
- if iorder = 4,
- if iorder = 6,

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), tr00ab(), and allglobal::wrtend().

#### **7.43.2.12 iprecon** integer inputlist::iprecon = 0

controls iterative solution to sparse matrix arising in real-space transformation to the straight-fieldline angle; only relevant if Lsparse.eq.2;

See also

tr00ab() for details

- if iprecon = 0, the preconditioner is 'N'
- if iprecon = 1, the preconditioner is 'J'
- if iprecon = 2, the preconditioner is 'S'

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), tr00ab(), and allglobal::wrtend().

# **7.43.2.13 mregular** integer inputlist::mregular = -1 maximum regularization factor

• if Mregular.ge.2, then regumm  $_i$  = Mregular /2 where m  $_i$  > Mregular

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

#### **7.43.2.14 Irzaxis** integer inputlist::lrzaxis = 1

controls the guess of geometry axis in the innermost volume or initialization of interfaces

- if iprecon = 1, the centroid is used
- if iprecon = 2, the Jacobian m = 1 harmonic elimination method is used

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), rzaxis(), and allglobal::wrtend().

# 7.44 locallist

The namelist locallist controls the construction of the Beltrami fields in each volume. Collaboration diagram for locallist:



#### **Variables**

• integer inputlist::lbeltrami = 4

Control flag for solution of Beltrami equation.

• integer inputlist::linitgues = 1

controls how initial guess for Beltrami field is constructed

• integer inputlist::lposdef = 0

redundant;

• real inputlist::maxrndgues = 1.0

the maximum random number of the Beltrami field if Linitgues = 3

• integer inputlist::Imatsolver = 3

1 for LU factorization, 2 for GMRES, 3 for GMRES matrix-free

• integer inputlist::nitergmres = 200

number of max iteration for GMRES

real inputlist::epsgmres = 1e-14

the precision of GMRES

• integer inputlist::lgmresprec = 1

type of preconditioner for GMRES, 1 for ILU sparse matrix

real inputlist::epsilu = 1e-12

the precision of incomplete LU factorization for preconditioning

#### 7.44.1 Detailed Description

The namelist locallist controls the construction of the Beltrami fields in each volume.

The transformation to straight-fieldline coordinates is singular when the rotational-transform of the interfaces is rational; however, the rotational-transform is still well defined.

## 7.44.2 Variable Documentation

#### 7.44.2.1 | Ibeltrami integer inputlist::lbeltrami = 4

Control flag for solution of Beltrami equation.

- if LBeltrami = 1,3,5 or 7, (SQP) then the Beltrami field in each volume is constructed by minimizing the magnetic energy with the constraint of fixed helicity; this is achieved by using sequential quadratic programming as provided by E04UFF. This approach has the benefit (in theory) of robustly constructing minimum energy solutions when multiple, i.e. bifurcated, solutions exist.
- if LBeltrami = 2,3,6 or 7, (Newton) then the Beltrami fields are constructed by employing a standard Newton method for locating an extremum of  $F \equiv \int B^2 dv \mu (\int \mathbf{A} \cdot \mathbf{B} dv \mathcal{K})$ , where  $\mu$  is treated as an independent degree of freedom similar to the parameters describing the vector potential and  $\mathcal{K}$  is the required value of the helicity; this is the standard Lagrange multipler approach for locating the constrained minimum; this method cannot distinguish saddle-type extrema from minima, and which solution that will be obtained depends on the initial guess;
- if LBeltrami = 4,5,6 or 7, (linear) it is assumed that the Beltrami fields are parameterized by  $\mu$ ; in this case, it is only required to solve  $\nabla \times \mathbf{B} = \mu \mathbf{B}$  which reduces to a system of linear equations;  $\mu$  may or may not be adjusted iteratively, depending on Lconstraint, to satisfy either rotational-transform or helicity constraints;
- for flexibility and comparison, each of the above methods can be employed; for example:
  - if LBeltrami = 1, only the SQP method will be employed;
  - if LBeltrami = 2, only the Newton method will be employed;
  - if LBeltrami = 4, only the linear method will be employed;
  - if LBeltrami = 3, the SQP and the Newton method are used;

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- if LBeltrami = 5, the SQP and the linear method are used;
- if LBeltrami = 6, the Newton and the linear method are used;
- if LBeltrami = 7, all three methods will be employed;

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

**7.44.2.2 linitgues** integer inputlist::linitgues = 1 controls how initial guess for Beltrami field is constructed

- only relevant for routines that require an initial guess for the Beltrami fields, such as the SQP and Newton methods, or the sparse linear solver;
- if Linitques = 0, the initial guess for the Beltrami field is trivial
- if Linitgues = 1, the initial guess for the Beltrami field is an integrable approximation
- if Linitgues = 2, the initial guess for the Beltrami field is read from file
- if Linitgues = 3, the initial guess for the Beltrami field will be randomized with the maximum maxrndgues

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

## 7.45 globallist

The namelist globallist controls the search for global force-balance. Collaboration diagram for globallist:



## **Variables**

• integer inputlist::lfindzero = 0

use Newton methods to find zero of force-balance, which is computed by dforce()

• real inputlist::escale = 0.0

controls the weight factor, BBweight, in the force-imbalance harmonics

• real inputlist::opsilon = 1.0

weighting of force-imbalance

real inputlist::pcondense = 2.0

spectral condensation parameter

• real inputlist::epsilon = 0.0

weighting of spectral-width constraint

• real inputlist::wpoloidal = 1.0

"star-like" poloidal angle constraint radial exponential factor used in preset() to construct sweight

• real inputlist::upsilon = 1.0

weighting of "star-like" poloidal angle constraint used in preset() to construct sweight

• real inputlist::forcetol = 1.0e-10

required tolerance in force-balance error; only used as an initial check

• real inputlist::c05xmax = 1.0e-06

required tolerance in position,  $\mathbf{x} \equiv \{R_{i,v}, Z_{i,v}\}$ 

• real inputlist::c05xtol = 1.0e-12

required tolerance in position,  $\mathbf{x} \equiv \{R_{i,v}, Z_{i,v}\}$ 

• real inputlist::c05factor = 1.0e-02

used to control initial step size in CO5NDF and CO5PDF

• logical inputlist::lreadgf = .true.

read  $\nabla_{\mathbf{x}}\mathbf{F}$  from file <code>ext.GF</code>

• integer inputlist::mfreeits = 0

maximum allowed free-boundary iterations

• real inputlist::bnstol = 1.0e-06

redundant;

• real inputlist::bnsblend = 0.666

redundant;

• real inputlist::gbntol = 1.0e-06

required tolerance in free-boundary iterations

• real inputlist::gbnbld = 0.666

normal blend

• real inputlist::vcasingeps = 1.e-12

regularization of Biot-Savart; see bnorml(), casing()

• real inputlist::vcasingtol = 1.e-08

accuracy on virtual casing integral; see bnorml(), casing()

• integer inputlist::vcasingits = 8

minimum number of calls to adaptive virtual casing routine; see casing()

integer inputlist::vcasingper = 1

periods of integragion in adaptive virtual casing routine; see <a href="casing">casing</a>()

• integer inputlist::mcasingcal = 8

minimum number of calls to adaptive virtual casing routine; see casing(); redundant;

#### 7.45.1 Detailed Description

The namelist  ${\tt globallist}$  controls the search for global force-balance. Comments:

• The "force" vector, **F**, which is constructed in dforce(), is a combination of pressure-imbalance Fourier harmonics.

$$F_{i,v} \equiv [[p+B^2/2]]_{i,v} \times \exp\left[-\operatorname{escale}(m_i^2 + n_i^2)\right] \times \operatorname{opsilon}, \tag{255}$$

and spectral-condensation constraints,  $I_{i,v}$ , and the "star-like" angle constraints,  $S_{i,v}$ , (see Iforce() for details)

$$F_{i,v} \equiv \operatorname{epsilon} \times I_{i,v} + \operatorname{upsilon} \times \left(\psi_v^{\omega} S_{i,v,1} - \psi_{v+1}^{\omega} S_{i,v+1,0}\right), \tag{256}$$

where  $\psi_v \equiv \text{normalized toroidal flux}$ , tflux, and  $\omega \equiv \text{wpoloidal}$ .

#### 7.45.2 Variable Documentation

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**7.45.2.1 Ifindzero** integer inputlist::lfindzero = 0

use Newton methods to find zero of force-balance, which is computed by dforce()

 if Lfindzero = 0, then dforce() is called once to compute the Beltrami fields consistent with the given geometry and constraints

- if Lfindzero = 1, then call CO5NDF (uses function values only), which iteratively calls dforce()
- if Lfindzero = 2, then call COSPDF (uses derivative information), which iteratively calls dforce()

Referenced by brcast(), allglobal::broadcast\_inputs(), allglobal::check\_inputs(), dfp200(), fcn1(), fcn2(), hesian(), sphdf5::mirror\_input\_to\_outfile(), newton(), packxi(), preset(), spec(), and allglobal::wrtend().

**7.45.2.2 escale** real inputlist::escale = 0.0

controls the weight factor, BBweight, in the force-imbalance harmonics

- BBweight (i)  $\equiv$  opsilon  $\times \exp\left[-\operatorname{escale} \times (m_i^2 + n_i^2)\right]$
- defined in preset(); used in dforce()
- also see Eqn. (255)

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

**7.45.2.3 opsilon** real inputlist::opsilon = 1.0 weighting of force-imbalance

• used in dforce(); also see Eqn. (255)

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

**7.45.2.4 pcondense** real inputlist::pcondense = 2.0 spectral condensation parameter

- used in preset() to define mmpp (i)  $\equiv m_i^p$ , where  $p \equiv p$  condense
- the angle freedom is exploited to minimize  $\operatorname{epsilon} \sum_i m_i^p (R_i^2 + Z_i^2)$  with respect to tangential variations in the interface geometry
- also see Eqn. (256)

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), preset(), and allglobal::wrtend().

**7.45.2.5 epsilon** real inputlist::epsilon = 0.0 weighting of spectral-width constraint

• used in dforce(); also see Eqn. (256)

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), dforce(), dfp200(), evaluate\_dbb(), sphdf5::mirror\_input\_to\_outfile(), pc00ab(), and allglobal::wrtend().

**7.45.2.6 forcetol** real inputlist::forcetol = 1.0e-10

required tolerance in force-balance error; only used as an initial check

- if the initially supplied interfaces are consistent with force-balance to within forcetol then the geometry of the interfaces is not altered
- if not, then the geometry of the interfaces is changed in order to bring the configuration into force balance so that the geometry of interfaces is within c05xtol, defined below, of the true solution
- to force execution of either  ${\tt C05NDF}$  or  ${\tt C05PDF}$ , regardless of the initial force imbalance, set  ${\tt forcetol} < {\tt 0}$

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), fcn1(), fcn2(), sphdf5::mirror\_input\_to\_outfile(), newton(), pc00aa(), pc00ab(), preset(), and allglobal::wrtend().

**7.45.2.7 c05xtol** real inputlist::c05xtol = 1.0e-12 required tolerance in position,  $\mathbf{x} \equiv \{R_{i,v}, Z_{i,v}\}$ 

- used by both C05NDF and C05PDF; see the NAG documents for further details on how the error is defined
- constraint c05xtol > 0.0

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), fcn1(), fcn2(), sphdf5::mirror\_input\_to\_outfile(), newton(), and allglobal::wrtend().

**7.45.2.8 c05factor** real inputlist::c05factor = 1.0e-02 used to control initial step size in C05NDF and C05PDF

- constraint c05factor > 0.0
- only relevant if Lfindzero > 0

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), fcn1(), fcn2(), sphdf5::mirror\_input\_to\_outfile(), newton(), and allglobal::wrtend().

- only used if Lfindzero = 2
- only used in newton()

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), fcn1(), fcn2(), sphdf5::mirror\_input\_to\_outfile(), newton(), and allglobal::wrtend().

**7.45.2.10 mfreeits** integer inputlist::mfreeits = 0 maximum allowed free-boundary iterations

- only used if Lfreebound = 1
- only used in xspech()

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), spec(), and allglobal::wrtend().

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**7.45.2.11 gbntol** real inputlist::gbntol = 1.0e-06 required tolerance in free-boundary iterations

- only used if Lfreebound = 1
- only used in xspech()

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), spec(), and allglobal::wrtend().

**7.45.2.12 gbnbld** real inputlist::gbnbld = 0.666 normal blend

• The "new" magnetic field at the computational boundary produced by the plasma currents is updated using a Picard scheme:

$$(\mathbf{B} \cdot \mathbf{n})^{j+1} = gBnbld \times (\mathbf{B} \cdot \mathbf{n})^j + (1 - gBnbld) \times (\mathbf{B} \cdot \mathbf{n})^*,$$
(257)

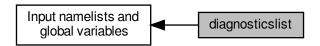
where j labels free-boundary iterations, and  $(\mathbf{B} \cdot \mathbf{n})^*$  is computed by virtual casing.

- only used if Lfreebound = 1
- only used in xspech()

Referenced by allglobal::broadcast\_inputs(), allglobal::check\_inputs(), sphdf5::mirror\_input\_to\_outfile(), spec(), and allglobal::wrtend().

# 7.46 diagnosticslist

The namelist diagnosticslist controls post-processor diagnostics, such as Poincaré plot resolution, etc. Collaboration diagram for diagnosticslist:



### **Variables**

• real inputlist::odetol = 1.0e-07

o.d.e. integration tolerance for all field line tracing routines

• real inputlist::absreq = 1.0e-08

redundant

• real inputlist::relreq = 1.0e-08

redundant

• real inputlist::absacc = 1.0e-04

redundant

• real inputlist::epsr = 1.0e-08

redundant

• integer inputlist::nppts = 0

number of toroidal transits used (per trajectory) in following field lines for constructing Poincaré plots; if nPpts < 1, no Poincaré plot is constructed;

• real inputlist::ppts = 0.0

stands for Poincare plot theta start. Chose at which angle (normalized over  $\pi$ ) the Poincare field-line tracing start.

• integer, dimension(1:mnvol+1) inputlist::nptrj = -1

number of trajectories in each annulus to be followed in constructing Poincaré plot

• logical inputlist::lhevalues = .false.

to compute eigenvalues of  $abla \mathbf{F}$ 

• logical inputlist::lhevectors = .false.

to compute eigenvectors (and also eigenvalues) of  $\nabla \mathbf{F}$ 

• logical inputlist::Ihmatrix = .false.

to compute and write to file the elements of  $\nabla \mathbf{F}$ 

• integer inputlist::lperturbed = 0

to compute linear, perturbed equilibrium

• integer inputlist::dpp = -1

perturbed harmonic

• integer inputlist::dqq = -1

perturbed harmonic

• integer inputlist::lerrortype = 0

the type of error output for Lcheck=1

integer inputlist::ngrid = -1

the number of points to output in the grid, -1 for Lrad(vvol)

• real inputlist::drz = 1E-5

difference in geometry for finite difference estimate (debug only)

• integer inputlist::lcheck = 0

implement various checks

• logical inputlist::Itiming = .false.

to check timing

• real inputlist::fudge = 1.0e-00

redundant

• real inputlist::scaling = 1.0e-00

redundant

#### 7.46.1 Detailed Description

The namelist diagnosticslist controls post-processor diagnostics, such as Poincaré plot resolution, etc.

#### 7.46.2 Variable Documentation

**7.46.2.1 nptrj** integer, dimension(1:mnvol+1) inputlist::nptrj = -1 number of trajectories in each annulus to be followed in constructing Poincaré plot

if nPtrj(1) <0, then nPtrj(1) = Ni(l), where Ni(1) is the grid resolution used to construct the Beltrami field in volume l</li>

Referenced by allglobal::broadcast\_inputs(), final\_diagnostics(), sphdf5::mirror\_input\_to\_outfile(), pp00aa(), spec(), and allglobal::wrtend().

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**7.46.2.2 Icheck** integer inputlist::lcheck = 0 implement various checks

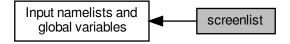
- if Lcheck = 0, no additional check on the calculation is performed
- if Lcheck = 1, the error in the current, i.e.  $\nabla \times \mathbf{B} \mu \mathbf{B}$  is computed as a post-diagnostic
- if Lcheck = 2, the analytic derivatives of the interface transform w.r.t. the helicity multiplier,  $\mu$ , and the enclosed poloidal flux,  $\Delta \psi_p$ , are compared to a finite-difference estimate
  - only if Lconstraint==1
  - only for dspec executable, i.e. must compile with DFLAGS = "-D DEBUG"
- if Lcheck = 3, the analytic derivatives of the volume w.r.t. interface Fourier harmonic is compared to a finite-difference estimate
  - must set Lfindzero = 2
  - set forcetol sufficiently small and set LreadGF = F, so that the matrix of second derivatives is calculated
  - only for dspec executable, i.e. must compile with DFLAGS = "-D DEBUG"
- if Lcheck = 4, the analytic calculation of the derivatives of the magnetic field,  $B^2$ , at the interfaces is compared to a finite-difference estimate
  - must set Lfindzero = 2
  - set forcetol sufficiently small
  - set LreadGF=F
  - only for dspec executable, i.e. must compile with DFLAGS = "-D DEBUG"
- if Lcheck = 5, the analytic calculation of the matrix of the derivatives of the force imbalance is compared to a finite-difference estimate
- if Lcheck = 6, the virtual casing calculation is compared to xdiagno (Lazerson 2013 [6])
  - the input file for xdiagno is written by bnorml()
  - this provides the Cartesian coordinates on the computational boundary where the virtual casing routine casing() computes the magnetic field, with the values of the magnetic field being written to the screen for comparison
  - must set Freebound=1, Lfindzero>0, mfreeits!=0
  - xdiagno must be executed manually

Referenced by bnorml(), allglobal::broadcast\_inputs(), allglobal::check\_inputs(), dforce(), dfp200(), evaluate\_dbb(), evaluate\_dmupfdx(), fcn1(), fcn2(), final\_diagnostics(), sphdf5::hdfint(), hesian(), lforce(), ma02aa(), sphdf5::mirror\_input\_to\_outfile(), newton(), preset(), rzaxis(), spec(), and allglobal::wrtend().

## 7.47 screenlist

The namelist screenlist controls screen output. Every subroutine, e.g. xy00aa.h, has its own write flag, Wxy00aa.

Collaboration diagram for screenlist:



#### **Variables**

- logical inputlist::wmanual = .false.
- logical inputlist::wrzaxis = .false.
- logical inputlist::wpackxi = .false.
- logical inputlist::wvolume = .false.
- logical inputlist::wcoords = .false.
- logical inputlist::wbasefn = .false.
- logical inputlist::wmemory = .false.
- logical inputlist::wmetrix = .false.
- logical inputlist::wma00aa = .false.
- logical inputlist::wmatrix = .false.
- logical inputlist::wspsmat = .false.
- logical inputlist::wspsint = .false.
- logical inputlist::wmp00ac = .false.
- logical inputlist::wma02aa = .false.
- logical inputlist::wpackab = .false.
- logical inputlist::wtr00ab = .false.
- logical inputlist::wcurent = .false.
- logical inputlist::wdf00ab = .false.
- logical inputlist::wlforce = .false.
- logical inputlist::wintghs = .false.
- logical inputlist::wmtrxhs = .false.
- logical inputlist::wlbpol = .false.
- logical inputlist::wbrcast = .false.
- logical inputlist::wdfp100 = .false.
- logical inputlist::wdfp200 = .false.
- logical inputlist::wdforce = .false.
- logical inputlist::wnewton = .false.
- logical inputlist::wcasing = .false.
- logical inputlist::wbnorml = .false.
- logical inputlist::wjo00aa = .false.
- logical inputlist::wpp00aa = .false.
- logical inputlist::wpp00ab = .false.
- logical inputlist::wbfield = .false.
- logical inputlist::wstzxyz = .false.
- logical inputlist::whesian = .false.
- logical inputlist::wra00aa = .false.
- logical inputlist::wnumrec = .false.
  logical inputlist::wdcuhre = .false.
- logical inputist::wdcurre = :laise.
   logical inputist::wminpack = .false.
- logical inputlist::wiqpack = .false.
- logical inputlist::wrksuite = .false.
- logical inputlist::wi1mach = .false.
- logical inputlist::wd1mach = .false.
- logical inputlist::wilut = .false.
- logical inputlist::witers = .false.
- logical inputlist::wsphdf5 = .false.
- logical inputlist::wpreset = .false.
- logical inputlist::wglobal = .false.
- logical inputlist::wxspech = .false.
- logical inputlist::wbuild\_vector\_potential = .false.
- logical inputlist::wreadin = .false.

write screen output of readin()

• logical inputlist::wwrtend = .false.

write screen output of wrtend()

logical inputlist::wmacros = .false.

write screen output from expanded macros

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## 7.47.1 Detailed Description

The namelist screenlist controls screen output. Every subroutine, e.g. xy00aa.h, has its own write flag, Wxy00aa.h

#### 7.47.2 Variable Documentation

7.47.2.1 wbuild\_vector\_potential = .false.

Todo: what is this?

## 8 Module Documentation

# 8.1 aligiobal Module Reference

global variable storage used as "workspace" throughout the code

#### **Functions/Subroutines**

- subroutine build\_vector\_potential (Ivol, iocons, aderiv, tderiv)
- subroutine set\_mpi\_comm (comm)
- subroutine read\_inputlists\_from\_file ()
- subroutine check\_inputs ()
- · subroutine broadcast inputs
- · subroutine wrtend

The restart file is written.

subroutine ismyvolume (vvol)

Check if volume vvol is associated to the corresponding MPI node.

subroutine whichcpuid (vvol, cpu\_id)

Returns which MPI node is associated to a given volume.

#### Variables

· integer myid

MPI rank of current CPU.

• integer ncpu

number of MPI tasks

• integer ismyvolumevalue

flag to indicate if a CPU is operating on its assigned volume

real cpus

initial time

• integer mpi\_comm\_spec

SPEC MPI communicator.

- logical **skip\_write** = .false.
- real pi2nfp
- · real pi2pi2nfp
- · real pi2pi2nfphalf
- · real pi2pi2nfpquart
- character(len=1000) ext
- · real forceerr

total force-imbalance

real energy

MHD energy.

- · real, dimension(:), allocatable ipdt
- · real, dimension(:,:), allocatable ipdtdpf

Toroidal pressure-driven current.

- · integer mvol
- logical yesstellsym

internal shorthand copies of Istellsym, which is an integer input;

logical notstellsym

internal shorthand copies of Istellsym, which is an integer input;

- · logical yesmatrixfree
- logical notmatrixfree

to use matrix-free method or not

real, dimension(:,:), allocatable cheby

local workspace for evaluation of Chebychev polynomials

real, dimension(:,:,:), allocatable zernike

local workspace for evaluation of Zernike polynomials

• real, dimension(:,:,:), allocatable tt

derivatives of Chebyshev polynomials at the inner and outer interfaces;

real, dimension(:,:,:,:), allocatable rtt

derivatives of Zernike polynomials at the inner and outer interfaces;

· real, dimension(:,:), allocatable rtm

 $\boldsymbol{r}^{m}$  term of Zernike polynomials at the origin

· real, dimension(:), allocatable zernikedof

Zernike degree of freedom for each m.

· integer mne

enhanced resolution for metric elements

• integer, dimension(:), allocatable ime

enhanced poloidal mode numbers for metric elements

· integer, dimension(:), allocatable ine

enhanced toroidal mode numbers for metric elements

integer mns

enhanced resolution for straight field line transformation

• integer, dimension(:), allocatable ims

enhanced poloidal mode numbers for straight field line transformation

• integer, dimension(:), allocatable ins

enhanced toroidal mode numbers for straight field line transformation

integer Impol

what is this?

· integer Intor

what is this?

integer smpol

what is this?

· integer sntor

what is this?

• real xoffset = 1.0

used to normalize NAG routines (which ones exacly where?)

logical, dimension(:), allocatable imagneticok

used to indicate if Beltrami fields have been correctly constructed;

logical iconstraintok

Used to break iteration loops of slaves in the global constraint minimization.

• real, dimension(:,:), allocatable beltramierror

to store the integral of |curlB-mu\*B| computed by jo00aa;

integer mn

total number of Fourier harmonics for coordinates/fields; calculated from Mpol, Ntor in readin()

· integer, dimension(:), allocatable im

poloidal mode numbers for Fourier representation

• integer, dimension(:), allocatable in

toroidal mode numbers for Fourier representation

· real, dimension(:), allocatable halfmm

I saw this already somewhere...

• real, dimension(:), allocatable regumm

I saw this already somewhere...

· real rscale

no idea

real, dimension(:,:), allocatable psifactor

no idea

real, dimension(:,:), allocatable inifactor

no idea

• real, dimension(:), allocatable bbweight

weight on force-imbalance harmonics; used in dforce()

• real, dimension(:), allocatable mmpp

spectral condensation factors

• real, dimension(:,:), allocatable irbc

cosine R harmonics of interface surface geometry; stellarator symmetric

real, dimension(:,:), allocatable izbs

sine Z harmonics of interface surface geometry; stellarator symmetric

real, dimension(:,:), allocatable irbs

sine R harmonics of interface surface geometry; non-stellarator symmetric

real, dimension(:,:), allocatable izbc

cosine Z harmonics of interface surface geometry; non-stellarator symmetric

• real, dimension(:,:), allocatable drbc

cosine R harmonics of interface surface geometry; stellarator symmetric; linear deformation

real, dimension(:,:), allocatable dzbs

sine Z harmonics of interface surface geometry; stellarator symmetric; linear deformation

• real, dimension(:,:), allocatable drbs

sine R harmonics of interface surface geometry; non-stellarator symmetric; linear deformation

• real, dimension(:,:), allocatable dzbc

cosine Z harmonics of interface surface geometry; non-stellarator symmetric; linear deformation

• real, dimension(:,:), allocatable irij

interface surface geometry; real space

• real, dimension(:,:), allocatable izij

interface surface geometry; real space

• real, dimension(:,:), allocatable drij

interface surface geometry; real space

real, dimension(:,:), allocatable dzij

interface surface geometry; real space

• real, dimension(:,:), allocatable trij

interface surface geometry; real space

real, dimension(:,:), allocatable tzij

interface surface geometry; real space

· real, dimension(:), allocatable ivns

sine harmonics of vacuum normal magnetic field on interfaces; stellarator symmetric

• real, dimension(:), allocatable ibns

sine harmonics of plasma normal magnetic field on interfaces; stellarator symmetric

· real, dimension(:), allocatable ivnc

cosine harmonics of vacuum normal magnetic field on interfaces; non-stellarator symmetric

· real, dimension(:), allocatable ibnc

cosine harmonics of plasma normal magnetic field on interfaces; non-stellarator symmetric

• real, dimension(:), allocatable Irbc

local workspace

• real, dimension(:), allocatable lzbs

local workspace

· real, dimension(:), allocatable Irbs

local workspace

• real, dimension(:), allocatable lzbc

local workspace

- integer num\_modes
- integer, dimension(:), allocatable mmrzrz
- integer, dimension(:), allocatable nnrzrz
- real, dimension(:,:,:), allocatable allrzrz
- integer nt

discrete resolution along  $\theta$  of grid in real space

• integer nz

discrete resolution along  $\zeta$  of grid in real space

· integer ntz

discrete resolution; Ntz=Nt\*Nz shorthand

· integer hnt

discrete resolution; Ntz=Nt\*Nz shorthand

· integer hnz

discrete resolution; Ntz=Nt\*Nz shorthand

real sontz

one / sqrt (one\*Ntz); shorthand

• real, dimension(:,:,:), allocatable rij

real-space grid; R

• real, dimension(:,:,:), allocatable zij

real-space grid; Z

• real, dimension(:,:,:), allocatable xij

what is this?

• real, dimension(:,:,:), allocatable yij

what is this?

• real, dimension(:,:), allocatable sg

real-space grid; jacobian and its derivatives

• real, dimension(:,:,:,:), allocatable guvij

real-space grid; metric elements

• real, dimension(:,:,:), allocatable gvuij

real-space grid; metric elements (?); 10 Dec 15;

real, dimension(:,:,:,:), allocatable guvijsave

what is this?

• integer, dimension(:,:), allocatable ki

identification of Fourier modes

integer, dimension(:,:,:), allocatable kijs

identification of Fourier modes

• integer, dimension(:,:,:), allocatable kija

identification of Fourier modes

 integer, dimension(:), allocatable iotakkii identification of Fourier modes

integer, dimension(:,:), allocatable iotaksub

identification of Fourier modes

integer, dimension(:,:), allocatable iotakadd

identification of Fourier modes

• integer, dimension(:,:), allocatable iotaksgn

identification of Fourier modes

real, dimension(:), allocatable efmn

Fourier harmonics; dummy workspace.

· real, dimension(:), allocatable ofmn

Fourier harmonics; dummy workspace.

real, dimension(:), allocatable cfmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable sfmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable evmn

Fourier harmonics; dummy workspace.

· real, dimension(:), allocatable odmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable comn

Fourier harmonics; dummy workspace.

real, dimension(:), allocatable simn

Fourier harmonics; dummy workspace.

· real, dimension(:), allocatable ijreal

what is this?

· real, dimension(:), allocatable ijimag

what is this?

• real, dimension(:), allocatable **jireal** 

what is this?

• real, dimension(:), allocatable jiimag

what is this?

• real, dimension(:), allocatable jkreal

what is this?

• real, dimension(:), allocatable jkimag

what is this?

• real, dimension(:), allocatable kjreal

what is this?

real, dimension(:), allocatable kjimag

what is this?

• real, dimension(:,:,:), allocatable **bsupumn** 

tangential field on interfaces;  $\theta$ -component; required for virtual casing construction of field; 11 Oct 12

real, dimension(:,:,:), allocatable bsupvmn

tangential field on interfaces;  $\zeta$  -component; required for virtual casing construction of field; 11 Oct 12

• real, dimension(:,:), allocatable **goomne** 

described in preset()

real, dimension(:,:), allocatable goomno

described in preset()

• real, dimension(:,:), allocatable gssmne

described in preset()

real, dimension(:,:), allocatable gssmno

described in preset()

```
• real, dimension(:,:), allocatable gstmne
      described in preset()

    real, dimension(:,:), allocatable gstmno

      described in preset()
• real, dimension(:,:), allocatable gszmne
      described in preset()

    real, dimension(:,:), allocatable gszmno

      described in preset()
• real, dimension(:,:), allocatable gttmne
      described in preset()
• real, dimension(:,:), allocatable gttmno
      described in preset()
• real, dimension(:,:), allocatable gtzmne
      described in preset()

    real, dimension(:,:), allocatable gtzmno

      described in preset()
• real, dimension(:,:), allocatable gzzmne
      described in preset()

    real, dimension(:,:), allocatable gzzmno

      described in preset()

    real, dimension(:,:,:,:), allocatable dtoocc

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable dtoocs

      volume-integrated Chebychev-metrics; see matrix()
• real, dimension(:,:,:), allocatable dtoosc
      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable dtooss

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ttsscc

      volume-integrated Chebychev-metrics; see matrix()
• real, dimension(:,:,:,:), allocatable ttsscs
      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ttsssc

      volume-integrated Chebychev-metrics; see matrix()
• real, dimension(:,:,:,:), allocatable ttssss
      volume-integrated Chebychev-metrics; see matrix()
• real, dimension(:,:,:,:), allocatable tdstcc
      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable tdstcs

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable tdstsc

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable tdstss

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable tdszcc

      volume-integrated Chebychev-metrics; see matrix()
• real, dimension(:,:,:,:), allocatable tdszcs
      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable tdszsc

      volume-integrated Chebychev-metrics; see matrix()
```

```
    real, dimension(:,:,:,:), allocatable tdszss

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ddttcc

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ddttcs

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ddttsc

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ddttss

      volume-integrated Chebychev-metrics; see matrix()
• real, dimension(:,:,:,:), allocatable ddtzcc
      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ddtzcs

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ddtzsc

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ddtzss

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ddzzcc

      volume-integrated Chebychev-metrics; see matrix()
• real, dimension(:,:,:,:), allocatable ddzzcs
      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ddzzsc

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:,:,:), allocatable ddzzss

      volume-integrated Chebychev-metrics; see matrix()

    real, dimension(:,:), allocatable tsc

      what is this?
• real, dimension(:,:), allocatable tss
      what is this?
• real, dimension(:,:), allocatable dtc
      what is this?
· real, dimension(:,:), allocatable dts
      what is this?
• real, dimension(:,:), allocatable dzc
      what is this?

    real, dimension(:,:), allocatable dzs

      what is this?

    real, dimension(:,:), allocatable ttc

      what is this?

    real, dimension(:,:), allocatable tzc

      what is this?
• real, dimension(:,:), allocatable tts
      what is this?
• real, dimension(:,:), allocatable tzs
      what is this?

    real, dimension(:), allocatable dtflux

      \delta \psi_{toroidal} in each annulus
· real, dimension(:), allocatable dpflux
      \delta\psi_{poloidal} in each annulus

    real, dimension(:), allocatable sweight
```

minimum poloidal length constraint weightinteger, dimension(:), allocatable nadof

· integer, dimension(:), allocatable nfielddof

type(subgrid), dimension(:,:,:), allocatable ate

type(subgrid), dimension(:,:,:), allocatable aze

• type(subgrid), dimension(:,:,:), allocatable ato

degrees of freedom in Beltrami fields in each annulus

degrees of freedom in Beltrami fields in each annulus, field only, no Lagrange multipliers

magnetic vector potential cosine Fourier harmonics; stellarator-symmetric

magnetic vector potential cosine Fourier harmonics; stellarator-symmetric

magnetic vector potential sine Fourier harmonics; non-stellarator-symmetric

```
    type(subgrid), dimension(:,:,:), allocatable azo

      magnetic vector potential sine Fourier harmonics; non-stellarator-symmetric
• integer, dimension(:,:), allocatable Ima
     Lagrange multipliers (?)
• integer, dimension(:,:), allocatable Imb
     Lagrange multipliers (?)
• integer, dimension(:,:), allocatable Imc
     Lagrange multipliers (?)

    integer, dimension(:,:), allocatable Imd

     Lagrange multipliers (?)
• integer, dimension(:,:), allocatable Ime
     Lagrange multipliers (?)

    integer, dimension(:,:), allocatable Imf

      Lagrange multipliers (?)
• integer, dimension(:,:), allocatable Img
      Lagrange multipliers (?)
• integer, dimension(:,:), allocatable Imh
     Lagrange multipliers (?)
• real, dimension(:,:), allocatable Imavalue
      what is this?
• real, dimension(:,:), allocatable Imbvalue
      what is this?
• real, dimension(:,:), allocatable Imcvalue
      what is this?
• real, dimension(:,:), allocatable Imdvalue
      what is this?
• real, dimension(:,:), allocatable Imevalue
      what is this?
• real, dimension(:,:), allocatable Imfvalue
      what is this?

    real, dimension(:,:), allocatable Imgvalue

      what is this?
• real, dimension(:,:), allocatable Imhvalue
      what is this?

    integer, dimension(:,:), allocatable fso

      what is this?
• integer, dimension(:,:), allocatable fse
      what is this?

    logical lcoordinatesingularity

      set by LREGION macro; true if inside the innermost volume
```

• logical Iplasmaregion

set by LREGION macro; true if inside the plasma region

logical Ivacuumregion

set by LREGION macro; true if inside the vacuum region

· logical Isavedguvij

flag used in matrix free

· logical localconstraint

what is this?

• real, dimension(:,:), allocatable dma

energy and helicity matrices; quadratic forms

• real, dimension(:,:), allocatable dmb

energy and helicity matrices; quadratic forms

real, dimension(:,:), allocatable dmd

energy and helicity matrices; quadratic forms

· real, dimension(:), allocatable dmas

sparse version of dMA, data

• real, dimension(:), allocatable dmds

sparse version of dMD, data

• integer, dimension(:), allocatable idmas

sparse version of dMA and dMD, indices

• integer, dimension(:), allocatable jdmas

sparse version of dMA and dMD, indices

integer, dimension(:), allocatable ndmasmax

number of elements for sparse matrices

· integer, dimension(:), allocatable ndmas

number of elements for sparse matrices

• real, dimension(:), allocatable dmg

what is this?

• real, dimension(:), allocatable adotx

the matrix-vector product

• real, dimension(:), allocatable ddotx

the matrix-vector product

• real, dimension(:,:), allocatable solution

this is allocated in dforce; used in mp00ac and ma02aa; and is passed to packab

• real, dimension(:,;,:), allocatable gmreslastsolution

used to store the last solution for restarting GMRES

• real, dimension(:), allocatable mbpsi

matrix vector products

· logical liluprecond

whether to use ILU preconditioner for GMRES

• real, dimension(:,:), allocatable beltramiinverse

Beltrami inverse matrix.

real, dimension(:,:,:), allocatable diotadxup

measured rotational transform on inner/outer interfaces for each volume; d(transform)/dx; (see dforce)

• real, dimension(:,:,:), allocatable ditgpdxtp

measured toroidal and poloidal current on inner/outer interfaces for each volume; d(ltor, Gpol)/dx; (see dforce)

real, dimension(:,:,:,:), allocatable glambda

save initial guesses for iterative calculation of rotational-transform

· integer Imns

what is this?

real, dimension(:,:,:), allocatable bemn

```
force vector; stellarator-symmetric (?)
• real, dimension(:,:), allocatable iomn
      force vector; stellarator-symmetric (?)

    real, dimension(:,:,:), allocatable somn

      force vector; non-stellarator-symmetric (?)

    real, dimension(:,:,:), allocatable pomn

      force vector; non-stellarator-symmetric (?)

    real, dimension(:,:,:), allocatable bomn

      force vector; stellarator-symmetric (?)

    real, dimension(:,:), allocatable iemn

      force vector; stellarator-symmetric (?)
• real, dimension(:,:,:), allocatable semn
      force vector; non-stellarator-symmetric (?)
• real, dimension(:,:,:), allocatable pemn
      force vector; non-stellarator-symmetric (?)
· real, dimension(:), allocatable bbe
      force vector (?); stellarator-symmetric (?)
· real, dimension(:), allocatable iio
      force vector (?); stellarator-symmetric (?)
· real, dimension(:), allocatable bbo
      force vector (?); non-stellarator-symmetric (?)
· real, dimension(:), allocatable iie
      force vector (?); non-stellarator-symmetric (?)

    real, dimension(:,:,:), allocatable btemn

      covariant \theta cosine component of the tangential field on interfaces; stellarator-symmetric
• real, dimension(:,:,:), allocatable bzemn
      covariant \zeta cosine component of the tangential field on interfaces; stellarator-symmetric
• real, dimension(:,:,:), allocatable btomn
      covariant \theta sine component of the tangential field on interfaces; non-stellarator-symmetric

    real, dimension(:,:,:), allocatable bzomn

      covariant \zeta sine component of the tangential field on interfaces; non-stellarator-symmetric

    real, dimension(:,:), allocatable bloweremn

      covariant field for Hessian computation

    real, dimension(:,:), allocatable bloweromn

      covariant field for Hessian computation

    integer Igdof

      geometrical degrees of freedom associated with each interface
· integer ngdof
      total geometrical degrees of freedom

    real, dimension(:,:,:), allocatable dbbdrz

      derivative of magnetic field w.r.t. geometry (?)

    real, dimension(:,:), allocatable diidrz

      derivative of spectral constraints w.r.t. geometry (?)

    real, dimension(:,:,:,:), allocatable dffdrz

      derivatives of B<sup>\(\)</sup>2 at the interfaces wrt geometry

    real, dimension(:,:,:,:), allocatable dbbdmp

      derivatives of B^{\wedge}2 at the interfaces wrt mu and dpflux

    real, dimension(:,:,:,:), allocatable dmupfdx

      derivatives of mu and dpflux wrt geometry at constant interface transform

    logical Ihessianallocated
```

flag to indicate that force gradient matrix is allocated (?)

- real, dimension(:,:), allocatable hessian force gradient matrix (?)
- real, dimension(:,:), allocatable dessian derivative of force gradient matrix (?)
- real, dimension(:,:), allocatable cosi some precomputed cosines
- real, dimension(:,:), allocatable sini some precomputed sines
- real, dimension(:), allocatable **gteta** something related to  $\sqrt{g}$  and  $\theta$ ?
- real, dimension(:), allocatable gzeta
   something related to √g and ζ?
   real, dimension(:), allocatable gits
- real, dimension(:), allocatable ajk definition of coordinate axis
- real, dimension(:,:,:), allocatable dradr derivatives of coordinate axis
- real, dimension(:,:,:), allocatable dradz
   derivatives of coordinate axis
- real, dimension(:,:,:,:), allocatable dzadr derivatives of coordinate axis
- real, dimension(:,:,:,:), allocatable dzadz
   derivatives of coordinate axis
- real, dimension(:,:,:), allocatable drodr
   derivatives of coordinate axis
- real, dimension(:,:,:), allocatable drodz
   derivatives of coordinate axis
- real, dimension(:,:,:), allocatable dzodr
   derivatives of coordinate axis
- real, dimension(:,:,:), allocatable dzodz
   derivatives of coordinate axis
- integer, dimension(:,:), allocatable djkp for calculating cylindrical volume
- integer, dimension(:,:), allocatable djkm for calculating cylindrical volume
- real, dimension(:), allocatable **lbbintegral** *B.B integral*.
- real, dimension(:), allocatable labintegral
- A.B integral.
   real, dimension(:), allocatable **vvolume**
- volume integral of  $\sqrt{g}$ ; computed in volume real **dvolume**
- · rear uvoiume

derivative of volume w.r.t. interface geometry

integer ivol

labels volume; some subroutines (called by NAG) are fixed argument list but require the volume label

real gbzeta

toroidal (contravariant) field; calculated in bfield; required to convert  $\dot{\theta}$  to  $B^{\theta}$ ,  $\dot{s}$  to  $B^{s}$ 

- integer, dimension(:), allocatable iquad
  - internal copy of Nquad
- real, dimension(:,:), allocatable gaussianweight

weights for Gaussian quadrature

real, dimension(:,:), allocatable gaussianabscissae

abscissae for Gaussian quadrature

· logical Iblinear

controls selection of Beltrami field solver; depends on LBeltrami

logical Ibnewton

controls selection of Beltrami field solver; depends on LBeltrami

logical Ibsequad

controls selection of Beltrami field solver; depends on LBeltrami

real, dimension(1:3) orzp

used in mg00aa() to determine  $(s, \theta, \zeta)$  given  $(R, Z, \varphi)$ 

• type(derivative) dbdx

 $d\mathbf{B}/d\mathbf{X}$  (?)

integer globaljk

labels position

• real, dimension(:,:), allocatable dxyz

computational boundary; position

• real, dimension(:,:), allocatable nxyz

computational boundary; normal

• real, dimension(:,:), allocatable jxyz

plasma boundary; surface current

real, dimension(1:2) tetazeta

what is this?

• real virtualcasingfactor = -one / (four\*pi)

this agrees with diagno

· integer iberror

for computing error in magnetic field

• integer nfreeboundaryiterations

number of free-boundary iterations already performed

• integer, parameter **node** = 2

best to make this global for consistency between calling and called routines

• logical first\_free\_bound = .false.

flag to indicate that this is the first free-boundary iteration

# 8.1.1 Detailed Description

global variable storage used as "workspace" throughout the code

### 8.1.2 Function/Subroutine Documentation

# **8.1.2.1 check\_inputs()** subroutine allglobal::check\_inputs reading of physicslist

- The internal variable, Mvol=Nvol+Lfreebound, gives the number of computational domains.
- The input value for the fluxes enclosed within each interface, tflux(1:Mvol) and tflux(1:Mvol), are immediately normalized:

```
tflux(1:Mvol) \rightarrow tflux(1:Mvol)/tflux(Nvol).
pflux(1:Mvol) \rightarrow pflux(1:Mvol)/tflux(Nvol).
```

The input  $\Phi_{edge} \equiv \text{phiedge}$  will provide the total toroidal flux; see preset().

• The input value for the toroidal current constraint (Isurf (1:Mvol)) and Ivolume (1:Mvol)) are also immediately normalized, using curror .  $Ivolume \rightarrow Ivolume \cdot \frac{curtor}{\sum_i Isurf_i + Ivolume_i} Isurf \rightarrow Isurf \cdot \frac{curtor}{\sum_i Isurf_i + Ivolume_i}$ 

#### **Current profiles normalization**

In case of a free boundary calculation (Lfreebound=1) and using a current constraint (Lconstraint=3), the current profiles are renormalized in order to match the linking current curtor. More specifically,

$$Isurf_{i} \rightarrow Isurf_{i} \cdot \frac{curtor}{\sum_{i=1}^{Mvol-1} Isurf_{i} + Ivol_{i}} Ivol_{i} \rightarrow Ivol_{i} \cdot \frac{curtor}{\sum_{i=1}^{Mvol-1} Isurf_{i} + Ivol_{i}}$$
(258)

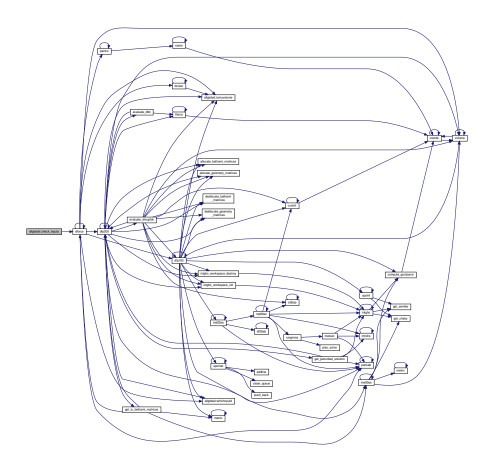
Finally, the volume current in the vacuum region is set to 0.

reading of numericlist reading of locallist reading of globallist reading of diagnosticslist reading of screenlist

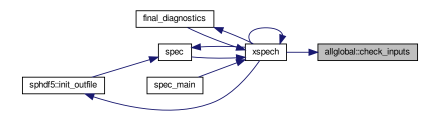
References inputlist::bnc, inputlist::c05factor, inputlist::c05xmax, inputlist::c05xtol, cpus, inputlist::curpol, inputlist::curtor, dforce(), inputlist::dpp, inputlist::dqq, inputlist::drz, inputlist::epsgmres, inputlist::epsilon, inputlist::epsilu, inputlist::escale, inputlist::forcetol, inputlist::gamma, inputlist::gbnbld, inputlist::gbntol, inputlist::helicity, inputlist::igeometry, inputlist::imethod, inputlist::impol, in, inputlist::intor, inputlist::iorder, inputlist::iotatol, inputlist::iprecon, inputlist::istellsym, inputlist::isurf, inputlist::ivolume, inputlist::ladiabatic, inputlist::lautoinitbn, inputlist::lbeltrami, inputlist::lcheck, inputlist::lconstraint, inputlist::lextrap, inputlist::lfindzero, inputlist::lfreebound, inputlist::lgmresprec, inputlist::lhevalues, inputlist::lhevectors, inputlist::lhmatrix, inputlist::linitgues, inputlist::lmatsolver, inputlist::lperturbed, inputlist::lrad, inputlist::lreadgf, inputlist::lreflect, inputlist::lrzaxis, inputlist::lsparse, inputlist::lsvdiota, inputlist::ltiming, inputlist::lzerovac, numerical::machprec, inputlist::mfreeits, inputlist::mmpol, inputlist::mntor, inputlist::mnvol, inputlist::mpol, inputlist::mregular, inputlist::mu, inputlist::mupfits, inputlist::mupftol, inputlist::ndiscrete, inputlist::nfp, inputlist::nitergmres, inputlist::nppts, inputlist::nquad, inputlist::ntor, inputlist::ntoraxis, inputlist::nvol, inputlist::odetol, constants::one, inputlist::opsilon, fileunits::ounit, inputlist::pcondense, inputlist::pflux, inputlist::phiedge, inputlist::pressure, inputlist::pscale, inputlist::rbs, inputlist::rpol, inputlist::rtor, inputlist::rws, numerical::small, inputlist::tflux, inputlist::upsilon, inputlist::vcasingeps, inputlist::vcasingits, inputlist::vcasingper, inputlist::vcasingtol, inputlist::vnc, numerical::vsmall, inputlist::wpoloidal, inputlist::wreadin, inputlist::zbc, constants::zero, and inputlist::zwc.

Referenced by xspech().

Here is the call graph for this function:



Here is the caller graph for this function:



# 8.1.2.2 broadcast\_inputs() subroutine allglobal::broadcast\_inputs broadcast physicslist broadcast numericlist broadcast globallist broadcast locallist broadcast diagnosticslist

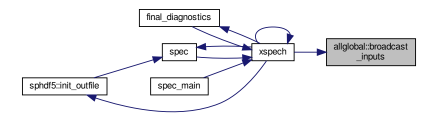
broadcast screenlist

References inputlist::adiabatic, inputlist::c05factor, inputlist::c05xmax, inputlist::c05xtol, cpus, inputlist::curpol, inputlist::curtor, inputlist::dpp, inputlist::dqq, inputlist::drz, inputlist::epsgmres, inputlist::epsilon, inputlist::epsilon, inputlist::epsilon, inputlist::igeometry, inputlist::igeomet

inputlist::imethod, inputlist::impol, inputlist::intor, inputlist::iorder, inputlist::iota, inputlist::iotatol, inputlist::iprecon, inputlist::istellsym, inputlist::isurf, inputlist::ivolume, inputlist::ladiabatic, inputlist::lautoinitbn, inputlist::lbeltrami, inputlist::lcheck, inputlist::lconstraint, inputlist::lerrortype, inputlist::lextrap, inputlist::lfindzero, inputlist::lfreebound, inputlist::lgmresprec, inputlist::lhevalues, inputlist::lhevectors, inputlist::lhmatrix, inputlist::linitgues, inputlist::linitgues, inputlist::lreadgf, inputlist::lreadgf, inputlist::lreadgf, inputlist::lreadgf, inputlist::lreadgf, inputlist::lreadgf, inputlist::lreadgf, inputlist::lreadgf, inputlist::lreadgf, inputlist::maxrndgues, inputlist::mfreeits, inputlist::mnvol, inputlist::mpol, inputlist::mregular, inputlist::mu, inputlist::mupfits, inputlist::mupftol, inputlist::ndiscrete, inputlist::nfp, inputlist::mgrid, inputlist::ndetol, inputlist::nppts, inputlist::nptrj, inputlist::nquad, inputlist::ntor, inputlist::ntoraxis, inputlist::nvol, inputlist::odetol, inputlist::opsilon, fileunits::ounit, inputlist::pcondense, inputlist::pflux, inputlist::phiedge, inputlist::pt, inputlist::pts, inputlist::pts, inputlist::pts, inputlist::pts, inputlist::rtor, inputli

Referenced by xspech().

Here is the caller graph for this function:



# **8.1.2.3** ismyvolume() subroutine allglobal::ismyvolume ( integer, intent(in) *vvol*)

Check if volume vvol is associated to the corresponding MPI node.

The global variable IsMyVolumeValue is updated to 0 or 1, depending on vvol. A value of -1 is set if an error occured.

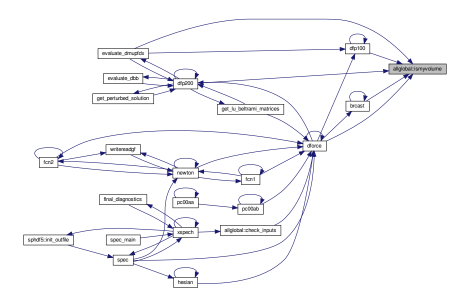
#### **Parameters**

1001	valume to sheek
vvol	volume to check

References is myvolume value, myid, and ncpu.

Referenced by brcast(), dforce(), dfp100(), dfp200(), and evaluate\_dmupfdx().

Here is the caller graph for this function:



## 8.2 constants Module Reference

some constants used throughout the code

# **Variables**

• real, parameter **zero** = 0.0

0

• real, parameter **one** = 1.0

1

• real, parameter **two** = 2.0

2

• real, parameter **three** = 3.0

3

• real, parameter **four** = 4.0

4

• real, parameter **five** = 5.0

5

• real, parameter **six** = 6.0

6

• real, parameter **seven** = 7.0

7

• real, parameter eight = 8.0

8

• real, parameter **nine** = 9.0

9

• real, parameter **ten** = 10.0

10

• real, parameter **eleven** = 11.0

11

• real, parameter **twelve** = 12.0

```
12
• real, parameter hundred = 100.0
• real, parameter thousand = 1000.0
     1000

 real, parameter half = one / two

 real, parameter third = one / three

• real, parameter quart = one / four
• real, parameter fifth = one / five
     1/5
• real, parameter sixth = one / six
• real, parameter pi2 = 6.28318530717958623
• real, parameter pi = pi2 / two

    real, parameter mu0 = 2.0E-07 * pi2

     4\pi \cdot 10^{-7}
• real, parameter goldenmean = 1.618033988749895
     golden mean = (1+\sqrt{5})/2;
• real, parameter version = 3.10
     version of SPEC
```

# 8.2.1 Detailed Description

• real tmanual = 0.0

some constants used throughout the code

# 8.3 cputiming Module Reference

timing variables

#### **Variables**

```
• real manualt = 0.0
• real trzaxis = 0.0
• real rzaxist = 0.0
• real tpackxi = 0.0

    real packxit = 0.0

• real tvolume = 0.0
• real volumet = 0.0
• real tcoords = 0.0
• real coordst = 0.0
• real tbasefn = 0.0
• real basefnt = 0.0
• real tmemory = 0.0
• real memoryt = 0.0
• real tmetrix = 0.0
• real metrixt = 0.0
• real tma00aa = 0.0
• real ma00aat = 0.0
```

- real tmatrix = 0.0
- real matrixt = 0.0
- real tspsmat = 0.0
- real spsmatt = 0.0
- real **tspsint** = 0.0
- real spsintt = 0.0
- real tmp00ac = 0.0
- real mp00act = 0.0
- real tma02aa = 0.0
- real ma02aat = 0.0
- real tpackab = 0.0
- real packabt = 0.0
- real ttr00ab = 0.0
- real **tr00abt** = 0.0
- real tcurent = 0.0
- real curentt = 0.0
- real tdf00ab = 0.0
- real df00abt = 0.0
- real tlforce = 0.0
- real Iforcet = 0.0
- real tintghs = 0.0
- real intghst = 0.0
- real tmtrxhs = 0.0
- real mtrxhst = 0.0
- real **tlbpol** = 0.0
- real **lbpolt** = 0.0
- real tbrcast = 0.0
- real brcastt = 0.0
- real tdfp100 = 0.0
- real dfp100t = 0.0
- real tdfp200 = 0.0
- real dfp200t = 0.0
- real **tdforce** = 0.0
- real dforcet = 0.0
- real **tnewton** = 0.0
- real **newtont** = 0.0
- real tcasing = 0.0
- real casingt = 0.0
- real **tbnorml** = 0.0
- real **bnormlt** = 0.0
- real tjo00aa = 0.0
- real jo00aat = 0.0
- real **tpp00aa** = 0.0
- real **pp00aat** = 0.0
- real **tpp00ab** = 0.0
- real **pp00abt** = 0.0
- real **tbfield** = 0.0
- real bfieldt = 0.0
- real tstzxyz = 0.0
- real stzxyzt = 0.0
- real thesian = 0.0
- real **hesiant** = 0.0
- real **tra00aa** = 0.0
- real **ra00aat** = 0.0
- real tnumrec = 0.0

- real **numrect** = 0.0
- real **tdcuhre** = 0.0
- real dcuhret = 0.0
- real tminpack = 0.0
- real minpackt = 0.0
- real tiqpack = 0.0
- real iqpackt = 0.0
- real trksuite = 0.0
- real rksuitet = 0.0
- real **ti1mach** = 0.0
- real **i1macht** = 0.0
- real **td1mach** = 0.0
- real **d1macht** = 0.0
- real **tilut** = 0.0
- real **ilutt** = 0.0
- real **titers** = 0.0
- real **iterst** = 0.0
- real tsphdf5 = 0.0
- real sphdf5t = 0.0
- real tpreset = 0.0
- real presett = 0.0
- real tglobal = 0.0
- real **globalt** = 0.0
- real txspech = 0.0
- real xspecht = 0.0
- real tinputlist = 0.0
- real inputlistt = 0.0
- real treadin = 0.0
- real twrtend = 0.0

#### 8.3.1 Detailed Description

timing variables

# 8.4 fftw\_interface Module Reference

Interface to FFTW library.

#### **Variables**

type(c\_ptr) planf

FFTW-related (?)

type(c\_ptr) planb

FFTW-related (?)

complex(c\_double\_complex), dimension(:,:,:), allocatable cplxin
 FFTW-related (?)

complex(c\_double\_complex), dimension(:,:,:), allocatable cplxout
 FFTW-related (?)

#### 8.4.1 Detailed Description

Interface to FFTW library.

# 8.5 fileunits Module Reference

central definition of file units to avoid conflicts

#### **Functions/Subroutines**

• subroutine mute (action)

#### **Variables**

• integer iunit = 10

input; used in global/readin:ext.sp, global/wrtend:ext.sp.end

• integer ounit = 6

screen output;

• integer gunit = 13

wall geometry; used in wa00aa

• integer aunit = 11

vector potential; used in ra00aa:.ext.AtAzmn;

• integer dunit = 12

derivative matrix; used in newton:.ext.GF;

• integer hunit = 14

eigenvalues of Hessian; under re-construction;

• integer munit = 14

matrix elements of Hessian;

• integer lunit = 20

local unit; used in lunit+myid: pp00aa:.ext.poincare,.ext.transform;

• integer **vunit** = 15

for examination of adaptive quadrature; used in casing:.ext.vcint;

## 8.5.1 Detailed Description

central definition of file units to avoid conflicts

# 8.6 laplaces Module Reference

...todo...

#### **Variables**

· logical stage1

what is this?

· logical exterior

what is this?

· logical dorm

what is this?

· integer nintervals

what is this?

• integer nsegments

what is this?

• integer ic

what is this?

· integer np4

what is this?

integer np1

what is this?

• integer, dimension(:), allocatable icint

what is this?

· real originalalpha

what is this?

• real, dimension(:), allocatable xpoly

what is this?

• real, dimension(:), allocatable ypoly

what is this?

• real, dimension(:), allocatable phi

what is this?

• real, dimension(:), allocatable phid

what is this?

• real, dimension(:,:), allocatable cc

what is this?

· integer ilength

what is this?

· real totallength

what is this?

· integer niterations

counter; eventually redundant; 24 Oct 12;

· integer iangle

angle; eventually redundant; 24 Oct 12;

· real rmid

used to define local polar coordinate; eventually redundant; 24 Oct 12;

real zmid

used to define local polar coordinate; eventually redundant; 24 Oct 12;

· real alpha

eventually redundant; 24 Oct 12;

#### 8.6.1 Detailed Description

...todo...

## 8.7 newtontime Module Reference

timing of Newton iterations

#### **Variables**

· integer nfcalls

number of calls to get function values (?)

integer ndcalls

number of calls to get derivative values (?)

· real lastcpu

last CPU that called this (?)

## 8.7.1 Detailed Description

timing of Newton iterations

#### 8.8 numerical Module Reference

platform-dependant numerical resolution

#### **Variables**

• real, parameter machprec = 1.11e-16

machine precision: 0.5\*epsilon(one) for 64 bit double precision

• real, parameter **vsmall** = 100\*machprec

very small number

• real, parameter **small** = 10000\*machprec

small number

real, parameter sqrtmachprec = sqrt(machprec)

square root of machine precision

• real, parameter logtolerance = 1.0e-32

this is used to avoid taking alog10(zero); see e.g. dforce;

#### 8.8.1 Detailed Description

platform-dependant numerical resolution

# 8.9 sphdf5 Module Reference

writing the HDF5 output file

#### **Functions/Subroutines**

subroutine init\_outfile

Initialize the interface to the HDF5 library and open the output file.

subroutine mirror\_input\_to\_outfile

Mirror input variables into output file.

subroutine init\_convergence\_output

Prepare convergence evolution output.

subroutine write\_convergence\_output (nDcalls, ForceErr)

Write convergence output (evolution of interface geometry, force, etc).

• subroutine write\_grid

Write the magnetic field on a grid.

• subroutine init\_flt\_output (numTrajTotal)

Initialize field line tracing output group and create array datasets.

• subroutine write\_poincare (offset, data, success)

Write a hyperslab of Poincare data corresponding to the output of one parallel worker.

• subroutine write\_transform (offset, length, Ivol, diotadxup, fiota)

Write the rotational transform output from field line following.

• subroutine finalize\_flt\_output

Finalize Poincare output.

subroutine write\_vector\_potential (sumLrad, allAte, allAze, allAto, allAzo)

Write the magnetic vector potential Fourier harmonics to the output file group /vector\_potential.

· subroutine hdfint

Write the final state of the equilibrium to the output file.

· subroutine finish\_outfile

Close all open HDF5 objects (we know of) and list any remaining still-open objects.

#### **Variables**

· logical, parameter hdfdebug = .false.

global flag to enable verbal diarrhea commenting HDF5 operations

• integer, parameter internalhdf5msg = 0

1: print internal HDF5 error messages; 0: only error messages from sphdf5

· integer hdfier

error flag for HDF5 library

· integer rank

rank of data to write using macros

• integer(hid\_t) file\_id

default file ID used in macros

• integer(hid\_t) space\_id

default dataspace ID used in macros

• integer(hid\_t) dset\_id

default dataset ID used in macros

• integer(hsize t), dimension(1:1) onedims

dimension specifier for one-dimensional data used in macros

• integer(hsize\_t), dimension(1:2) twodims

dimension specifier for two-dimensional data used in macros

integer(hsize t), dimension(1:3) threedims

dimension specifier for three-dimensional data used in macros

· logical grp\_exists

flags used to signal if a group already exists

logical var exists

flags used to signal if a variable already exists

• integer(hid\_t) iteration\_dset\_id

Dataset identifier for "iteration".

• integer(hid\_t) dataspace

dataspace for extension by 1 iteration object

• integer(hid\_t) memspace

memspace for extension by 1 iteration object

integer(hsize\_t), dimension(1) old\_data\_dims

current dimensions of "iterations" dataset

• integer(hsize t), dimension(1) data dims

new dimensions for "iterations" dataset

• integer(hsize\_t), dimension(1) max\_dims

maximum dimensions for "iterations" dataset

• integer(hid\_t) plist\_id

Property list identifier used to activate dataset transfer property.

integer(hid\_t) dt\_ndcalls\_id

Memory datatype identifier (for "nDcalls" dataset in "/grid")

• integer(hid\_t) dt\_energy\_id

Memory datatype identifier (for "Energy" dataset in "/grid")

• integer(hid\_t) dt\_forceerr\_id

Memory datatype identifier (for "ForceErr" dataset in "/grid")

• integer(hid\_t) dt\_irbc\_id

Memory datatype identifier (for "iRbc" dataset in "/grid")

• integer(hid\_t) dt\_izbs\_id

Memory datatype identifier (for "iZbs" dataset in "/grid")

integer(hid\_t) dt\_irbs\_id

Memory datatype identifier (for "iRbs" dataset in "/grid")

• integer(hid\_t) dt\_izbc\_id

Memory datatype identifier (for "iZbc" dataset in "/grid")

integer, parameter rankp =3

rank of Poincare data

integer, parameter rankt =2

rank of rotational transform data

integer(hid\_t) grppoincare

group for Poincare data

integer(hid\_t) dset\_id\_t

Dataset identifier for  $\theta$  coordinate of field line following.

integer(hid\_t) dset\_id\_s

Dataset identifier for s coordinate of field line following.

integer(hid\_t) dset\_id\_r

Dataset identifier for R coordinate of field line following.

integer(hid\_t) dset\_id\_z

Dataset identifier for  ${\cal Z}$  coordinate of field line following.

integer(hid\_t) dset\_id\_success

Dataset identifier for success flag of trajectories to follow.

integer(hid t) filespace t

Dataspace identifier in file for  $\theta$  coordinate of field line following.

• integer(hid\_t) filespace\_s

Dataspace identifier in file for s coordinate of field line following.

integer(hid\_t) filespace\_r

Dataspace identifier in file for R coordinate of field line following.

• integer(hid\_t) filespace\_z

Dataspace identifier in file for  ${\cal Z}$  coordinate of field line following.

integer(hid\_t) filespace\_success

Dataspace identifier in file for success flag of trajectories to follow.

integer(hid t) memspace t

Dataspace identifier in memory for  $\theta$  coordinate of field line following.

• integer(hid t) memspace s

Dataspace identifier in memory for s coordinate of field line following.

integer(hid\_t) memspace\_r

Dataspace identifier in memory for  ${\cal R}$  coordinate of field line following.

• integer(hid\_t) memspace\_z

Dataspace identifier in memory for  ${\cal Z}$  coordinate of field line following.

integer(hid\_t) memspace\_success

Dataspace identifier in memory for success flag of trajectories to follow.

integer(hid\_t) grptransform

group for rotational transform data

• integer(hid t) dset id diotadxup

Dataset identifier for diotadxup (derivative of rotational transform ?)

integer(hid\_t) dset\_id\_fiota

Dataset identifier for fiota ( rotational transform ?)

• integer(hid\_t) filespace\_diotadxup

Dataspace identifier in file for diotadxup.

• integer(hid\_t) filespace\_fiota

Dataspace identifier in file for fiota.

integer(hid\_t) memspace\_diotadxup

Dataspace identifier in memory for diotadxup.

integer(hid\_t) memspace\_fiota

Dataspace identifier in memory for fiota.

• character(len=15), parameter aname = "description"

Attribute name for descriptive info.

integer(hid\_t) attr\_id

Attribute identifier.

• integer(hid\_t) aspace\_id

Attribute Dataspace identifier.

integer(hid\_t) atype\_id

Attribute Datatype identifier.

• integer, parameter arank = 1

Attribure rank.

• integer(hsize\_t), dimension(arank) adims = (/1/)

Attribute dimension.

• integer(size\_t) attrlen

Length of the attribute string.

• character(len=:), allocatable attr\_data

Attribute data.

#### 8.9.1 Detailed Description

writing the HDF5 output file

# 8.10 typedefns Module Reference

type definitions for custom datatypes

#### **Data Types**

· type derivative

 $\mathrm{d}\mathbf{B}/\mathrm{d}\mathbf{X}$  (?) More...

- · type matrixlu
- · type subgrid

used for quantities which have different resolutions in different volumes, e.g. the vector potential More...

## 8.10.1 Detailed Description

type definitions for custom datatypes

## 8.10.2 Data Type Documentation

# 8.10.2.1 type typedefns::derivative $d\mathbf{B}/d\mathbf{X}$ (?)

#### **Class Members**

logical	1	what is this?
integer	vol	Used in coords(); required for global constraint force gradient evaluation.
integer	innout	what is this?
integer	ii	what is this?
integer	irz	what is this?
integer	issym	what is this?

#### **Class Members**

real, dimension(:,:), allocatable	mat	
integer, dimension(:), allocatable	ipivot	

#### 8.10.2.2 type typedefns::matrixlu

**8.10.2.3 type typedefns::subgrid** used for quantities which have different resolutions in different volumes, e.g. the vector potential

#### **Class Members**

real, dimension(:), allocatable	s	coefficients
integer, dimension(:), allocatable	i	indices

# 9 Data Type Documentation

# 9.1 intghs\_module::intghs\_workspace Type Reference

This calculates the integral of something related to matrix-vector-multiplication.

#### **Public Attributes**

real, dimension(:,:), allocatable efmn

This is efmn.

real, dimension(:,:), allocatable ofmn
 This is ofmn.

- real, dimension(:,:), allocatable cfmn
- real, dimension(:,:), allocatable sfmn
- real, dimension(:,:), allocatable evmn
- real, dimension(:,:), allocatable odmn
- real, dimension(:,:), allocatable ijreal
- real, dimension(:,:), allocatable jireal
- real, dimension(:,:), allocatable jkreal
- real, dimension(:,:), allocatable kjreal
- real, dimension(:,:,:), allocatable bloweremn
   real, dimension(:,:,:), allocatable bloweromn
- real, dimension(:,:,:), allocatable **gbupper**
- real, dimension(:,:,:), allocatable blower
- real, dimension(:,:,:,:), allocatable basis

#### 9.1.1 Detailed Description

This calculates the integral of something related to matrix-vector-multiplication.

**Todo** Zhisong might need to update the documentation of this type.

# 9.1.2 Member Data Documentation

**9.1.2.1 efmn** real, dimension(:,:), allocatable intghs\_module::intghs\_workspace::efmn This is efmn.

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```
9.1.2.2 ofmn real, dimension(:,:), allocatable intghs_module::intghs_workspace::ofmn
This is ofmn.
9.1.2.3 cfmn real, dimension(:,:), allocatable intghs_module::intghs_workspace::cfmn
9.1.2.4 sfmn real, dimension(:,:), allocatable intghs_module::intghs_workspace::sfmn
9.1.2.5 evmn real, dimension(:,:), allocatable intghs_module::intghs_workspace::evmn
9.1.2.6 odmn real, dimension(:,:), allocatable intghs_module::intghs_workspace::odmn
9.1.2.7 ijreal real, dimension(:,:), allocatable intghs_module::intghs_workspace::ijreal
9.1.2.8 jireal real, dimension(:,:), allocatable intghs_module::intghs_workspace::jireal
9.1.2.9 jkreal real, dimension(:,:), allocatable intghs_module::intghs_workspace::jkreal
9.1.2.10 kjreal real, dimension(:,:), allocatable intghs_module::intghs_workspace::kjreal
\textbf{9.1.2.11} \quad \textbf{bloweremn} \quad \texttt{real, dimension(:,:,:), allocatable intghs\_module::intghs\_workspace} \leftarrow
::bloweremn
9.1.2.12 bloweromn real, dimension(:,:,:), allocatable intghs_module::intghs_workspace←
::bloweromn
9.1.2.13 gbupper real, dimension(:,:,:), allocatable intghs_module::intghs_workspace::gbupper
9.1.2.14 blower real, dimension(:,:,:), allocatable intghs_module::intghs_workspace::blower
9.1.2.15 basis real, dimension(:,:,:,:), allocatable intghs_module::intghs_workspace::basis
The documentation for this type was generated from the following file:
```

# 10 File Documentation

## 10.1 src/basefn.f90 File Reference

Polynomials evaluation.

• src/intghs.f90

#### **Functions/Subroutines**

• subroutine get\_cheby (lss, lrad, cheby)

Get the Chebyshev polynomials with zeroth, first derivatives.

• subroutine get\_cheby\_d2 (lss, lrad, cheby)

Get the Chebyshev polynomials with zeroth, first and second derivatives The Chebyshev polynomial has been recombined and rescaled. See get\_cheby for more detail.

• subroutine get\_zernike (r, Irad, mpol, zernike)

Get the Zernike polynomials  $\hat{R}_l^m$  with zeroth, first derivatives.

• subroutine get\_zernike\_d2 (r, Irad, mpol, zernike)

Get the Zernike polynomials  $\hat{R}_{l}^{m}$  with zeroth, first, second derivatives.

• subroutine get\_zernike\_rm (r, Irad, mpol, zernike)

Get the Zernike polynomials  $\hat{R}_l^m/r^m$ .

#### 10.1.1 Detailed Description

Polynomials evaluation.

#### 10.1.2 Function/Subroutine Documentation

Get the Chebyshev polynomials with zeroth, first derivatives.

The Chebyshev polynomial has been recombined and rescaled. By doing so, the Chebyshev polynomial satisfy the zero Dirichlet boundary condition on the inner surface of the annulus with reduced ill-conditioning problem. Let  $T_l$  be the Chebyshev polynomial of the first kind with degree l. This subroutine computes

$$\bar{T}_0 = 1,$$

and

$$\bar{T}_l = \frac{T_l - (-1)^l}{l+1}.$$

 $T_l$  are computed iteratively.

$$T_0(s) = 1,$$

$$T_1(s) = s,$$

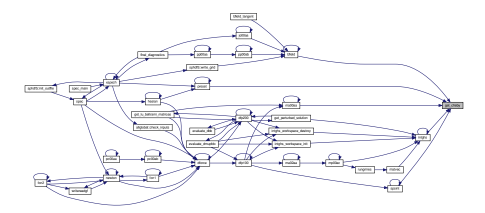
$$T_{l+1}(s) = 2sT_l(s) - T_{l-1}(s).$$

#### **Parameters**

in	Iss	coordinate input Iss
in	Irad	radial resolution
out	cheby	the value, first derivative of Chebyshev polynomial

References constants::one, constants::two, and constants::zero. Referenced by bfield(), intghs(), ma00aa(), preset(), and spsint().

Here is the caller graph for this function:



Get the Chebyshev polynomials with zeroth, first and second derivatives The Chebyshev polynomial has been recombined and rescaled. See get\_cheby for more detail.

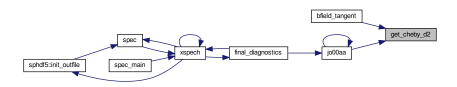
#### **Parameters**

in	Iss	coordinate input lss
in	Irad	radial resolution
out	cheby	the value, first and second derivative of Chebyshev polynomial

References constants::one, constants::two, and constants::zero.

Referenced by bfield tangent(), and jo00aa().

Here is the caller graph for this function:



Get the Zernike polynomials  $\hat{R}_l^m$  with zeroth, first derivatives.

The original Zernike polynomial is defined by The Zernike polynomials take the form

$$Z_l^{-m}(s,\theta) = R_l^m(s)\sin m\theta,$$
  

$$Z_l^m(s,\theta) = R_l^m(s)\cos m\theta,$$

where  $R_l^m(s)$  is a l-th order polynomial given by

$$R_l^m(s) = \sum_{k=0}^{\frac{l-m}{2}} \frac{(-1)^k (l-k)!}{k! \left[\frac{1}{2} (l+m) - k\right]! \left[\frac{1}{2} (l-m) - k\right]!} s^{l-2k},$$

and is only non-zero for  $l \geq m$  and even l-m.

In this subroutine,  $R_l^m(s)$  is computed using the iterative relationship

$$R_l^m(s) = \frac{2(l-1)(2l(l-2)s^2 - m^2 - l(l-2))R_{l-2}^m(s) - l(l+m-2)(l-m-2)R_{l-4}^m(s)}{(l+m)(l-m)(l-2)}$$

For m=0 and m=1, a basis recombination method is used by defining new radial basis functions as

$$\begin{split} \hat{R}_0^0 &= 1, \hat{R}_l^0 &= \frac{1}{l+1} R_l^0 - \frac{(-1)^{l/2}}{l+1}, \\ \hat{R}_1^1 &= s, \hat{R}_l^1 &= \frac{1}{l+1} R_l^1 - \frac{(-1)^{(l-1)/2}}{2} s. \end{split}$$

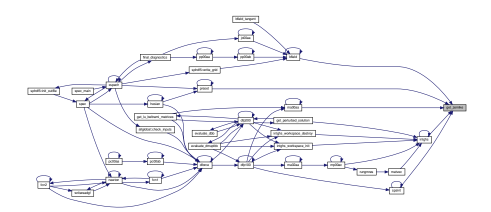
so that the basis scales as  $s^{m+2}$  except for  $\hat{R}_0^0$  and  $\hat{R}_1^1$ , which are excluded from the representation of  $A_{\theta,m,n}$ . For  $m \geq 2$ , the radial basis functions are only rescaled as

$$\hat{R}_l^m = \frac{1}{l+1} R_l^m.$$

#### **Parameters**

in	r	coordinate input, note that this is normalized to $\left[0,1\right]$
in	Irad	radial resolution
in	mpol	poloidal resolution
out	zernike	the value, first derivative of Zernike polynomial

References constants::one, constants::two, and constants::zero. Referenced by bfield(), intghs(), ma00aa(), preset(), and spsint(). Here is the caller graph for this function:



#### 10.1.2.4 get\_zernike\_d2() subroutine get\_zernike\_d2 (

```
real, intent(in) r,
integer, intent(in) lrad,
integer, intent(in) mpol,
real, dimension(0:lrad,0:mpol,0:2), intent(inout) zernike)
```

Get the Zernike polynomials  $\hat{R}^m_l$  with zeroth, first, second derivatives. See get\_zernike for more detail.

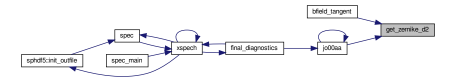
#### **Parameters**

in	r	coordinate input, note that this is normalized to $\left[0,1\right]$
in	Irad	radial resolution
in	mpol	poloidal resolution
out	zernike	the value, first/second derivative of Zernike polynomial

References constants::one, constants::two, and constants::zero.

Referenced by bfield\_tangent(), and jo00aa().

Here is the caller graph for this function:



```
10.1.2.5 \quad get\_zernike\_rm() \quad \texttt{subroutine get}\_zernike\_rm \ (
```

```
real, intent(in) r,
integer, intent(in) lrad,
integer, intent(in) mpol,
real, dimension(0:lrad,0:mpol), intent(inout) zernike)
```

Get the Zernike polynomials  $\hat{R}_l^m/r^m$ .

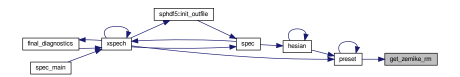
See get zernike for more detail.

#### **Parameters**

in	r	coordinate input, note that this is normalized to $\left[0,1\right]$			
in	Irad	radial resolution			
in	mpol	poloidal resolution			
out	zernike	the value			

References constants::one, constants::two, and constants::zero. Referenced by preset().

Here is the caller graph for this function:



#### 10.2 src/bfield.f90 File Reference

Returns  $\dot{s} \equiv B^s/B^{\zeta}$  and  $\dot{\theta} \equiv B^{\theta}/B^{\zeta}$ .

## **Functions/Subroutines**

• subroutine bfield (zeta, st, Bst)

Compute the magnetic field.

• subroutine bfield\_tangent (zeta, st, Bst) compute the tangential magnetic field

# 10.2.1 Detailed Description

Returns  $\dot{s} \equiv B^s/B^{\zeta}$  and  $\dot{\theta} \equiv B^{\theta}/B^{\zeta}$ .

#### 10.2.2 Function/Subroutine Documentation

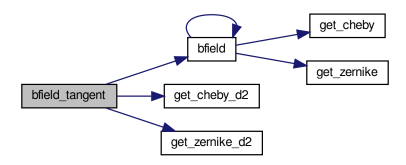
compute the tangential magnetic field

#### **Parameters**

in	zeta	toroidal angle
in	st	radial(s) and poloidal(theta) positions
out	Bst	tangential magnetic field

References allglobal::ate, allglobal::ato, allglobal::aze, allglobal::azo, bfield(), allglobal::cpus, allglobal::gbzeta, get\_cheby\_d2(), get\_zernike\_d2(), constants::half, allglobal::halfmm, allglobal::im, allglobal::in, allglobal::in, allglobal::in, allglobal::in, allglobal::myid, allglobal::ncpu, allglobal::node, allglobal::notstellsym, constants::one, fileunits::ounit, allglobal::regumm, numerical::small, constants::two, numerical::vsmall, inputlist::wmacros, and constants::zero.

Here is the call graph for this function:



# 10.3 src/bnorml.f90 File Reference

Computes  $\mathbf{B}_{Plasma} \cdot \mathbf{e}_{\theta} \times \mathbf{e}_{\zeta}$  on the computational boundary,  $\partial \mathcal{D}$ .

#### **Functions/Subroutines**

• subroutine bnorml (mn, Ntz, efmn, ofmn)  $\textit{Computes } \mathbf{B}_{Plasma} \cdot \mathbf{e}_{\theta} \times \mathbf{e}_{\zeta} \textit{ on the computational boundary, } \partial \mathcal{D}.$ 

## 10.3.1 Detailed Description

Computes  $\mathbf{B}_{Plasma}\cdot\mathbf{e}_{\theta} imes\mathbf{e}_{\zeta}$  on the computational boundary,  $\partial\mathcal{D}$ .

# 10.4 src/brcast.f90 File Reference

Broadcasts Beltrami fields, profiles, . . .

## **Functions/Subroutines**

• subroutine brcast (Ivol)

Broadcasts Beltrami fields, profiles, . . .

#### 10.4.1 Detailed Description

Broadcasts Beltrami fields, profiles, . . .

# 10.5 src/casing.f90 File Reference

Constructs the field created by the plasma currents, at an arbitrary, external location using virtual casing.

#### **Functions/Subroutines**

- subroutine casing (teta, zeta, gBn, icasing)
   Constructs the field created by the plasma currents, at an arbitrary, external location using virtual casing.
- subroutine dvcfield (Ndim, tz, Nfun, vcintegrand)

  Differential virtual casing integrand.

# 10.5.1 Detailed Description

Constructs the field created by the plasma currents, at an arbitrary, external location using virtual casing.

#### 10.6 src/coords.f90 File Reference

Calculates coordinates,  $\mathbf{x}(s, \theta, \zeta) \equiv R \mathbf{e}_R + Z \mathbf{e}_Z$ , and metrics, using FFTs.

#### **Functions/Subroutines**

• subroutine coords (Ivol, Iss, Lcurvature, Ntz, mn) Calculates coordinates,  $\mathbf{x}(s,\theta,\zeta) \equiv R\,\mathbf{e}_R + Z\,\mathbf{e}_Z$ , and metrics, using FFTs.

#### 10.6.1 Detailed Description

Calculates coordinates,  $\mathbf{x}(s, \theta, \zeta) \equiv R \mathbf{e}_R + Z \mathbf{e}_Z$ , and metrics, using FFTs.

#### 10.7 src/curent.f90 File Reference

Computes the plasma current,  $I \equiv \int B_{\theta} d\theta$ , and the "linking" current,  $G \equiv \int B_{\zeta} d\zeta$ .

#### **Functions/Subroutines**

• subroutine curent (Ivol, mn, Nt, Nz, iflag, IdItGp)

Computes the plasma current,  $I \equiv \int B_{\theta} d\theta$ , and the "linking" current,  $G \equiv \int B_{\zeta} d\zeta$ .

#### 10.7.1 Detailed Description

Computes the plasma current,  $I \equiv \int B_{\theta} d\theta$ , and the "linking" current,  $G \equiv \int B_{\zeta} d\zeta$ .

# 10.8 src/df00ab.f90 File Reference

Evaluates volume integrals, and their derivatives w.r.t. interface geometry, using "packed" format.

#### **Functions/Subroutines**

subroutine df00ab (pNN, xi, Fxi, DFxi, Ldfjac, iflag)
 Evaluates volume integrals, and their derivatives w.r.t. interface geometry, using "packed" format.

#### 10.8.1 Detailed Description

Evaluates volume integrals, and their derivatives w.r.t. interface geometry, using "packed" format.

#### 10.9 src/dforce.f90 File Reference

Calculates  $\mathbf{F}(\mathbf{x})$ , where  $\mathbf{x} \equiv \{\text{geometry}\} \equiv \{R_{i,v}, Z_{i,v}\}$  and  $\mathbf{F} \equiv [[p+B^2/2]] + \{\text{spectral constraints}\}$ , and  $\nabla \mathbf{F}$ .

#### **Functions/Subroutines**

- subroutine dforce (NGdof, position, force, LComputeDerivatives, LComputeAxis)  $\textit{Calculates} \ \mathbf{F}(\mathbf{x}), \textit{where} \ \mathbf{x} \equiv \{\textit{geometry}\} \equiv \{R_{i,v}, Z_{i,v}\} \textit{ and } \mathbf{F} \equiv [[p+B^2/2]] + \{\textit{spectral constraints}\}, \textit{and } \nabla \mathbf{F}.$
- subroutine fndiff\_dforce (NGdof)

#### 10.9.1 Detailed Description

Calculates  $\mathbf{F}(\mathbf{x})$ , where  $\mathbf{x} \equiv \{\text{geometry}\} \equiv \{R_{i,v}, Z_{i,v}\}$  and  $\mathbf{F} \equiv [[p+B^2/2]] + \{\text{spectral constraints}\}$ , and  $\nabla \mathbf{F}$ .

# 10.10 src/dfp100.f90 File Reference

Split the work between MPI nodes and evaluate the global constraint.

#### Functions/Subroutines

subroutine dfp100 (Ndofgl, x, Fvec, LComputeDerivatives)
 Split the work between MPI nodes and evaluate the global constraint.

## 10.10.1 Detailed Description

Split the work between MPI nodes and evaluate the global constraint.

#### 10.10.2 Function/Subroutine Documentation

Split the work between MPI nodes and evaluate the global constraint.

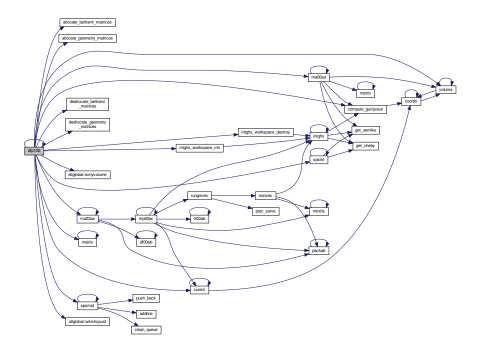
#### **Parameters**

Ndofgl	
X	
Fvec	
LComputeDerivatives	

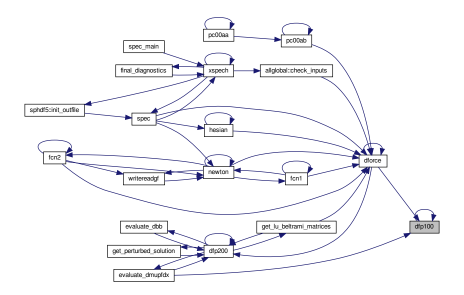
References allocate beltrami matrices(), allocate geometry matrices(), compute guvijsave(), allglobal::cpus, curent(), inputlist::curpol, allglobal::dbdx, allglobal::ddttcc, allglobal::ddttcs, allglobal::ddttcs, allglobal::ddttcs, allglobal::ddtzcc, allglobal::ddtzcs, allglobal::ddtzsc, allglobal::ddtzsc, allglobal::ddtzsc, allglobal::ddtzcs, allglobal::ddzzsc, allglobal::ddzzss, deallocate beltrami matrices(), deallocate geometry matrices(), dfp100(), allglobal::dma, allglobal::dmb, allglobal::dmd, allglobal::dmg, allglobal::dpflux, allglobal::dtoocc, allglobal::dtoocc, allglobal::dtoosc, allglobal::dtooss, allglobal::guvijsave, constants::half, allglobal::iconstraintok, inputlist::igeometry, allglobal::imagneticok, allglobal::in, intghs workspace destroy(), intghs workspace init(), allglobal::ipdtdpf, allglobal::iguad, allglobal::ismyvolume(), allglobal::ismyvolumevalue, inputlist::isurf, allglobal::izbs, inputlist::lconstraint, allglobal::lcoordinatesingularity, inputlist::lfreebound, allglobal::liluprecond, allglobal::localconstraint, allglobal::lplasmaregion, inputlist::lrad, allglobal::lsavedguvij, allglobal::lvacuumregion, ma00aa(), ma02aa(), matrix(), allglobal::mbpsi, allglobal::mn, allglobal::mpi\_comm\_spec, constants::mu0, allglobal::myid, allglobal::nadof, allglobal::ncpu, allglobal::notmatrixfree, allglobal::nt, inputlist::nvol, allglobal::nz, constants::one, fileunits::ounit, constants::pi, constants::pi2, allglobal::solution, spsint(), spsmat(), allglobal::tdstcc, allglobal::tdstcs, allglobal::tdstcs, allglobal::tdstss, allglobal::tdszcc, allglobal::tdszcs, allglobal::tdszsc, allglobal::tdszss, allglobal::ttsscc, allglobal::ttsscs, allglobal::ttsssc, allglobal::ttsssc, constants::two, volume(), allglobal::whichcpuid(), inputlist::wmacros, allglobal::xoffset, and constants::zero.

Referenced by dforce(), dfp100(), and evaluate dmupfdx().

Here is the call graph for this function:



Here is the caller graph for this function:



# 10.11 src/dfp200.f90 File Reference

Given the field consistent with the constraints and the geometry, computes local quantites related to the force evaluation.

## **Functions/Subroutines**

• subroutine dfp200 (LcomputeDerivatives, vvol)

Given the field consistent with the constraints and the geometry, computes local quantites related to the force evaluation.

subroutine get lu beltrami matrices (vvol, oBl, NN)

get LU Beltrami matrices

subroutine get\_perturbed\_solution (vvol, oBI, NN)

This routine evaluates the value of the magnetic field once the interface is perturbed using matrix perturbation theory.

• subroutine evaluate dmupfdx (innout, idof, ii, issym, irz)

Evaluate mu and psip derivatives and store them in dmupfdx.

• subroutine evaluate\_dbb (Ivol, idof, innout, issym, irz, ii, dBB, XX, YY, length, dRR, dZZ, dII, dLL, dPP, Ntz, LcomputeDerivatives)

Evaluate the derivative of the square of the magnetic field modulus. Add spectral constraint derivatives if required.

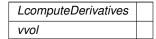
## 10.11.1 Detailed Description

Given the field consistent with the constraints and the geometry, computes local quantites related to the force evaluation.

#### 10.11.2 Function/Subroutine Documentation

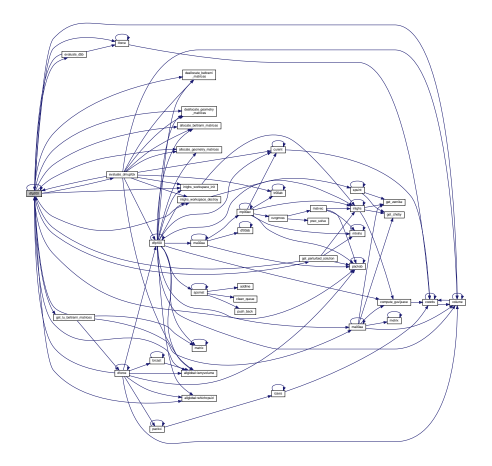
Given the field consistent with the constraints and the geometry, computes local quantites related to the force evaluation.

#### **Parameters**

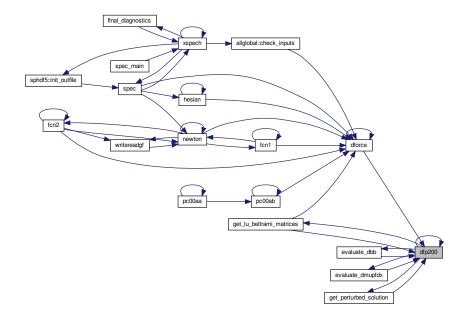


References inputlist::adiabatic, allocate\_beltrami\_matrices(), allocate\_geometry\_matrices(), allglobal::ato, allglobal::aze, allglobal::bbweight, allglobal::bemn, allglobal::bomn, allglobal::btemn, allglobal::cfmn, allglobal::comn, allglobal::cosi, allglobal::cpus, allglobal::dbbdmp, allglobal::dbdx, deallocate beltrami matrices(), deallocate\_geometry\_matrices(), allglobal::dessian, allglobal::dffdrz, dfp200(), allglobal::diotadxup, allglobal::ditgpdxtp, allglobal::dma, allglobal::dmb, allglobal::dmd, allglobal::dmg, allglobal::dmupfdx, allglobal::dpflux, allglobal::drij, allglobal::drodr, allglobal::drodz, allglobal::dtflux, allglobal::dvolume, allglobal::dzij, allglobal::dzodr, allglobal::dzodr, allglobal::efmn, inputlist::epsilon, evaluate\_dbb(), evaluate\_dmupfdx(), allglobal::evmn, inputlist::gamma, get\_lu\_beltrami\_matrices(), get\_perturbed\_solution(), allglobal::guvij, constants::half, allglobal::hessian, allglobal::iemn, inputlist::igeometry, allglobal::ijreal, allglobal::im, allglobal::in, intghs workspace destroy(), intghs workspace init(), allglobal::iomn, allglobal::iguad, allglobal::irbc, allglobal::irbs, allglobal::irij, allglobal:ismyvolume(), allglobal::ismyvolumevalue, allglobal::izbc, allglobal::izbs, allglobal::izij, inputlist::lcheck, inputlist::lconstraint, allglobal::lcoordinatesingularity, inputlist::lextrap, inputlist::lfindzero, lforce(), inputlist::lfreebound, allglobal::lgdof, allglobal::lhessianallocated, allglobal::localconstraint, allglobal::lplasmaregion, inputlist::lrad, allglobal::lvacuumregion, allglobal::lmns. allglobal::mn, allglobal::mne, allglobal::mns, allglobal::mpi\_comm\_spec, inputlist::mpol, allglobal::mmpp. inputlist::mu, allglobal::myid, allglobal::nadof, allglobal::ncpu, allglobal::notstellsym, allglobal::nt, inputlist::ntor, allglobal::ntz, inputlist::nvol, allglobal::nz, allglobal::odmn, allglobal::ofmn, constants::one, fileunits::ounit, packab(), inputlist::pscale, allglobal::psifactor, allglobal::rij, allglobal::semn, allglobal::sfmn, allglobal::sg, allglobal::simn, allglobal::sini, numerical::small, allglobal::solution, allglobal::somn, allglobal::sweight, inputlist::tflux, allglobal::trij, constants::two, allglobal::tzij, volume(), allglobal::vvolume, allglobal::whichcpuid(), inputlist::wmacros, allglobal::yesstellsym, constants::zero, and allglobal::zij.

Here is the call graph for this function:



Here is the caller graph for this function:



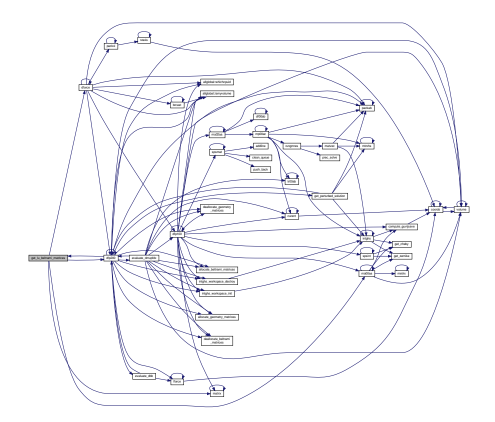
# Parameters

vvol	
oBI	
NN	

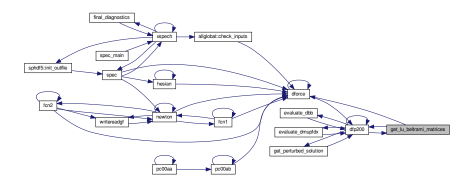
References allglobal::cpus, allglobal::dbdx, dforce(), dfp200(), allglobal::dma, allglobal::dmb, allglobal::dmd, allglobal::dmg, constants::half, allglobal::iquad, inputlist::lconstraint, allglobal::lcoordinatesingularity, allglobal::lplasmaregion, inputlist::lrad, allglobal::lsavedguvij, allglobal::lvacuumregion, ma00aa(), matrix(), allglobal::mn, allglobal::mne, allglobal::mpi\_comm\_spec, inputlist::mu, allglobal::myid, allglobal::ncpu, allglobal::nt, allglobal::nz, constants::one, fileunits::ounit, allglobal::solution, constants::two, inputlist::wmacros, and constants::zero.

Referenced by dfp200().

Here is the call graph for this function:



Here is the caller graph for this function:



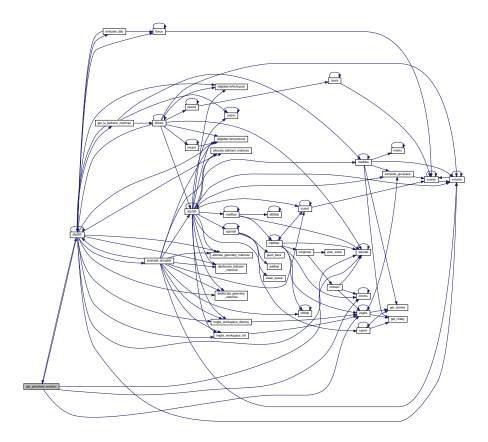
This routine evaluates the value of the magnetic field once the interface is perturbed using matrix perturbation theory.

## **Parameters**

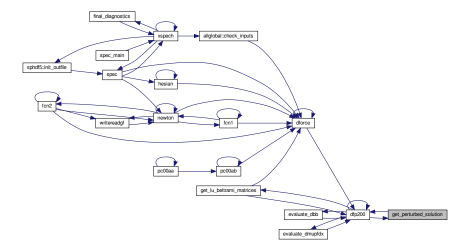
vvol	
oBI	
NN	

References allglobal::cpus, allglobal::dbdx, dfp200(), allglobal::dma, allglobal::dmb, allglobal::dmd, allglobal::dmg, allglobal::dpflux, allglobal::dtflux, constants::half, intghs(), allglobal::iquad, inputlist::lconstraint, inputlist::lrad, allglobal::mn, allglobal::mpi\_comm\_spec, mtrxhs(), inputlist::mu, allglobal::myid, allglobal::nadof, allglobal::ncpu, constants::one, fileunits::ounit, packab(), allglobal::solution, constants::two, inputlist::wmacros, and constants::zero. Referenced by dfp200().

Here is the call graph for this function:



Here is the caller graph for this function:



integer issym,
integer irz )

Evaluate mu and psip derivatives and store them in dmupfdx.

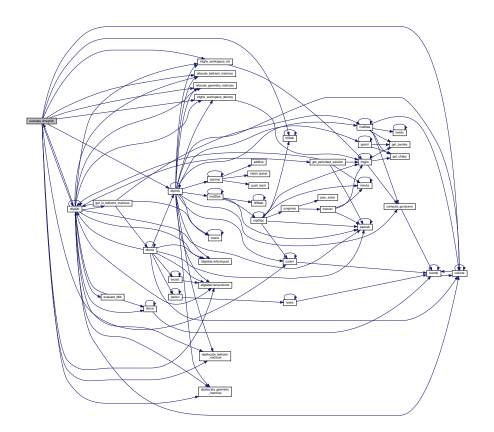
#### **Parameters**

innout	
idof	
ii	
issym	
irz	

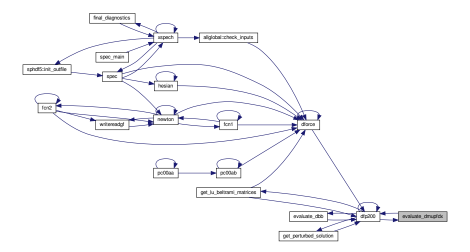
References allocate\_beltrami\_matrices(), allocate\_geometry\_matrices(), allglobal::ate, allglobal::ato, allglobal::ate, allglobal::ate, allglobal::ate, allglobal::ate, allglobal::ate, allglobal::ate, allglobal::dbdx, deallocate\_beltrami\_matrices(), deallocate\_geometry\_matrices(), dfp100(), dfp200(), allglobal::diotadxup, allglobal::ditgpdxtp, allglobal::dma, allglobal::dmb, allglobal::dmd, allglobal::dmg, allglobal::dmupfdx, allglobal::dpflux, inputlist::drz, allglobal::dtflux, allglobal::dvolume, constants::half, inputlist::igeometry, allglobal::im, allglobal::in, intghs\_workspace\_destroy(), intghs\_workspace\_init(), allglobal::ipdtdpf, allglobal::iquad, allglobal::irbc, allglobal::irbs, allglobal::ismyvolume(), allglobal::ismyvolumevalue, allglobal::izbc, allglobal::izbs, inputlist::lcheck, inputlist::lconstraint, allglobal::lcoordinatesingularity, inputlist::lfreebound, allglobal::lmns, allglobal::localconstraint, allglobal::lplasmaregion, inputlist::lrad, allglobal::lvacuumregion, allglobal::mn, allglobal::mn, allglobal::mn, allglobal::mn, allglobal::mn, allglobal::mpi\_comm\_spec, inputlist::mu, inputlist::mupftol, allglobal::myid, allglobal::ncpu, allglobal::ngdof, allglobal::nt, inputlist::nvol, allglobal::nz, constants::one, fileunits::ounit, allglobal::psifactor, allglobal::rij, numerical::small, allglobal::sweight, tr00ab(), constants::two, volume(), allglobal::vvolume, inputlist::wmacros, allglobal::xoffset, constants::zero, and allglobal::zij.

Referenced by dfp200().

Here is the call graph for this function:



Here is the caller graph for this function:



```
integer idof,
integer innout,
integer issym,
integer irz,
integer ii,
real, dimension(1:ntz,-1:2) dBB,
real, dimension(1:ntz) XX,
real, dimension(1:ntz) YY,
real, dimension(1:ntz) length,
real, dimension(1:ntz,-1:2) dRR,
real, dimension(1:ntz,-1:2) dZZ,
real, dimension(1:ntz) dII,
real, dimension(1:ntz) dLL,
real, dimension(1:ntz) dPP,
integer Ntz,
logical, intent(in) LcomputeDerivatives )
```

Evaluate the derivative of the square of the magnetic field modulus. Add spectral constraint derivatives if required.

### **Parameters**

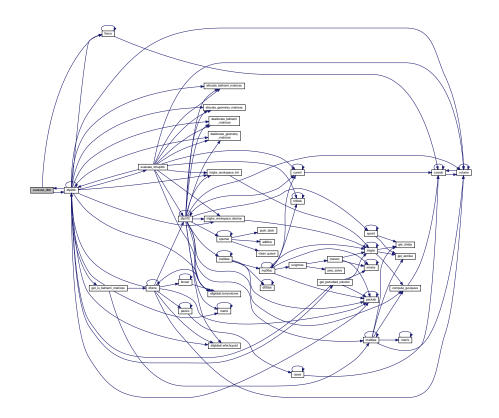
Ivol	
idof	
innout	
issym	
irz	
ii	
dBB	
XX	
YY	
length	
dRR	
dZZ	

#### **Parameters**

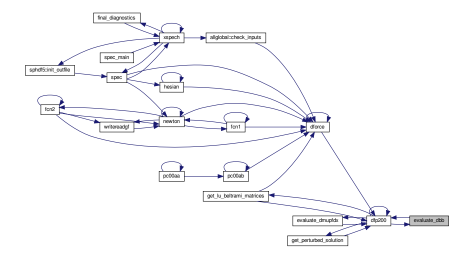
dII	
dLL	
dPP	
Ntz	
LcomputeDerivatives	

References inputlist::adiabatic, allglobal::bbweight, allglobal::cfmn, allglobal::comn, allglobal::cosi, allglobal::cpus, allglobal::dbbdmp, allglobal::dbdx, allglobal::dffdrz, dfp200(), allglobal::dpflux, allglobal::drij, allglobal::drodr, allglobal::drodz, inputlist::drz, allglobal::dvolume, allglobal::dzij, allglobal::dzodr, allglobal::dzodz, allglobal::efmn, inputlist::epsilon, allglobal::evmn, inputlist::gamma, allglobal::guvij, constants::half, inputlist::igeometry, allglobal::ijreal, allglobal::im, allglobal::iquad, allglobal::irbc, allglobal::irbs, allglobal::irij, allglobal::izbc, allglobal::izbc, allglobal::izbc, allglobal::izbc, allglobal::izbc, allglobal::iguad, allglobal::ilcoordinatesingularity, lforce(), allglobal::lgdof, allglobal::localconstraint, allglobal::lplasmaregion, inputlist::lrad, allglobal::lvacuumregion, allglobal::mmpp, allglobal::mnn, allglobal::mns, allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::nadof, allglobal::ncpu, allglobal::ngdof, allglobal::notstellsym, allglobal::nt, inputlist::ntor, inputlist::nvol, allglobal::nz, allglobal::odmn, allglobal::ofmn, constants::one, fileunits::ounit, inputlist::pscale, allglobal::psifactor, allglobal::rij, allglobal::sfmn, allglobal::spin, allglobal::simn, allglobal::simi, numerical::small, allglobal::sweight, allglobal::trij, constants::two, allglobal::tzij, allglobal::vvolume, inputlist::wmacros, allglobal::xoffset, constants::zero, and allglobal::zij. Referenced by dfp200().

Here is the call graph for this function:



Here is the caller graph for this function:



# 10.12 src/global.f90 File Reference

Defines input namelists and global variables, and opens some output files.

## **Data Types**

- · type typedefns::subgrid
  - used for quantities which have different resolutions in different volumes, e.g. the vector potential More...
- · type typedefns::matrixlu
- type typedefns::derivative
  - $\mathrm{d}\mathbf{B}/\mathrm{d}\mathbf{X}$  (?) More...

# Modules

- module constants
  - some constants used throughout the code
- · module numerical
  - platform-dependant numerical resolution
- module fileunits
  - central definition of file units to avoid conflicts
- · module cputiming
  - timing variables
- module typedefns
  - type definitions for custom datatypes
- · module allglobal
  - global variable storage used as "workspace" throughout the code
- module fftw\_interface
  - Interface to FFTW library.

### **Functions/Subroutines**

- subroutine fileunits::mute (action)
- subroutine allglobal::build\_vector\_potential (Ivol, iocons, aderiv, tderiv)
- subroutine allglobal::set\_mpi\_comm (comm)

- subroutine allglobal::read\_inputlists\_from\_file ()
- subroutine allglobal::check\_inputs ()
- · subroutine allglobal::broadcast inputs
- · subroutine allglobal::wrtend

The restart file is written.

• subroutine allglobal::ismyvolume (vvol)

Check if volume vvol is associated to the corresponding MPI node.

• subroutine allglobal::whichcpuid (vvol, cpu\_id)

Returns which MPI node is associated to a given volume.

### **Variables**

```
• real, parameter constants::zero = 0.0
```

• real, parameter constants::one = 1.0

1

• real, parameter constants::two = 2.0

2

• real, parameter constants::three = 3.0

.3

• real, parameter constants::four = 4.0

4

• real, parameter constants::five = 5.0

5

• real, parameter constants::six = 6.0

6

• real, parameter constants::seven = 7.0

7

• real, parameter constants::eight = 8.0

8

• real, parameter constants::nine = 9.0

9

• real, parameter constants::ten = 10.0

10

• real, parameter constants::eleven = 11.0

11

• real, parameter constants::twelve = 12.0

12

• real, parameter constants::hundred = 100.0

100

• real, parameter constants::thousand = 1000.0

1000

• real, parameter **constants::half** = one / two

.,\_

• real, parameter **constants::third** = one / three

1/3

• real, parameter **constants::quart** = one / four

1/4

• real, parameter constants::fifth = one / five

1/5

• real, parameter constants::sixth = one / six

```
1/6
• real, parameter constants::pi2 = 6.28318530717958623 2\pi
• real, parameter constants::pi = pi2 / two
```

• real, parameter **constants::mu0** = 2.0E-07\*pi2 $4\pi\cdot10^{-7}$ 

real, parameter constants::goldenmean = 1.618033988749895

golden mean =  $(1 + \sqrt{5})/2$ ;

• real, parameter constants::version = 3.10

version of SPEC

• real, parameter numerical::machprec = 1.11e-16

machine precision: 0.5\*epsilon(one) for 64 bit double precision

• real, parameter **numerical::vsmall** = 100\*machprec

very small number

small number

• real, parameter **numerical::small** = 10000\*machprec

• real, parameter **numerical::sqrtmachprec** = sqrt(machprec)

square root of machine precision

real, parameter numerical::logtolerance = 1.0e-32

this is used to avoid taking alog10(zero); see e.g. dforce;

• integer fileunits::iunit = 10

input; used in global/readin:ext.sp, global/wrtend:ext.sp.end

• integer fileunits::ounit = 6

screen output;

• integer fileunits::gunit = 13

wall geometry; used in wa00aa

• integer fileunits::aunit = 11

vector potential; used in ra00aa:.ext.AtAzmn;

• integer fileunits::dunit = 12

derivative matrix; used in newton:.ext.GF;

• integer fileunits::hunit = 14

eigenvalues of Hessian; under re-construction;

• integer fileunits::munit = 14

matrix elements of Hessian;

• integer fileunits::lunit = 20

local unit; used in lunit+myid: pp00aa:.ext.poincare,.ext.transform;

• integer fileunits::vunit = 15

for examination of adaptive quadrature; used in casing:.ext.vcint;

- real cputiming::tmanual = 0.0
- real cputiming::manualt = 0.0
- real cputiming::trzaxis = 0.0
- real cputiming::rzaxist = 0.0
- real cputiming::tpackxi = 0.0
- real cputiming::packxit = 0.0
- real cputiming::tvolume = 0.0
- real cputiming::volumet = 0.0
- real cputiming::tcoords = 0.0
- real cputiming::coordst = 0.0
- real cputiming::tbasefn = 0.0
   real cputiming::basefnt = 0.0
- real cputiming::tmemory = 0.0

- real cputiming::memoryt = 0.0
- real cputiming::tmetrix = 0.0
- real cputiming::metrixt = 0.0
- real cputiming::tma00aa = 0.0
- real cputiming::ma00aat = 0.0
- real cputiming::tmatrix = 0.0
- real cputiming::matrixt = 0.0
- real cputiming::tspsmat = 0.0
- real cputiming::spsmatt = 0.0
- real cputiming::tspsint = 0.0
- real cputiming::spsintt = 0.0
- real cputiming::tmp00ac = 0.0
- real cputiming::mp00act = 0.0
- real cputiming::tma02aa = 0.0
- real cputiming::ma02aat = 0.0
- real cputiming::tpackab = 0.0
- real cputiming::packabt = 0.0
- real cputiming::ttr00ab = 0.0
- real cputiming::tr00abt = 0.0
- real cputiming::tcurent = 0.0
- real cputiming::curentt = 0.0
- real cputiming::tdf00ab = 0.0
- real cputiming::df00abt = 0.0
- real cputiming::tlforce = 0.0
- real cputiming::Iforcet = 0.0
- real cputiming::tintghs = 0.0
- real cputiming::intghst = 0.0
- real cputiming::tmtrxhs = 0.0
- real cputiming::mtrxhst = 0.0
- real cputiming::tlbpol = 0.0
- real cputiming::lbpolt = 0.0
- real cputiming::tbrcast = 0.0
- real cputiming::brcastt = 0.0
- real cputiming::tdfp100 = 0.0
- real cputiming::dfp100t = 0.0
- real cputiming::tdfp200 = 0.0
- real cputiming::dfp200t = 0.0
- real cputiming::tdforce = 0.0
- real cputiming::dforcet = 0.0
- real cputiming::tnewton = 0.0 real cputiming::newtont = 0.0
- real cputiming::tcasing = 0.0
- real cputiming::casingt = 0.0
- real cputiming::tbnorml = 0.0
- real cputiming::bnormlt = 0.0
- real cputiming::tjo00aa = 0.0
- real cputiming::jo00aat = 0.0
- real cputiming::tpp00aa = 0.0
- real cputiming::pp00aat = 0.0
- real cputiming::tpp00ab = 0.0
- real cputiming::pp00abt = 0.0
- real cputiming::tbfield = 0.0
- real cputiming::bfieldt = 0.0
- real cputiming::tstzxyz = 0.0
- real cputiming::stzxyzt = 0.0

- real cputiming::thesian = 0.0
- real cputiming::hesiant = 0.0
- real cputiming::tra00aa = 0.0
- real cputiming::ra00aat = 0.0
- real cputiming::tnumrec = 0.0
- real cputiming::numrect = 0.0
- real cputiming::tdcuhre = 0.0
- real cputiming::dcuhret = 0.0
- real cputiming::tminpack = 0.0
- real cputiming::minpackt = 0.0
- real cputiming::tiqpack = 0.0
- real cputiming::iqpackt = 0.0
- real cputiming::trksuite = 0.0
- real cputiming::rksuitet = 0.0
- real cputiming::ti1mach = 0.0
- real cputiming::i1macht = 0.0
- real cputiming::td1mach = 0.0
- real cputiming::d1macht = 0.0
- real cputiming::tilut = 0.0
- real cputiming::ilutt = 0.0
- real cputiming::titers = 0.0
- real cputiming::iterst = 0.0
- real cputiming::tsphdf5 = 0.0
- real cputiming::sphdf5t = 0.0
- real cputiming::tpreset = 0.0
- real cputiming::presett = 0.0
- real cputiming::tglobal = 0.0
- real cputiming::globalt = 0.0
- real cputiming::txspech = 0.0
- real cputiming::xspecht = 0.0
- real cputiming::tinputlist = 0.0
- real cputiming::inputlistt = 0.0
- real cputiming::treadin = 0.0
- real cputiming::twrtend = 0.0
- integer allglobal::myid

MPI rank of current CPU.

· integer allglobal::ncpu

number of MPI tasks

• integer allglobal::ismyvolumevalue

flag to indicate if a CPU is operating on its assigned volume

· real allglobal::cpus

initial time

• integer allglobal::mpi\_comm\_spec

SPEC MPI communicator.

- logical allglobal::skip\_write = .false.
- real allglobal::pi2nfp
- real allglobal::pi2pi2nfp
- real allglobal::pi2pi2nfphalf
- real allglobal::pi2pi2nfpquart
- character(len=1000) allglobal::ext
- real aligiobal::forceerr

total force-imbalance

real allglobal::energy

MHD energy.

- real, dimension(:), allocatable allglobal::ipdt
- real, dimension(:,:), allocatable allglobal::ipdtdpf

Toroidal pressure-driven current.

- · integer allglobal::mvol
- logical allglobal::yesstellsym

internal shorthand copies of Istellsym, which is an integer input;

· logical allglobal::notstellsym

internal shorthand copies of Istellsym, which is an integer input;

- · logical allglobal::yesmatrixfree
- logical allglobal::notmatrixfree

to use matrix-free method or not

real, dimension(:,:), allocatable allglobal::cheby

local workspace for evaluation of Chebychev polynomials

real, dimension(:,:,:), allocatable allglobal::zernike

local workspace for evaluation of Zernike polynomials

• real, dimension(:,:,:), allocatable allglobal::tt

derivatives of Chebyshev polynomials at the inner and outer interfaces;

• real, dimension(:,:,:,:), allocatable allglobal::rtt

derivatives of Zernike polynomials at the inner and outer interfaces;

• real, dimension(:,:), allocatable allglobal::rtm

 $r^{m}$  term of Zernike polynomials at the origin

· real, dimension(:), allocatable allglobal::zernikedof

Zernike degree of freedom for each m.

• integer allglobal::mne

enhanced resolution for metric elements

• integer, dimension(:), allocatable allglobal::ime

enhanced poloidal mode numbers for metric elements

• integer, dimension(:), allocatable allglobal::ine

enhanced toroidal mode numbers for metric elements

integer allglobal::mns

enhanced resolution for straight field line transformation

• integer, dimension(:), allocatable allglobal::ims

enhanced poloidal mode numbers for straight field line transformation

• integer, dimension(:), allocatable allglobal::ins

enhanced toroidal mode numbers for straight field line transformation

• integer allglobal::Impol

what is this?

• integer allglobal::Intor

what is this?

· integer allglobal::smpol

what is this?

integer allglobal::sntor

what is this?

• real allglobal::xoffset = 1.0

used to normalize NAG routines (which ones exacly where?)

logical, dimension(:), allocatable allglobal::imagneticok

used to indicate if Beltrami fields have been correctly constructed;

logical allglobal::iconstraintok

Used to break iteration loops of slaves in the global constraint minimization.

real, dimension(:,:), allocatable allglobal::beltramierror

to store the integral of |curlB-mu\*B| computed by jo00aa;

integer allglobal::mn

total number of Fourier harmonics for coordinates/fields; calculated from Mpol, Ntor in readin()

integer, dimension(:), allocatable allglobal::im

poloidal mode numbers for Fourier representation

• integer, dimension(:), allocatable allglobal::in

toroidal mode numbers for Fourier representation

real, dimension(:), allocatable allglobal::halfmm

I saw this already somewhere...

real, dimension(:), allocatable allglobal::regumm

I saw this already somewhere...

• real aligiobal::rscale

no idea

• real, dimension(:,:), allocatable allglobal::psifactor

no idea

real, dimension(:,:), allocatable allglobal::inifactor

no idea

• real, dimension(:), allocatable allglobal::bbweight

weight on force-imbalance harmonics; used in dforce()

real, dimension(:), allocatable allglobal::mmpp

spectral condensation factors

• real, dimension(:,:), allocatable allglobal::irbc

cosine R harmonics of interface surface geometry; stellarator symmetric

• real, dimension(:,:), allocatable allglobal::izbs

sine Z harmonics of interface surface geometry; stellarator symmetric

• real, dimension(:,:), allocatable allglobal::irbs

sine R harmonics of interface surface geometry; non-stellarator symmetric

real, dimension(:,:), allocatable allglobal::izbc

cosine Z harmonics of interface surface geometry; non-stellarator symmetric

real, dimension(:,:), allocatable allglobal::drbc

cosine R harmonics of interface surface geometry; stellarator symmetric; linear deformation

real, dimension(:,:), allocatable allglobal::dzbs

sine Z harmonics of interface surface geometry; stellarator symmetric; linear deformation

real, dimension(:,:), allocatable allglobal::drbs

sine R harmonics of interface surface geometry; non-stellarator symmetric; linear deformation

real, dimension(:,:), allocatable allglobal::dzbc

cosine Z harmonics of interface surface geometry; non-stellarator symmetric; linear deformation

• real, dimension(:,:), allocatable allglobal::irij

interface surface geometry; real space

• real, dimension(:,:), allocatable allglobal::izij

interface surface geometry; real space

• real, dimension(:,:), allocatable allglobal::drij

interface surface geometry; real space

real, dimension(:,:), allocatable allglobal::dzij

interface surface geometry; real space

• real, dimension(:,:), allocatable allglobal::trij

interface surface geometry; real space

real, dimension(:,:), allocatable allglobal::tzij

interface surface geometry; real space

real, dimension(:), allocatable allglobal::ivns

sine harmonics of vacuum normal magnetic field on interfaces; stellarator symmetric

 real, dimension(:), allocatable allglobal::ibns sine harmonics of plasma normal magnetic field on interfaces; stellarator symmetric • real, dimension(:), allocatable allglobal::ivnc cosine harmonics of vacuum normal magnetic field on interfaces; non-stellarator symmetric real, dimension(:), allocatable allglobal::ibnc cosine harmonics of plasma normal magnetic field on interfaces; non-stellarator symmetric • real, dimension(:), allocatable allglobal::Irbc local workspace • real, dimension(:), allocatable allglobal::lzbs local workspace • real, dimension(:), allocatable allglobal::Irbs local workspace • real, dimension(:), allocatable allglobal::lzbc local workspace • integer allglobal::num modes • integer, dimension(:), allocatable allglobal::mmrzrz • integer, dimension(:), allocatable allglobal::nnrzrz • real, dimension(:,:,:), allocatable allglobal::allrzrz • integer allglobal::nt discrete resolution along  $\theta$  of grid in real space • integer allglobal::nz discrete resolution along  $\zeta$  of grid in real space • integer allglobal::ntz discrete resolution; Ntz=Nt\*Nz shorthand integer allglobal::hnt discrete resolution; Ntz=Nt\*Nz shorthand · integer allglobal::hnz discrete resolution; Ntz=Nt\*Nz shorthand real aliglobal::sontz one / sqrt (one\*Ntz); shorthand real, dimension(:,:,:), allocatable allglobal::rij real-space grid; R • real, dimension(:,:,:), allocatable allglobal::zij real-space grid; Z • real, dimension(:,:,:), allocatable allglobal::xij what is this? • real, dimension(:,:,:), allocatable allglobal::yij what is this? • real, dimension(:,:), allocatable allglobal::sg real-space grid; jacobian and its derivatives • real, dimension(:,:,:,:), allocatable allglobal::guvij real-space grid; metric elements real, dimension(:,:,:), allocatable allglobal::gvuij real-space grid; metric elements (?); 10 Dec 15; real, dimension(:,:,:,:), allocatable allglobal::guvijsave what is this?

 integer, dimension(:,:), allocatable allglobal::ki identification of Fourier modes integer, dimension(:,:,:), allocatable allglobal::kijs identification of Fourier modes integer, dimension(:,:,:), allocatable allglobal::kija

identification of Fourier modes

• integer, dimension(:), allocatable allglobal::iotakkii

identification of Fourier modes

integer, dimension(:,:), allocatable allglobal::iotaksub

identification of Fourier modes

• integer, dimension(:,:), allocatable allglobal::iotakadd

identification of Fourier modes

integer, dimension(:,:), allocatable allglobal::iotaksgn

identification of Fourier modes

• real, dimension(:), allocatable allglobal::efmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::ofmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::cfmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::sfmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::evmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::odmn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::comn

Fourier harmonics; dummy workspace.

real, dimension(:), allocatable allglobal::simn

Fourier harmonics; dummy workspace.

• real, dimension(:), allocatable allglobal::ijreal

what is this?

• real, dimension(:), allocatable allglobal::ijimag

what is this?

• real, dimension(:), allocatable allglobal::jireal

what is this?

• real, dimension(:), allocatable allglobal::jiimag

what is this?

• real, dimension(:), allocatable allglobal::jkreal

what is this?

• real, dimension(:), allocatable allglobal::jkimag

what is this?

• real, dimension(:), allocatable allglobal::kjreal

what is this?

• real, dimension(:), allocatable allglobal::kjimag

what is this?

• real, dimension(:,:,:), allocatable allglobal::bsupumn

tangential field on interfaces;  $\theta$ -component; required for virtual casing construction of field; 11 Oct 12

real, dimension(:,:,:), allocatable allglobal::bsupvmn

tangential field on interfaces; ζ -component; required for virtual casing construction of field; 11 Oct 12

• real, dimension(:,:), allocatable allglobal::goomne

described in preset()

• real, dimension(:,:), allocatable allglobal::goomno

described in preset()

• real, dimension(:,:), allocatable allglobal::gssmne

described in preset()

- real, dimension(:,:), allocatable allglobal::gssmno described in preset()
- real, dimension(:,:), allocatable allglobal::gstmne described in preset()
- real, dimension(:,:), allocatable allglobal::gstmno described in preset()
- real, dimension(:,:), allocatable allglobal::gszmne described in preset()
- real, dimension(:,:), allocatable allglobal::gszmno described in preset()
- real, dimension(:,:), allocatable allglobal::gttmne described in preset()
- real, dimension(:,:), allocatable allglobal::gttmno described in preset()
- real, dimension(:,:), allocatable allglobal::gtzmne described in preset()
- real, dimension(:,:), allocatable allglobal::gtzmno described in preset()
- real, dimension(:,:), allocatable allglobal::gzzmne described in preset()
- real, dimension(:,:), allocatable allglobal::gzzmno described in preset()
- real, dimension(:,:,:,:), allocatable allglobal::dtoocc
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::dtoocs
   volume-integrated Chebychev-metrics: see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::dtoosc
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::dtooss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ttsscc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,::), allocatable allglobal::ttsscs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ttsssc
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ttssss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdstcc
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdstcs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdstsc
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdstss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdszcc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdszcs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdszsc

- volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::tdszss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddttcc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddttcs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddttsc
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddttss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddtzcc
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddtzcs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddtzsc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddtzss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddzzcc
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddzzcs
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddzzsc volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:,:,:), allocatable allglobal::ddzzss
   volume-integrated Chebychev-metrics; see matrix()
- real, dimension(:,:), allocatable allglobal::tsc what is this?
- real, dimension(:,:), allocatable aliglobal::tss what is this?
- real, dimension(:,:), allocatable allglobal::dtc what is this?
- real, dimension(:,:), allocatable allglobal::dts what is this?
- real, dimension(:,:), allocatable allglobal::dzc what is this?
- real, dimension(:,:), allocatable allglobal::dzs what is this?
- real, dimension(:,:), allocatable allglobal::ttc what is this?
- real, dimension(:,:), allocatable allglobal::tzc what is this?
- real, dimension(:,:), allocatable allglobal::tts what is this?
- real, dimension(:,:), allocatable allglobal::tzs what is this?
- real, dimension(:), allocatable allglobal::dtflux  $\delta\psi_{toroidal}$  in each annulus
- real, dimension(:), allocatable allglobal::dpflux  $\delta\psi_{poloidal} \ \emph{in each annulus}$

- real, dimension(:), allocatable allglobal::sweight
   minimum poloidal length constraint weight
- integer, dimension(:), allocatable allglobal::nadof

degrees of freedom in Beltrami fields in each annulus

• integer, dimension(:), allocatable allglobal::nfielddof

degrees of freedom in Beltrami fields in each annulus, field only, no Lagrange multipliers

• type(subgrid), dimension(:,:,:), allocatable allglobal::ate

magnetic vector potential cosine Fourier harmonics; stellarator-symmetric

• type(subgrid), dimension(:,:,:), allocatable allglobal::aze

magnetic vector potential cosine Fourier harmonics; stellarator-symmetric

type(subgrid), dimension(:,:,:), allocatable allglobal::ato

magnetic vector potential sine Fourier harmonics; non-stellarator-symmetric

type(subgrid), dimension(:,:,:), allocatable allglobal::azo

magnetic vector potential sine Fourier harmonics; non-stellarator-symmetric

• integer, dimension(:,:), allocatable allglobal::lma

Lagrange multipliers (?)

• integer, dimension(:,:), allocatable allglobal::Imb

Lagrange multipliers (?)

• integer, dimension(:,:), allocatable allglobal::Imc

Lagrange multipliers (?)

• integer, dimension(:,:), allocatable allglobal::Imd

Lagrange multipliers (?)

• integer, dimension(:,:), allocatable allglobal::Ime

Lagrange multipliers (?)

• integer, dimension(:,:), allocatable allglobal::Imf

Lagrange multipliers (?)

• integer, dimension(:,:), allocatable allglobal::Img

Lagrange multipliers (?)

• integer, dimension(:,:), allocatable allglobal::Imh

Lagrange multipliers (?)

• real, dimension(:,:), allocatable allglobal::Imavalue

what is this?

• real, dimension(:,:), allocatable allglobal::Imbvalue

what is this?

• real, dimension(:,:), allocatable allglobal::Imcvalue

what is this?

real, dimension(:,:), allocatable allglobal::Imdvalue

what is this?

• real, dimension(:,:), allocatable allglobal::Imevalue

what is this?

• real, dimension(:,:), allocatable allglobal::Imfvalue

what is this?

• real, dimension(:,:), allocatable allglobal::Imgvalue

what is this?

• real, dimension(:,:), allocatable allglobal::Imhvalue

what is this?

• integer, dimension(:,:), allocatable allglobal::fso

what is this?

integer, dimension(:,:), allocatable allglobal::fse

what is this?

logical allglobal::lcoordinatesingularity

set by LREGION macro; true if inside the innermost volume

logical allglobal::lplasmaregion

set by LREGION macro; true if inside the plasma region

logical allglobal::lvacuumregion

set by LREGION macro; true if inside the vacuum region

logical allglobal::lsavedguvij

flag used in matrix free

logical aliglobal::localconstraint

what is this?

• real, dimension(:,:), allocatable allglobal::dma

energy and helicity matrices; quadratic forms

• real, dimension(:,:), allocatable allglobal::dmb

energy and helicity matrices; quadratic forms

• real, dimension(:,:), allocatable allglobal::dmd

energy and helicity matrices; quadratic forms

• real, dimension(:), allocatable allglobal::dmas

sparse version of dMA, data

real, dimension(:), allocatable allglobal::dmds

sparse version of dMD, data

integer, dimension(:), allocatable allglobal::idmas

sparse version of dMA and dMD, indices

integer, dimension(:), allocatable allglobal::jdmas

sparse version of dMA and dMD, indices

integer, dimension(:), allocatable allglobal::ndmasmax

number of elements for sparse matrices

• integer, dimension(:), allocatable allglobal::ndmas

number of elements for sparse matrices

• real, dimension(:), allocatable allglobal::dmg

what is this?

real, dimension(:), allocatable allglobal::adotx

the matrix-vector product

real, dimension(:), allocatable allglobal::ddotx

the matrix-vector product

real, dimension(:,:), allocatable allglobal::solution

this is allocated in dforce; used in mp00ac and ma02aa; and is passed to packab

• real, dimension(:,:,:), allocatable allglobal::gmreslastsolution

used to store the last solution for restarting GMRES

• real, dimension(:), allocatable allglobal::mbpsi

matrix vector products

logical allglobal::liluprecond

whether to use ILU preconditioner for GMRES

• real, dimension(:,:), allocatable allglobal::beltramiinverse

Beltrami inverse matrix.

real, dimension(:,:,:), allocatable allglobal::diotadxup

measured rotational transform on inner/outer interfaces for each volume; d(transform)/dx; (see dforce)

• real, dimension(:,:,:), allocatable allglobal::ditgpdxtp

measured toroidal and poloidal current on inner/outer interfaces for each volume; d(Itor,Gpol)/dx; (see dforce)

• real, dimension(:,:,:,:), allocatable allglobal::glambda

save initial guesses for iterative calculation of rotational-transform

integer allglobal::lmns

what is this?

real, dimension(:,:,:), allocatable allglobal::bemn force vector; stellarator-symmetric (?)
 real, dimension(:,:), allocatable allglobal::iomn

```
force vector; stellarator-symmetric (?)

    real, dimension(:,:,:), allocatable allglobal::somn

      force vector; non-stellarator-symmetric (?)
• real, dimension(:,:,:), allocatable allglobal::pomn
      force vector; non-stellarator-symmetric (?)

    real, dimension(:,:,:), allocatable allglobal::bomn

      force vector; stellarator-symmetric (?)
• real, dimension(:,:), allocatable allglobal::iemn
      force vector; stellarator-symmetric (?)

    real, dimension(:,:,:), allocatable allglobal::semn

      force vector; non-stellarator-symmetric (?)

    real, dimension(:,:,:), allocatable allglobal::pemn

      force vector; non-stellarator-symmetric (?)

    real, dimension(:), allocatable allglobal::bbe

      force vector (?); stellarator-symmetric (?)

    real, dimension(:), allocatable allglobal::iio

      force vector (?); stellarator-symmetric (?)
• real, dimension(:), allocatable allglobal::bbo
      force vector (?); non-stellarator-symmetric (?)

    real, dimension(:), allocatable allglobal::iie

      force vector (?); non-stellarator-symmetric (?)

    real, dimension(:,:,:), allocatable allglobal::btemn

      covariant \theta cosine component of the tangential field on interfaces; stellarator-symmetric

    real, dimension(:,:,:), allocatable allglobal::bzemn

      covariant \zeta cosine component of the tangential field on interfaces; stellarator-symmetric

    real, dimension(:,:,:), allocatable allglobal::btomn

      covariant \theta sine component of the tangential field on interfaces; non-stellarator-symmetric
• real, dimension(:,:,:), allocatable allglobal::bzomn
      covariant \zeta sine component of the tangential field on interfaces; non-stellarator-symmetric
• real, dimension(:,:), allocatable allglobal::bloweremn
      covariant field for Hessian computation

    real, dimension(:,:), allocatable allglobal::bloweromn

      covariant field for Hessian computation

    integer allglobal::lgdof

      geometrical degrees of freedom associated with each interface

    integer allglobal::ngdof

      total geometrical degrees of freedom

    real, dimension(:,:,:), allocatable allglobal::dbbdrz

      derivative of magnetic field w.r.t. geometry (?)
• real, dimension(:,:), allocatable allglobal::diidrz
      derivative of spectral constraints w.r.t. geometry (?)
• real, dimension(:,:,:,:), allocatable allglobal::dffdrz
      derivatives of B^{\wedge}2 at the interfaces wrt geometry
• real, dimension(:,:,:,:), allocatable allglobal::dbbdmp
      derivatives of B<sup>\(\)</sup>2 at the interfaces wrt mu and dpflux

    real, dimension(:,:,:,::), allocatable allglobal::dmupfdx

      derivatives of mu and dpflux wrt geometry at constant interface transform

    logical allglobal::Ihessianallocated
```

flag to indicate that force gradient matrix is allocated (?)

• real, dimension(:,:), allocatable allglobal::hessian

force gradient matrix (?)

• real, dimension(:,:), allocatable allglobal::dessian

derivative of force gradient matrix (?)

• real, dimension(:,:), allocatable allglobal::cosi

some precomputed cosines

• real, dimension(:,:), allocatable allglobal::sini

some precomputed sines

• real, dimension(:), allocatable allglobal::gteta

something related to  $\sqrt{q}$  and  $\theta$ ?

• real, dimension(:), allocatable allglobal::gzeta

something related to  $\sqrt{g}$  and  $\zeta$ ?

• real, dimension(:), allocatable allglobal::ajk

definition of coordinate axis

• real, dimension(:,:,:), allocatable allglobal::dradr

derivatives of coordinate axis

real, dimension(:,:,:,:), allocatable allglobal::dradz

derivatives of coordinate axis

real, dimension(:,:,:,:), allocatable allglobal::dzadr

derivatives of coordinate axis

real, dimension(:,:,:,:), allocatable allglobal::dzadz

derivatives of coordinate axis

real, dimension(:,:,:), allocatable allglobal::drodr

derivatives of coordinate axis

• real, dimension(:,:,:), allocatable allglobal::drodz

derivatives of coordinate axis

• real, dimension(:,:,:), allocatable allglobal::dzodr

derivatives of coordinate axis

real, dimension(:,:,:), allocatable allglobal::dzodz

derivatives of coordinate axis

• integer, dimension(:,:), allocatable allglobal::djkp

for calculating cylindrical volume

• integer, dimension(:,:), allocatable allglobal::djkm

for calculating cylindrical volume

• real, dimension(:), allocatable allglobal::lbbintegral

B.B integral.

• real, dimension(:), allocatable allglobal::labintegral

A.B integral.

• real, dimension(:), allocatable allglobal::vvolume

volume integral of  $\sqrt{g}$ ; computed in volume

· real allglobal::dvolume

derivative of volume w.r.t. interface geometry

integer allglobal::ivol

labels volume; some subroutines (called by NAG) are fixed argument list but require the volume label

• real allglobal::gbzeta

toroidal (contravariant) field; calculated in bfield; required to convert  $\dot{\theta}$  to  $B^{\theta}$ ,  $\dot{s}$  to  $B^{s}$ 

• integer, dimension(:), allocatable allglobal::iquad

internal copy of Nquad

• real, dimension(:,:), allocatable allglobal::gaussianweight

weights for Gaussian quadrature

real, dimension(:,:), allocatable allglobal::gaussianabscissae

abscissae for Gaussian quadrature

· logical allglobal::lblinear

controls selection of Beltrami field solver; depends on LBeltrami

logical allglobal::lbnewton

controls selection of Beltrami field solver; depends on LBeltrami

logical allglobal::lbsequad

controls selection of Beltrami field solver; depends on LBeltrami

real, dimension(1:3) allglobal::orzp

used in mg00aa() to determine  $(s,\theta,\zeta)$  given  $(R,Z,\varphi)$ 

• type(derivative) allglobal::dbdx

 $d\mathbf{B}/d\mathbf{X}$  (?)

· integer allglobal::globaljk

labels position

real, dimension(:,:), allocatable allglobal::dxyz

computational boundary; position

real, dimension(:,:), allocatable allglobal::nxyz

computational boundary; normal

real, dimension(:,:), allocatable allglobal::jxyz

plasma boundary; surface current

• real, dimension(1:2) allglobal::tetazeta

what is this?

• real allglobal::virtualcasingfactor = -one / (four\*pi)

this agrees with diagno

integer allglobal::iberror

for computing error in magnetic field

• integer allglobal::nfreeboundaryiterations

number of free-boundary iterations already performed

• integer, parameter allglobal::node = 2

best to make this global for consistency between calling and called routines

• logical allglobal::first\_free\_bound = .false.

flag to indicate that this is the first free-boundary iteration

• type(c\_ptr) fftw\_interface::planf

FFTW-related (?)

• type(c\_ptr) fftw\_interface::planb

FFTW-related (?)

complex(c double complex), dimension(:,..;), allocatable fftw interface::cplxin

FFTW-related (?)

complex(c\_double\_complex), dimension(:,:,:), allocatable fftw\_interface::cplxout

FFTW-related (?)

## 10.12.1 Detailed Description

Defines input namelists and global variables, and opens some output files.

Note that all variables in namelist need to be broadcasted in readin.

### Input geometry

• The geometry of the l-th interface, for l=0,N where  $N\equiv$  Nvol, is described by a set of Fourier harmonics, using an arbitrary poloidal angle,

$$R_l(\theta,\zeta) = \sum_{j} R_{j,l} \cos(m_j \theta - n_j \zeta), \tag{259}$$

$$R_{l}(\theta,\zeta) = \sum_{j} R_{j,l} \cos(m_{j}\theta - n_{j}\zeta), \qquad (259)$$

$$Z_{l}(\theta,\zeta) = \sum_{j} Z_{j,l} \sin(m_{j}\theta - n_{j}\zeta). \qquad (260)$$

• These harmonics are read from the ext.sp file and come directly after the namelists described above. The required format is as follows:

- The coordinate axis corresponds to j=0 and the outermost boundary corresponds to j= Nvol.
- An arbitrary selection of harmonics may be inluded in any order, but only those within the range specified by Mpol and Ntor will be used.
- The geometry of all the interfaces, i.e. l=0,N, including the degenerate "coordinate-axis" interface, must be given.

#### 10.12.2 Data Type Documentation

**10.12.2.1 type typedefns::subgrid** used for quantities which have different resolutions in different volumes, e.g. the vector potential

#### **Class Members**

real, dimension(:), allocatable	s	coefficients
integer, dimension(:), allocatable	i	indices

### **Class Members**

re	al, dimension(:,:), allocatable	mat	
inte	ger, dimension(:), allocatable	ipivot	

## 10.12.2.2 type typedefns::matrixlu

### 10.12.2.3 type typedefns::derivative dB/dX (?)

#### **Class Members**

logical	I	what is this?
integer	vol	Used in coords(); required for global constraint force gradient evaluation.
integer	innout	what is this?
integer	ii	what is this?
integer	irz	what is this?
integer	issym	what is this?

# 10.13 src/hesian.f90 File Reference

Computes eigenvalues and eigenvectors of derivative matrix,  $\nabla_{\xi} \mathbf{F}$ .

## **Functions/Subroutines**

• subroutine hesian (NGdof, position, Mvol, mn, LGdof) Computes eigenvalues and eigenvectors of derivative matrix,  $\nabla_{\varepsilon} \mathbf{F}$ .

## 10.13.1 Detailed Description

Computes eigenvalues and eigenvectors of derivative matrix,  $\nabla_{\varepsilon} \mathbf{F}$ .

# 10.14 src/inputlist.f90 File Reference

Input namelists.

#### **Functions/Subroutines**

• subroutine inputlist::initialize\_inputs

#### **Variables**

```
• integer, parameter inputlist::mnvol = 256
```

The maximum value of Nvol is MNvol=256.

• integer, parameter inputlist::mmpol = 128

The maximum value of Mpol is MNpol=64.

• integer, parameter inputlist::mntor = 128

The maximum value of Ntor is MNtor=64.

• integer inputlist::igeometry = 3

selects Cartesian, cylindrical or toroidal geometry;

• integer inputlist::istellsym = 1

stellarator symmetry is enforced if Istellsym==1

• integer inputlist::Ifreebound = 0

compute vacuum field surrounding plasma

• real inputlist::phiedge = 1.0

total enclosed toroidal magnetic flux;

• real inputlist::curtor = 0.0

total enclosed (toroidal) plasma current;

• real inputlist::curpol = 0.0

total enclosed (poloidal) linking current;

• real inputlist::gamma = 0.0

adiabatic index; cannot set  $|\gamma| = 1$ 

• integer inputlist::nfp = 1

field periodicity

• integer inputlist::nvol = 1

number of volumes

integer inputlist::mpol = 0

number of poloidal Fourier harmonics

• integer inputlist::ntor = 0

number of toroidal Fourier harmonics

integer, dimension(1:mnvol+1) inputlist::lrad = 4

Chebyshev resolution in each volume.

• integer inputlist::lconstraint = -1

selects constraints; primarily used in ma02aa() and mp00ac().

• real, dimension(1:mnvol+1) inputlist::tflux = 0.0

toroidal flux,  $\psi_t$ , enclosed by each interface

• real, dimension(1:mnvol+1) inputlist::pflux = 0.0

poloidal flux,  $\psi_p$ , enclosed by each interface

• real, dimension(1:mnvol) inputlist::helicity = 0.0

helicity, K, in each volume,  $V_i$ 

• real inputlist::pscale = 0.0

```
pressure scale factor
• real, dimension(1:mnvol+1) inputlist::pressure = 0.0
      pressure in each volume
• integer inputlist::ladiabatic = 0
      logical flag
• real, dimension(1:mnvol+1) inputlist::adiabatic = 0.0
      adiabatic constants in each volume
real, dimension(1:mnvol+1) inputlist::mu = 0.0
      helicity-multiplier, \mu, in each volume
• real, dimension(1:mnvol+1) inputlist::ivolume = 0.0
       Toroidal current constraint normalized by \mu_0 ( I_{volume} = \mu_0 \cdot [A]), in each volume. This is a cumulative quantity:
      I_{\mathcal{V},i} = \int_0^{\psi_{t,i}} \mathbf{J} \cdot \mathbf{dS}. Physically, it represents the sum of all non-pressure driven currents.
• real, dimension(1:mnvol) inputlist::isurf = 0.0
       Toroidal current normalized by \mu_0 at each interface (cumulative). This is the sum of all pressure driven currents.

    integer, dimension(0:mnvol) inputlist::pl = 0

       "inside" interface rotational-transform is \iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.

    integer, dimension(0:mnvol) inputlist::ql = 0

       "inside" interface rotational-transform is \iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.

    integer, dimension(0:mnvol) inputlist::pr = 0

       "inside" interface rotational-transform is \iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.

    integer, dimension(0:mnvol) inputlist::qr = 0

       "inside" interface rotational-transform is \iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.
• real, dimension(0:mnvol) inputlist::iota = 0.0
      rotational-transform, t, on inner side of each interface

    integer, dimension(0:mnvol) inputlist::lp = 0

       "outer" interface rotational-transform is \,\iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.
• integer, dimension(0:mnvol) inputlist::lq = 0
       "outer" interface rotational-transform is \,\iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.

    integer, dimension(0:mnvol) inputlist::rp = 0

       "outer" interface rotational-transform is \iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.

    integer, dimension(0:mnvol) inputlist::rg = 0

       "outer" interface rotational-transform is \,\iota=(p_l+\gamma p_r)/(q_l+\gamma q_r), where \gamma is the golden mean, \gamma=(1+\sqrt{5})/2.

    real, dimension(0:mnvol) inputlist::oita = 0.0

      rotational-transform, ε, on outer side of each interface
real inputlist::mupftol = 1.0e-14
      accuracy to which \mu and \Delta\psi_p are required
integer inputlist::mupfits = 8
      an upper limit on the transform/helicity constraint iterations;
real inputlist::rpol = 1.0
      poloidal extent of slab (effective radius)
real inputlist::rtor = 1.0
      toroidal extent of slab (effective radius)
• integer inputlist::lreflect = 0
       =1 reflect the upper and lower bound in slab, =0 do not reflect
• real, dimension(0:mntor) inputlist::rac = 0.0
      stellarator symmetric coordinate axis;
• real, dimension(0:mntor) inputlist::zas = 0.0
      stellarator symmetric coordinate axis;
• real, dimension(0:mntor) inputlist::ras = 0.0
      non-stellarator symmetric coordinate axis;
• real, dimension(0:mntor) inputlist::zac = 0.0
```

non-stellarator symmetric coordinate axis;

- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::rbc = 0.0 stellarator symmetric boundary components;
- real, dimension(-mntor:mntor,-mmpol:mmpol) **inputlist::zbs** = 0.0 stellarator symmetric boundary components;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::rbs = 0.0
   non-stellarator symmetric boundary components;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::zbc = 0.0
   non-stellarator symmetric boundary components;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::rwc = 0.0 stellarator symmetric boundary components of wall;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::zws = 0.0 stellarator symmetric boundary components of wall;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::rws = 0.0
   non-stellarator symmetric boundary components of wall;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::zwc = 0.0
   non-stellarator symmetric boundary components of wall;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::vns = 0.0 stellarator symmetric normal field at boundary; vacuum component;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::bns = 0.0 stellarator symmetric normal field at boundary; plasma component;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::vnc = 0.0
   non-stellarator symmetric normal field at boundary; vacuum component;
- real, dimension(-mntor:mntor,-mmpol:mmpol) inputlist::bnc = 0.0
   non-stellarator symmetric normal field at boundary; plasma component;
- integer inputlist::linitialize = 0

Used to initialize geometry using a regularization / extrapolation method.

• integer inputlist::lautoinitbn = 1

Used to initialize  $B_{ns}$  using an initial fixed-boundary calculation.

• integer inputlist::lzerovac = 0

Used to adjust vacuum field to cancel plasma field on computational boundary.

• integer inputlist::ndiscrete = 2

resolution of the real space grid on which fast Fourier transforms are performed is given by Ndiscrete\*Mpol\*4

integer inputlist::nquad = -1

Resolution of the Gaussian quadrature.

• integer inputlist::impol = -4

Fourier resolution of straight-fieldline angle on interfaces.

• integer inputlist::intor = -4

Fourier resolution of straight-fieldline angle on interfaces;.

• integer inputlist::lsparse = 0

controls method used to solve for rotational-transform on interfaces

• integer inputlist::lsvdiota = 0

controls method used to solve for rotational-transform on interfaces; only relevant if Lsparse = 0

integer inputlist::imethod = 3

controls iterative solution to sparse matrix arising in real-space transformation to the straight-fieldline angle; only relevant if Lsparse.eq.2;

integer inputlist::iorder = 2

controls real-space grid resolution for constructing the straight-fieldline angle; only relevant if Lsparse>0

• integer inputlist::iprecon = 0

controls iterative solution to sparse matrix arising in real-space transformation to the straight-fieldline angle; only relevant if Lsparse.eq.2;

```
real inputlist::iotatol = -1.0
      tolerance required for iterative construction of straight-fieldline angle; only relevant if Lsparse.ge.2
• integer inputlist::lextrap = 0
      geometry of innermost interface is defined by extrapolation
integer inputlist::mregular = -1
      maximum regularization factor
• integer inputlist::lrzaxis = 1
      controls the guess of geometry axis in the innermost volume or initialization of interfaces
• integer inputlist::ntoraxis = 3
      the number of n harmonics used in the Jacobian m=1 harmonic elimination method; only relevant if \texttt{Lrzaxis.} \leftarrow
      ge.1.

    integer inputlist::lbeltrami = 4

      Control flag for solution of Beltrami equation.

    integer inputlist::linitgues = 1

      controls how initial guess for Beltrami field is constructed
• integer inputlist::lposdef = 0
      redundant;
• real inputlist::maxrndgues = 1.0
      the maximum random number of the Beltrami field if Linitgues = 3
• integer inputlist::Imatsolver = 3
      1 for LU factorization, 2 for GMRES, 3 for GMRES matrix-free
• integer inputlist::nitergmres = 200
      number of max iteration for GMRES
real inputlist::epsgmres = 1e-14
      the precision of GMRES
• integer inputlist::lgmresprec = 1
      type of preconditioner for GMRES, 1 for ILU sparse matrix
• real inputlist::epsilu = 1e-12
      the precision of incomplete LU factorization for preconditioning
• integer inputlist::lfindzero = 0
      use Newton methods to find zero of force-balance, which is computed by dforce()
• real inputlist::escale = 0.0
      controls the weight factor, BBweight, in the force-imbalance harmonics
• real inputlist::opsilon = 1.0
      weighting of force-imbalance

    real inputlist::pcondense = 2.0

      spectral condensation parameter

    real inputlist::epsilon = 0.0

      weighting of spectral-width constraint
• real inputlist::wpoloidal = 1.0
       star-like" poloidal angle constraint radial exponential factor used in preset() to construct <code>sweight</code>
• real inputlist::upsilon = 1.0
      weighting of "star-like" poloidal angle constraint used in preset() to construct sweight
• real inputlist::forcetol = 1.0e-10
      required tolerance in force-balance error; only used as an initial check
• real inputlist::c05xmax = 1.0e-06
      required tolerance in position, \mathbf{x} \equiv \{R_{i,v}, Z_{i,v}\}
real inputlist::c05xtol = 1.0e-12
      required tolerance in position, \mathbf{x} \equiv \{R_{i,v}, Z_{i,v}\}

    real inputlist::c05factor = 1.0e-02

      used to control initial step size in CO5NDF and CO5PDF
```

```
• logical inputlist::lreadgf = .true.
      \operatorname{read} \nabla_{\mathbf{x}} \mathbf{F} from file \operatorname{ext} . \operatorname{GF}
• integer inputlist::mfreeits = 0
      maximum allowed free-boundary iterations
• real inputlist::bnstol = 1.0e-06
      redundant;
• real inputlist::bnsblend = 0.666
      redundant;
• real inputlist::gbntol = 1.0e-06
      required tolerance in free-boundary iterations
• real inputlist::gbnbld = 0.666
      normal blend
real inputlist::vcasingeps = 1.e-12
      regularization of Biot-Savart; see bnorml(), casing()
• real inputlist::vcasingtol = 1.e-08
      accuracy on virtual casing integral; see bnorml(), casing()
• integer inputlist::vcasingits = 8
      minimum number of calls to adaptive virtual casing routine; see casing()
• integer inputlist::vcasingper = 1
      periods of integragion in adaptive virtual casing routine; see <a href="casing">casing</a>()
• integer inputlist::mcasingcal = 8
      minimum number of calls to adaptive virtual casing routine; see casing(); redundant;
• real inputlist::odetol = 1.0e-07
      o.d.e. integration tolerance for all field line tracing routines
• real inputlist::absreq = 1.0e-08
      redundant
• real inputlist::relreq = 1.0e-08
      redundant
• real inputlist::absacc = 1.0e-04
      redundant
• real inputlist::epsr = 1.0e-08
      redundant
• integer inputlist::nppts = 0
      number of toroidal transits used (per trajectory) in following field lines for constructing Poincaré plots; if nPpts<1,
      no Poincaré plot is constructed;
• real inputlist::ppts = 0.0
      stands for Poincare plot theta start. Chose at which angle (normalized over \pi) the Poincare field-line tracing start.

    integer, dimension(1:mnvol+1) inputlist::nptrj = -1

      number of trajectories in each annulus to be followed in constructing Poincaré plot
• logical inputlist::lhevalues = .false.
      to compute eigenvalues of 
abla \mathbf{F}
• logical inputlist::Ihevectors = .false.
      to compute eigenvectors (and also eigenvalues) of 
abla \mathbf{F}
• logical inputlist::lhmatrix = .false.
      to compute and write to file the elements of \nabla \mathbf{F}
• integer inputlist::lperturbed = 0
      to compute linear, perturbed equilibrium
integer inputlist::dpp = -1
      perturbed harmonic
integer inputlist::dqq = -1
      perturbed harmonic
```

• integer inputlist::lerrortype = 0

the type of error output for Lcheck=1

• integer inputlist::ngrid = -1

the number of points to output in the grid, -1 for Lrad(vvol)

• real inputlist::drz = 1E-5

difference in geometry for finite difference estimate (debug only)

• integer inputlist::lcheck = 0

implement various checks

• logical inputlist::Itiming = .false.

to check timing

• real inputlist::fudge = 1.0e-00

redundant

• real inputlist::scaling = 1.0e-00

redundant

- logical inputlist::wmanual = .false.
- logical inputlist::wrzaxis = .false.
- logical inputlist::wpackxi = .false.
- logical inputlist::wvolume = .false.
- logical inputlist::wcoords = .false.
- logical inputlist::wbasefn = .false.
- logical inputlist::wmemory = .false.
- logical inputlist::wmetrix = .false.
- logical inputlist::wma00aa = .false.
- logical inputlist::wmatrix = .false.
- logical inputlist::wspsmat = .false.
- logical inputlist::wspsint = .false.
- logical inputlist::wmp00ac = .false.
- logical inputlist::wma02aa = .false.
- logical inputlist::wpackab = .false.
- logical inputlist::wtr00ab = .false.
- logical inputlist::wcurent = .false.
- logical inputlist::wdf00ab = .false.
- logical inputlist::wlforce = .false.
- logical inputlist::wintghs = .false.
- logical inputlist::wmtrxhs = .false.
- logical inputlist::wlbpol = .false.
- logical inputlist::wbrcast = .false.
- logical inputlist::wdfp100 = .false.
- logical inputlist::wdfp200 = .false.
- logical inputlist::wdforce = .false.
- logical inputlist::wnewton = .false.
- logical inputlist::wcasing = .false.
- logical inputlist::wbnorml = .false.
- logical inputlist::wjo00aa = .false.
- logical inputlist::wpp00aa = .false.
- logical inputlist::wpp00ab = .false.
- logical inputlist::wbfield = .false.
- logical inputlist::wstzxyz = .false.
- logical inputlist::whesian = .false.
- logical inputlist::wra00aa = .false.
- logical inputlist::wnumrec = .false.
- logical inputlist::wdcuhre = .false.
- logical inputlist::wminpack = .false.

- logical inputlist::wiqpack = .false.
- logical inputlist::wrksuite = .false.
- logical inputlist::wi1mach = .false.
- logical inputlist::wd1mach = .false.
- logical inputlist::wilut = .false.
- logical inputlist::witers = .false.
- logical inputlist::wsphdf5 = .false.
- logical inputlist::wpreset = .false.
- logical inputlist::wglobal = .false.
- logical inputlist::wxspech = .false.
- logical inputlist::wbuild vector potential = .false.
- logical inputlist::wreadin = .false.

write screen output of readin()

• logical inputlist::wwrtend = .false.

write screen output of wrtend()

• logical inputlist::wmacros = .false.

write screen output from expanded macros

### 10.14.1 Detailed Description

Input namelists.

# 10.15 src/intghs.f90 File Reference

Calculates volume integrals of Chebyshev-polynomials and covariant field for Hessian computation.

### **Data Types**

· type intghs\_module::intghs\_workspace

This calculates the integral of something related to matrix-vector-multiplication.

### **Functions/Subroutines**

• subroutine intghs (Iquad, mn, Ivol, Irad, idx)

Calculates volume integrals of Chebyshev polynomials and covariant field products.

· subroutine intghs workspace init (Ivol)

init workspace

subroutine intghs\_workspace\_destroy ()

free workspace

### **Variables**

• type(intghs\_workspace) intghs\_module::wk

This is an instance of the intghs\_workspace type.

### 10.15.1 Detailed Description

Calculates volume integrals of Chebyshev-polynomials and covariant field for Hessian computation.

## 10.15.2 Function/Subroutine Documentation

Calculates volume integrals of Chebyshev polynomials and covariant field products.

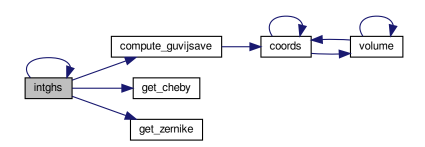
#### **Parameters**

lquad	
mn	
Ivol	
Irad	
idx	

References allglobal::ate, allglobal::ato, allglobal::aze, allglobal::aze, compute\_guvijsave(), allglobal::cpus, allglobal::dtc, allglobal::dts, allglobal::dzc, allglobal::dzs, allglobal::gaussianabscissae, allglobal::gaussianweight, get\_cheby(), get\_zernike(), allglobal::guvij, allglobal::guvijsave, constants::half, allglobal::im, allglobal::in, intghs(), allglobal::lcoordinatesingularity, allglobal::lsavedguvij, allglobal::mne, allglobal::mpi\_comm\_spec, inputlist::mpol, allglobal::myid, allglobal::ncpu, allglobal::notstellsym, allglobal::nt, allglobal::nt, allglobal::nz, constants::one, fileunits::ounit, constants::pi, constants::pi2, allglobal::g, numerical::small, numerical::sqrtmachprec, allglobal::tsc, allglobal::tts, allglobal::tts, allglobal::tts, allglobal::tts, allglobal::tts, allglobal::tts, numerical::vsmall, inputlist::wmacros, allglobal::yesstellsym, and constants::zero.

Referenced by get\_perturbed\_solution(), intghs(), intghs\_workspace\_destroy(), intghs\_workspace\_init(), matvec(), and mp00ac().

Here is the call graph for this function:



Here is the caller graph for this function:



# 

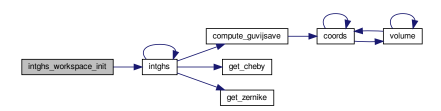
init workspace

### **Parameters**

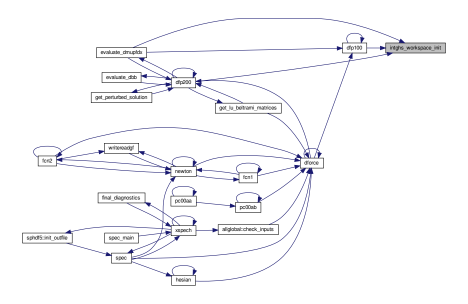
Ivol

References allglobal::cpus, intghs(), allglobal::iquad, inputlist::lrad, allglobal::mn, allglobal::mpi\_comm\_spec, inputlist::mpol, allglobal::myid, allglobal::ncpu, allglobal::ntz, fileunits::ounit, inputlist::wmacros, and constants::zero. Referenced by dfp100(), dfp200(), and evaluate\_dmupfdx().

Here is the call graph for this function:



Here is the caller graph for this function:



**10.15.2.3 intghs\_workspace\_destroy()** subroutine intghs\_workspace\_destroy free workspace

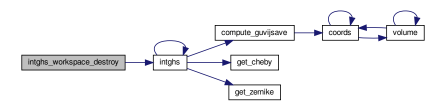
#### **Parameters**

Ivol

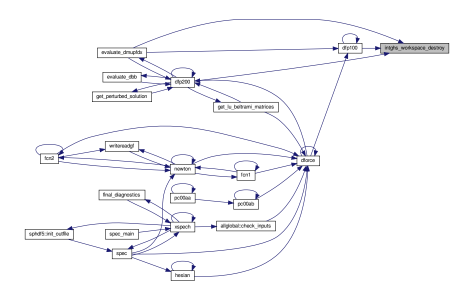
References allglobal::cpus, intghs(), allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ncpu, fileunits::ounit, and inputlist::wmacros.

Referenced by dfp100(), dfp200(), and evaluate\_dmupfdx().

Here is the call graph for this function:



Here is the caller graph for this function:



# 10.16 src/jo00aa.f90 File Reference

Measures error in Beltrami equation,  $\nabla \times \mathbf{B} - \mu \mathbf{B}$ .

## **Functions/Subroutines**

• subroutine jo00aa (Ivol, Ntz, Iquad, mn) Measures error in Beltrami equation,  $\nabla \times \mathbf{B} - \mu \mathbf{B}$ .

## 10.16.1 Detailed Description

Measures error in Beltrami equation,  $\nabla \times \mathbf{B} - \mu \mathbf{B}$ .

# 10.17 src/lforce.f90 File Reference

Computes  $B^2$ , and the spectral condensation constraints if required, on the interfaces,  $\mathcal{I}_i$ .

#### **Functions/Subroutines**

subroutine Iforce (Ivol, iocons, ideriv, Ntz, dBB, XX, YY, length, DDI, MMI, iflag)
 Computes B<sup>2</sup>, and the spectral condensation constraints if required, on the interfaces, \( \mathcal{I}\_i \).

#### 10.17.1 Detailed Description

Computes  $B^2$ , and the spectral condensation constraints if required, on the interfaces,  $\mathcal{I}_i$ .

#### 10.18 src/ma00aa.f90 File Reference

Calculates volume integrals of Chebyshev polynomials and metric element products.

#### **Functions/Subroutines**

subroutine ma00aa (Iquad, mn, Ivol, Irad)
 Calculates volume integrals of Chebyshev polynomials and metric element products.

#### 10.18.1 Detailed Description

Calculates volume integrals of Chebyshev polynomials and metric element products.

#### 10.19 src/ma02aa.f90 File Reference

Constructs Beltrami field in given volume consistent with flux, helicity, rotational-transform and/or parallel-current constraints.

#### **Functions/Subroutines**

• subroutine ma02aa (Ivol, NN)

Constructs Beltrami field in given volume consistent with flux, helicity, rotational-transform and/or parallel-current constraints.

## 10.19.1 Detailed Description

Constructs Beltrami field in given volume consistent with flux, helicity, rotational-transform and/or parallel-current constraints.

## 10.20 src/matrix.f90 File Reference

Constructs energy and helicity matrices that represent the Beltrami linear system.

#### **Functions/Subroutines**

• subroutine matrix (Ivol, mn, Irad)

Constructs energy and helicity matrices that represent the Beltrami linear system. gauge conditions

subroutine matrixbg (Ivol, mn, Irad)

#### 10.20.1 Detailed Description

Constructs energy and helicity matrices that represent the Beltrami linear system.

## 10.21 src/memory.f90 File Reference

memory management module

#### **Functions/Subroutines**

- subroutine allocate\_beltrami\_matrices (vvol, LcomputeDerivatives) allocate Beltrami matrices
- subroutine deallocate\_beltrami\_matrices (LcomputeDerivatives)
- subroutine allocate\_geometry\_matrices (vvol, LcomputeDerivatives)
  - allocate geometry matrices

deallocate Beltrami matrices

• subroutine deallocate\_geometry\_matrices (LcomputeDerivatives) deallocate geometry matrices

#### 10.21.1 Detailed Description

memory management module

#### 10.21.2 Function/Subroutine Documentation

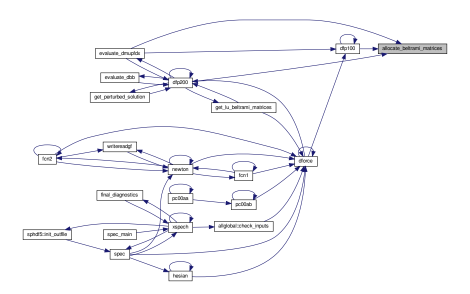
#### anocate Dentami matrici

**Parameters** 

vvol	
LcomputeDerivatives	

References allglobal::adotx, allglobal::ddotx, allglobal::dma, allglobal::dmas, allglobal::dmb, allglobal::dmb

Here is the caller graph for this function:



## 10.21.2.2 deallocate\_beltrami\_matrices() subroutine deallocate\_beltrami\_matrices (

logical, intent(in) LcomputeDerivatives )

deallocate Beltrami matrices

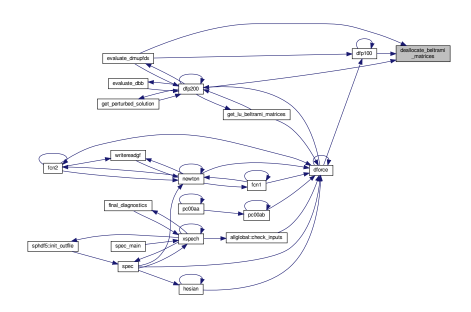
#### **Parameters**

LcomputeDerivatives

References allglobal::adotx, allglobal::dma, allglobal::dmas, allglobal::dmb, allglobal::dmb, allglobal::dmb, allglobal::dmb, allglobal::dmb, allglobal::dmb, allglobal::dmbpsi, allglobal::notmatrixfree, allglobal::solution, and inputlist::wmacros.

Referenced by dfp100(), dfp200(), and evaluate\_dmupfdx().

Here is the caller graph for this function:



## $\textbf{10.21.2.3} \quad \textbf{allocate\_geometry\_matrices()} \quad \texttt{subroutine allocate\_geometry\_matrices} \ \ ($

integer *vvol*,

logical, intent(in) LcomputeDerivatives )

allocate geometry matrices

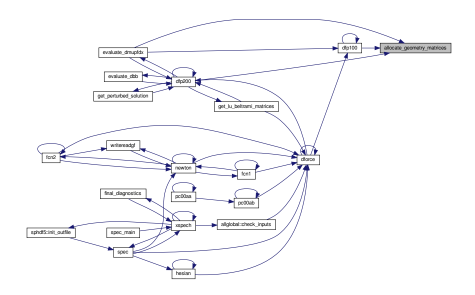
#### **Parameters**

vvol	
LcomputeDerivatives	

References allglobal::ddttcc, allglobal::ddttcs, allglobal::ddttsc, allglobal::dtsc, allglobal::dtsc, allglobal::dtsc, allglobal::dtsc, allglobal::dtsc, allglobal::dtsc, allglobal::mn, inputlist::mpol, allglobal::notstellsym, allglobal::ntt, allglobal::tdstcc, allglobal::tdstcs, allglobal::tdstsc, allglobal::tdstsc, allglobal::tdstsc, allglobal::tsc, allglobal::tsc, allglobal::tsc, allglobal::tsc, allglobal::tsc, allglobal::tsc, allglobal::ttsc, allgloba

Referenced by dfp100(), dfp200(), and evaluate\_dmupfdx().

Here is the caller graph for this function:



# 10.21.2.4 deallocate\_geometry\_matrices() subroutine deallocate\_geometry\_matrices ( logical, intent(in) LcomputeDerivatives )

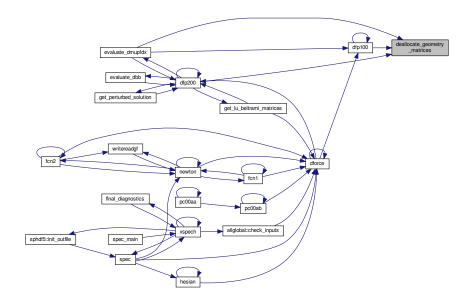
deallocate geometry matrices

## **Parameters**

LcomputeDerivatives

Referenced by dfp100(), dfp200(), and evaluate\_dmupfdx().

Here is the caller graph for this function:



## 10.22 src/metrix.f90 File Reference

Calculates the metric quantities,  $\sqrt{g}\,g^{\mu\nu}$ , which are required for the energy and helicity integrals.

## **Functions/Subroutines**

- subroutine metrix (Iquad, IvoI) Calculates the metric quantities,  $\sqrt{g}\,g^{\mu\nu}$ , which are required for the energy and helicity integrals.
- subroutine compute\_guvijsave (Iquad, vvol, ideriv, Lcurvature) compute guvijsave

## 10.22.1 Detailed Description

Calculates the metric quantities,  $\sqrt{g}\,g^{\mu\nu}$ , which are required for the energy and helicity integrals.

#### 10.22.2 Function/Subroutine Documentation

## **Parameters**

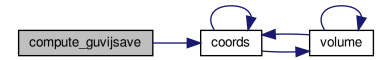
lquad	
vvol	
ideriv	
Lcurvature	

References coords(), allglobal::gaussianabscissae, allglobal::guvij, allglobal::guvijsave, allglobal::mn, allglobal::ntz,

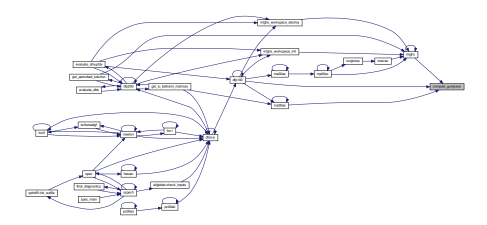
and allglobal::sg.

Referenced by dfp100(), intghs(), and ma00aa().

Here is the call graph for this function:



Here is the caller graph for this function:



## 10.23 src/mp00ac.f90 File Reference

Solves Beltrami/vacuum (linear) system, given matrices.

## **Functions/Subroutines**

- subroutine mp00ac (Ndof, Xdof, Fdof, Ddof, Ldfjac, iflag)
   Solves Beltrami/vacuum (linear) system, given matrices.
   unpacking fluxes, helicity multiplier
- subroutine rungmres (n, nrestart, mu, vvol, rhs, sol, ipar, fpar, wk, nw, guess, a, au, jau, ju, iperm, ierr) run GMRES
- subroutine matvec (n, x, ax, a, mu, vvol) compute a.x by either by coumputing it directly, or using a matrix free method
- subroutine prec\_solve (n, vecin, vecout, au, jau, ju, iperm)
   apply the preconditioner

## 10.23.1 Detailed Description

Solves Beltrami/vacuum (linear) system, given matrices.

## 10.23.2 Function/Subroutine Documentation

#### 10.23.2.1 rungmres() subroutine rungmres (

```
integer n,
integer nrestart,
real mu,
integer vvol,
real, dimension(1:n) rhs,
real, dimension(1:n) sol,
integer, dimension(16) ipar,
real, dimension(16) fpar,
real, dimension(1:nw) wk,
integer nw,
real, dimension(n) guess,
real, dimension(*) a,
real, dimension(*) au,
integer, dimension(*) jau,
integer, dimension(*) ju,
integer, dimension(*) iperm,
integer ierr )
```

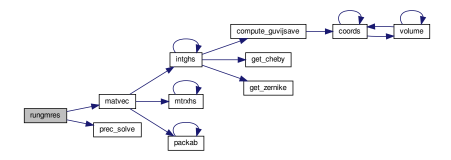
## run GMRES

#### **Parameters**

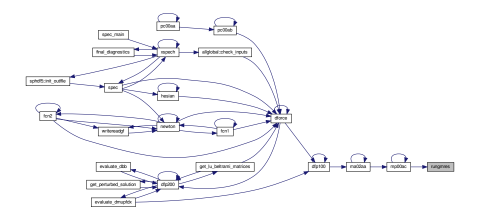
References inputlist::epsgmres, allglobal::liluprecond, matvec(), inputlist::nitergmres, constants::one, prec\_solve(), and constants::zero.

Referenced by mp00ac().

Here is the call graph for this function:



Here is the caller graph for this function:



compute a.x by either by coumputing it directly, or using a matrix free method

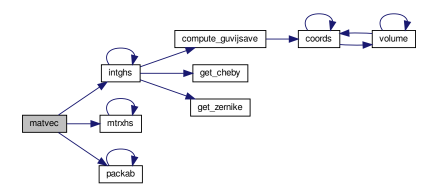
#### **Parameters**

n	
X	
ax	
а	
mu	
vvol	

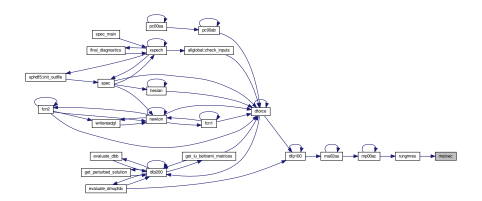
References allglobal::dmd, intghs(), allglobal::iquad, inputlist::lrad, allglobal::mn, mtrxhs(), allglobal::notmatrixfree, constants::one, packab(), and constants::zero.

Referenced by rungmres().

Here is the call graph for this function:



Here is the caller graph for this function:

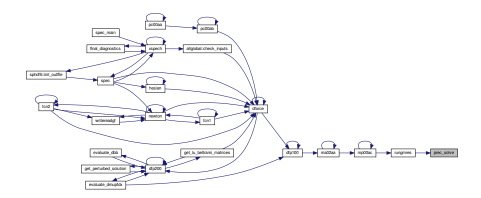


## Parameters

n	
vecin	
vecout	
au	
jau	
ju	
iperm	

Referenced by rungmres().

Here is the caller graph for this function:



#### 10.24 src/mtrxhs.f90 File Reference

Constructs matrices that represent the Beltrami linear system, matrix-free.

## **Functions/Subroutines**

subroutine mtrxhs (Ivol, mn, Irad, resultA, resultD, idx)
 Constructs matrices that represent the Beltrami linear system, matrix-free.

#### 10.24.1 Detailed Description

Constructs matrices that represent the Beltrami linear system, matrix-free.

#### 10.25 src/newton.f90 File Reference

Employs Newton method to find  $\mathbf{F}(\mathbf{x}) = 0$ , where  $\mathbf{x} \equiv \{\text{geometry}\}\$ and  $\mathbf{F}$  is defined in dforce() .

#### Modules

• module newtontime timing of Newton iterations

## **Functions/Subroutines**

• subroutine newton (NGdof, position, ihybrd)

Employs Newton method to find F(x) = 0, where  $x \equiv \{\text{geometry}\}\$ and F is defined in dforce().

• subroutine writereadgf (readorwrite, NGdof, ireadhessian)

read or write force-derivative matrix

• subroutine fcn1 (NGdof, xx, fvec, irevcm)

fcn

subroutine fcn2 (NGdof, xx, fvec, fjac, Ldfjac, irevcm)

fcn2

## Variables

• integer newtontime::nfcalls

number of calls to get function values (?)

• integer newtontime::ndcalls

number of calls to get derivative values (?)

## · real newtontime::lastcpu

last CPU that called this (?)

#### 10.25.1 Detailed Description

Employs Newton method to find F(x) = 0, where  $x \equiv \{\text{geometry}\}\$  and F is defined in dforce().

## 10.26 src/packab.f90 File Reference

Packs, and unpacks, Beltrami field solution vector;  $\mathbf{a} \equiv \{A_{\theta,e,i,l}, A_{\zeta,e,i,l}, \text{etc.}\}.$ 

## **Functions/Subroutines**

subroutine packab (packorunpack, Ivol, NN, solution, ideriv)
 Packs and unpacks Beltrami field solution vector.

#### 10.26.1 Detailed Description

Packs, and unpacks, Beltrami field solution vector;  $\mathbf{a} \equiv \{A_{\theta,e,i,l}, A_{\zeta,e,i,l}, \text{etc.}\}.$ 

## 10.27 src/packxi.f90 File Reference

Packs, and unpacks, geometrical degrees of freedom; and sets coordinate axis.

#### **Functions/Subroutines**

 subroutine packxi (NGdof, position, Mvol, mn, iRbc, iZbs, iRbs, iZbc, packorunpack, LComputeDerivatives, LComputeAxis)

Packs, and unpacks, geometrical degrees of freedom; and sets coordinate axis.

#### 10.27.1 Detailed Description

Packs, and unpacks, geometrical degrees of freedom; and sets coordinate axis.

#### 10.28 src/pc00aa.f90 File Reference

Use preconditioned conjugate gradient method to find minimum of energy functional.

#### **Functions/Subroutines**

subroutine pc00aa (NGdof, position, Nvol, mn, ie04dgf)
 Use preconditioned conjugate gradient method to find minimum of energy functional.

## 10.28.1 Detailed Description

Use preconditioned conjugate gradient method to find minimum of energy functional.

#### 10.29 src/pc00ab.f90 File Reference

Returns the energy functional and it's derivatives with respect to geometry.

#### **Functions/Subroutines**

• subroutine pc00ab (mode, NGdof, Position, Energy, Gradient, nstate, iuser, ruser)

Returns the energy functional and it's derivatives with respect to geometry.

## 10.29.1 Detailed Description

Returns the energy functional and it's derivatives with respect to geometry.

## 10.30 src/pp00aa.f90 File Reference

Constructs Poincaré plot and "approximate" rotational-transform (driver).

#### **Functions/Subroutines**

• subroutine pp00aa

Constructs Poincaré plot and "approximate" rotational-transform (driver).

#### 10.30.1 Detailed Description

Constructs Poincaré plot and "approximate" rotational-transform (driver).

## 10.31 src/pp00ab.f90 File Reference

Follows magnetic fieldline using ode-integration routine from rksuite.f.

## **Functions/Subroutines**

• subroutine pp00ab (Ivol, sti, Nz, nPpts, poincaredata, fittedtransform, utflag)

Constructs Poincaré plot and "approximate" rotational-transform (for single field line).

## 10.31.1 Detailed Description

Follows magnetic fieldline using ode-integration routine from rksuite.f .

## 10.32 src/preset.f90 File Reference

Allocates and initializes internal arrays.

#### **Functions/Subroutines**

· subroutine preset

Allocates and initializes internal arrays.

#### 10.32.1 Detailed Description

Allocates and initializes internal arrays.

#### 10.33 src/ra00aa.f90 File Reference

Writes vector potential to .ext.sp.A .

#### **Functions/Subroutines**

• subroutine ra00aa (writeorread)

Writes vector potential to .ext.sp.A .

## 10.33.1 Detailed Description

Writes vector potential to .ext.sp.A .

#### 10.34 src/rzaxis.f90 File Reference

The coordinate axis is assigned via a poloidal average over an arbitrary surface.

## **Functions/Subroutines**

• subroutine rzaxis (Mvol, mn, inRbc, inZbs, inRbs, inZbc, ivol, LcomputeDerivatives)

The coordinate axis is assigned via a poloidal average over an arbitrary surface.

• subroutine fndiff\_rzaxis (Mvol, mn, ivol, jRbc, jRbs, jZbc, JZbs, imn, irz, issym)

## 10.34.1 Detailed Description

The coordinate axis is assigned via a poloidal average over an arbitrary surface.

## 10.35 src/sphdf5.f90 File Reference

Writes all the output information to ext.sp.h5.

#### **Modules**

module sphdf5

writing the HDF5 output file

#### **Functions/Subroutines**

· subroutine sphdf5::init\_outfile

Initialize the interface to the HDF5 library and open the output file.

• subroutine sphdf5::mirror\_input\_to\_outfile

Mirror input variables into output file.

• subroutine sphdf5::init\_convergence\_output

Prepare convergence evolution output.

• subroutine sphdf5::write\_convergence\_output (nDcalls, ForceErr)

Write convergence output (evolution of interface geometry, force, etc).

• subroutine sphdf5::write\_grid

Write the magnetic field on a grid.

• subroutine sphdf5::init\_flt\_output (numTrajTotal)

Initialize field line tracing output group and create array datasets.

subroutine sphdf5::write\_poincare (offset, data, success)

Write a hyperslab of Poincare data corresponding to the output of one parallel worker.

• subroutine sphdf5::write\_transform (offset, length, lvol, diotadxup, fiota)

Write the rotational transform output from field line following.

• subroutine sphdf5::finalize\_flt\_output

Finalize Poincare output.

• subroutine sphdf5::write vector potential (sumLrad, allAte, allAze, allAto, allAzo)

Write the magnetic vector potential Fourier harmonics to the output file group /vector\_potential.

subroutine sphdf5::hdfint

Write the final state of the equilibrium to the output file.

subroutine sphdf5::finish\_outfile

Close all open HDF5 objects (we know of) and list any remaining still-open objects.

#### **Variables**

• logical, parameter **sphdf5::hdfdebug** = .false.

global flag to enable verbal diarrhea commenting HDF5 operations

• integer, parameter **sphdf5::internalhdf5msg** = 0

1: print internal HDF5 error messages; 0: only error messages from sphdf5

• integer sphdf5::hdfier

error flag for HDF5 library

integer sphdf5::rank

rank of data to write using macros

integer(hid t) sphdf5::file\_id

default file ID used in macros

integer(hid\_t) sphdf5::space\_id

default dataspace ID used in macros

• integer(hid t) sphdf5::dset id

default dataset ID used in macros

integer(hsize t), dimension(1:1) sphdf5::onedims

dimension specifier for one-dimensional data used in macros

integer(hsize t), dimension(1:2) sphdf5::twodims

dimension specifier for two-dimensional data used in macros

integer(hsize\_t), dimension(1:3) sphdf5::threedims

dimension specifier for three-dimensional data used in macros

logical sphdf5::grp\_exists

flags used to signal if a group already exists

logical sphdf5::var\_exists

flags used to signal if a variable already exists

· integer(hid t) sphdf5::iteration\_dset\_id

Dataset identifier for "iteration".

integer(hid\_t) sphdf5::dataspace

dataspace for extension by 1 iteration object

integer(hid t) sphdf5::memspace

memspace for extension by 1 iteration object

integer(hsize\_t), dimension(1) sphdf5::old\_data\_dims

current dimensions of "iterations" dataset

integer(hsize\_t), dimension(1) sphdf5::data\_dims

new dimensions for "iterations" dataset

• integer(hsize\_t), dimension(1) sphdf5::max\_dims

maximum dimensions for "iterations" dataset

integer(hid\_t) sphdf5::plist\_id

Property list identifier used to activate dataset transfer property.

integer(hid\_t) sphdf5::dt\_ndcalls\_id

Memory datatype identifier (for "nDcalls" dataset in "/grid")

integer(hid\_t) sphdf5::dt\_energy\_id

Memory datatype identifier (for "Energy" dataset in "/grid")

integer(hid\_t) sphdf5::dt\_forceerr\_id

Memory datatype identifier (for "ForceErr" dataset in "/grid")

integer(hid\_t) sphdf5::dt\_irbc\_id

Memory datatype identifier (for "iRbc" dataset in "/grid")

• integer(hid t) sphdf5::dt izbs id

Memory datatype identifier (for "iZbs" dataset in "/grid")

integer(hid\_t) sphdf5::dt\_irbs\_id

Memory datatype identifier (for "iRbs" dataset in "/grid")

• integer(hid t) sphdf5::dt izbc id

Memory datatype identifier (for "iZbc" dataset in "/grid")

integer, parameter sphdf5::rankp =3

rank of Poincare data

• integer, parameter sphdf5::rankt =2

rank of rotational transform data

integer(hid\_t) sphdf5::grppoincare

group for Poincare data

integer(hid\_t) sphdf5::dset\_id\_t

Dataset identifier for  $\theta$  coordinate of field line following.

integer(hid t) sphdf5::dset id s

Dataset identifier for s coordinate of field line following.

integer(hid t) sphdf5::dset\_id\_r

Dataset identifier for R coordinate of field line following.

integer(hid\_t) sphdf5::dset\_id\_z

Dataset identifier for Z coordinate of field line following.

integer(hid\_t) sphdf5::dset\_id\_success

Dataset identifier for success flag of trajectories to follow.

integer(hid\_t) sphdf5::filespace\_t

Dataspace identifier in file for  $\theta$  coordinate of field line following.

• integer(hid\_t) sphdf5::filespace\_s

Dataspace identifier in file for s coordinate of field line following.

integer(hid t) sphdf5::filespace\_r

Dataspace identifier in file for R coordinate of field line following.

• integer(hid\_t) sphdf5::filespace\_z

Dataspace identifier in file for Z coordinate of field line following.

integer(hid t) sphdf5::filespace\_success

Dataspace identifier in file for success flag of trajectories to follow.

integer(hid\_t) sphdf5::memspace\_t

Dataspace identifier in memory for  $\theta$  coordinate of field line following.

integer(hid\_t) sphdf5::memspace\_s

Dataspace identifier in memory for s coordinate of field line following.

integer(hid\_t) sphdf5::memspace\_r

Dataspace identifier in memory for  ${\cal R}$  coordinate of field line following.

integer(hid\_t) sphdf5::memspace\_z

Dataspace identifier in memory for  ${\cal Z}$  coordinate of field line following.

• integer(hid\_t) sphdf5::memspace\_success

Dataspace identifier in memory for success flag of trajectories to follow.

integer(hid t) sphdf5::grptransform

group for rotational transform data

integer(hid\_t) sphdf5::dset\_id\_diotadxup

Dataset identifier for diotadxup (derivative of rotational transform ?)

• integer(hid\_t) sphdf5::dset\_id\_fiota

Dataset identifier for fiota ( rotational transform ?)

integer(hid\_t) sphdf5::filespace\_diotadxup

Dataspace identifier in file for diotadxup.

integer(hid\_t) sphdf5::filespace\_fiota

Dataspace identifier in file for fiota.

• integer(hid t) sphdf5::memspace diotadxup

Dataspace identifier in memory for diotadxup.

integer(hid\_t) sphdf5::memspace\_fiota

Dataspace identifier in memory for fiota.

• character(len=15), parameter **sphdf5::aname** = "description"

Attribute name for descriptive info.

integer(hid\_t) sphdf5::attr\_id

Attribute identifier.

integer(hid\_t) sphdf5::aspace\_id

Attribute Dataspace identifier.

integer(hid\_t) sphdf5::atype\_id

Attribute Datatype identifier.

• integer, parameter sphdf5::arank = 1

Attribure rank.

integer(hsize\_t), dimension(arank) sphdf5::adims = (/1/)

Attribute dimension.

integer(size\_t) sphdf5::attrlen

Length of the attribute string.

character(len=:), allocatable sphdf5::attr\_data

Attribute data.

#### 10.35.1 Detailed Description

Writes all the output information to ext.sp.h5.

If the output file already exists, it will be deleted and replaced by an empty one, which gets filled in with the updated data. All calls to the HDF5 API are filtered to only happen from MPI rank-0 to be able to use the serial HDF5 library. Parallel HDF5 was considered in the past, but abandoned due to very subtle and irreproducible errors.

## 10.36 src/spsint.f90 File Reference

Calculates volume integrals of Chebyshev-polynomials and metric elements for preconditioner.

#### **Functions/Subroutines**

subroutine spsint (Iquad, mn, Ivol, Irad)
 Calculates volume integrals of Chebyshev-polynomials and metric elements for preconditioner.

#### 10.36.1 Detailed Description

Calculates volume integrals of Chebyshev-polynomials and metric elements for preconditioner.

#### 10.37 src/spsmat.f90 File Reference

Constructs matrices for the precondtioner.

#### **Functions/Subroutines**

• subroutine spsmat (Ivol, mn, Irad)

Constructs matrices for the precondtioner.

• subroutine push back (iq, nq, NN, vA, vD, vjA, qA, qD, qjA)

push a new element at the back of the queue

subroutine clean\_queue (nq, NN, qA, qD, qjA)

clean the queue

• subroutine addline (nq, NN, qA, qD, qjA, ns, nrow, dMAS, dMDS, jdMAS, idMAS)

add the content from the queue to the real matrices

#### 10.37.1 Detailed Description

Constructs matrices for the precondtioner.

#### 10.37.2 Function/Subroutine Documentation

```
10.37.2.1 push_back() subroutine push_back (
             integer, intent(in) iq,
             integer, dimension(4), intent(inout) nq,
             integer, intent(in) NN,
             real, intent(in) vA,
             real, intent(in) vD,
             integer, intent(in) vjA,
             real, dimension(nn,4), intent(inout) qA,
             real, dimension(nn,4), intent(inout) qD,
             integer, dimension(nn,4), intent(inout) qjA)
push a new element at the back of the queue
```

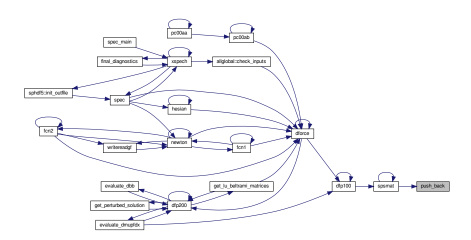
## **Parameters**

iq	
nq	
NN	
vΑ	
vD	
vjA	
qΑ	
qD	
qjA	

References constants::zero.

Referenced by spsmat().

Here is the caller graph for this function:



```
10.37.2.2 clean_queue() subroutine clean_queue (
             integer, dimension(4), intent(inout) nq,
             integer, intent(in) NN,
             real, dimension(nn,4), intent(inout) qA,
             real, dimension(nn,4), intent(inout) qD,
             integer, dimension(nn,4), intent(inout) qjA)
```

clean the queue

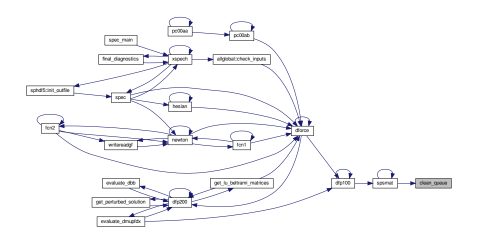
#### **Parameters**

nq	
NN	
qΑ	
qD	
qjA	

References constants::zero.

Referenced by spsmat().

Here is the caller graph for this function:



```
10.37.2.3 addline() subroutine addline ( integer, dimension(4), intent(inout) nq,
```

```
integer, dimension(4), intent(inout) nq,
integer, intent(inout) NN,
real, dimension(nn,4), intent(inout) qA,
real, dimension(nn,4), intent(inout) qD,
integer, dimension(nn,4), intent(inout) qjA,
integer, intent(inout) ns,
integer, intent(inout) nrow,
real, dimension(*) dMAS,
real, dimension(*) dMDS,
integer, dimension(*) jdMAS,
integer, dimension(*) idMAS)
```

add the content from the queue to the real matrices

#### **Parameters**

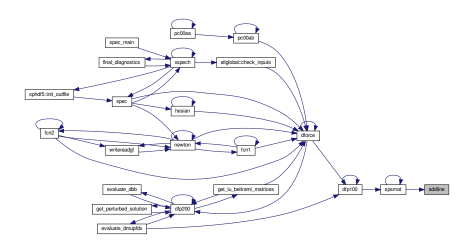
nq	
NN	
qΑ	
qD	
qjA	
ns	
nrow	
dMAS	
dMDS	

#### **Parameters**

jdMAS	
idMAS	

#### Referenced by spsmat().

Here is the caller graph for this function:



## 10.38 src/stzxyz.f90 File Reference

Calculates coordinates,  $\mathbf{x}(s, \theta, \zeta) \equiv R \mathbf{e}_R + Z \mathbf{e}_Z$ , and metrics, at given  $(s, \theta, \zeta)$ .

#### **Functions/Subroutines**

• subroutine stzxyz (Ivol, stz, RpZ) Calculates coordinates,  $\mathbf{x}(s,\theta,\zeta) \equiv R \, \mathbf{e}_R + Z \, \mathbf{e}_Z$ , and metrics, at given  $(s,\theta,\zeta)$ .

## 10.38.1 Detailed Description

Calculates coordinates,  $\mathbf{x}(s, \theta, \zeta) \equiv R \mathbf{e}_R + Z \mathbf{e}_Z$ , and metrics, at given  $(s, \theta, \zeta)$ .

## 10.39 src/tr00ab.f90 File Reference

Calculates rotational transform given an arbitrary tangential field.

#### **Functions/Subroutines**

• subroutine tr00ab (Ivol, mn, NN, Nt, Nz, iflag, Idiota)

Calculates rotational transform given an arbitrary tangential field.

#### 10.39.1 Detailed Description

Calculates rotational transform given an arbitrary tangential field.

## 10.40 src/volume.f90 File Reference

Computes volume of each region; and, if required, the derivatives of the volume with respect to the interface geometry.

#### **Functions/Subroutines**

• subroutine volume (Ivol, vflag)

Computes volume of each region; and, if required, the derivatives of the volume with respect to the interface geometry.

#### 10.40.1 Detailed Description

Computes volume of each region; and, if required, the derivatives of the volume with respect to the interface geometry.

## 10.41 src/wa00aa.f90 File Reference

Constructs smooth approximation to wall.

#### Modules

· module laplaces

...todo...

#### **Functions/Subroutines**

• subroutine wa00aa (iwa00aa)

Constructs smooth approximation to wall.

· subroutine vacuumphi (Nconstraints, rho, fvec, iflag)

Compute vacuum magnetic scalar potential (?)

#### **Variables**

· logical laplaces::stage1

what is this?

logical laplaces::exterior

what is this?

· logical laplaces::dorm

what is this?

• integer laplaces::nintervals

what is this?

• integer laplaces::nsegments

what is this?

• integer laplaces::ic

what is this?

• integer laplaces::np4

what is this?

• integer laplaces::np1

what is this?

integer, dimension(:), allocatable laplaces::icint

what is this?

· real laplaces::originalalpha

what is this?

• real, dimension(:), allocatable laplaces::xpoly

what is this?

• real, dimension(:), allocatable laplaces::ypoly

what is this?

• real, dimension(:), allocatable laplaces::phi

what is this?

· real, dimension(:), allocatable laplaces::phid

what is this?

• real, dimension(:,:), allocatable laplaces::cc

what is this?

• integer laplaces::ilength

what is this?

· real laplaces::totallength

what is this?

• integer laplaces::niterations

counter; eventually redundant; 24 Oct 12;

• integer laplaces::iangle

angle; eventually redundant; 24 Oct 12;

real laplaces::rmid

used to define local polar coordinate; eventually redundant; 24 Oct 12;

· real laplaces::zmid

used to define local polar coordinate; eventually redundant; 24 Oct 12;

· real laplaces::alpha

eventually redundant; 24 Oct 12;

## 10.41.1 Detailed Description

Constructs smooth approximation to wall.

## 10.42 src/xspech.f90 File Reference

Main program.

## **Functions/Subroutines**

• program spec\_main

Main program of SPEC.

· subroutine xspech

Main subroutine of SPEC.

subroutine read\_command\_args

Read command-line arguments; in particular, determine input file (name or extension).

• subroutine spec

This is the main "driver" for the physics part of SPEC.

• subroutine final\_diagnostics

Final diagnostics.

• subroutine ending

Closes output files, writes screen summary.

## 10.42.1 Detailed Description

Main program.

## 10.42.2 Function/Subroutine Documentation

#### 10.42.2.1 spec\_main() program spec\_main

Main program of SPEC.

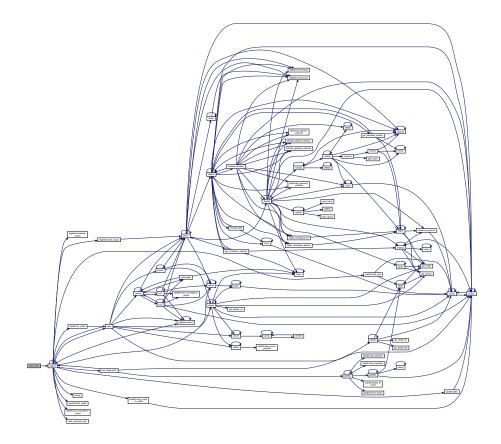
This only calls the xpech() subroutine to do a stand-alone SPEC run.

Returns

none

References xspech().

Here is the call graph for this function:



## 10.42.2.2 xspech() subroutine xspech

Main subroutine of SPEC.

This orchestrates a stand-alone SPEC run:

- · read the input file
- solve the MRxMHD equilibrium (see spec())
- · run some diagnostics on the results
- write the output file(s)

## reading input, allocating global variables

- The input namelists and geometry are read in via a call to readin(). A full description of the required input is given in global.f90.
- Most internal variables, global memory etc., are allocated in preset() .
- All quantities in the input file are mirrored into the output file's group  $/ \mathtt{input}$ .

#### preparing output file group iterations

• The group /iterations is created in the output file. This group contains the interface geometry at each iteration, which is useful for constructing movies illustrating the convergence. The data structure in use is an unlimited array of the following compound datatype:

```
DATATYPE H5T_COMPOUND {

H5T_NATIVE_INTEGER "nDcalls";

H5T_NATIVE_DOUBLE "Energy";

H5T_NATIVE_DOUBLE "ForceErr";

H5T_ARRAY { [Mvol+1] [mn] H5T_NATIVE_DOUBLE } "iRbc";

H5T_ARRAY { [Mvol+1] [mn] H5T_NATIVE_DOUBLE } "iZbs";

H5T_ARRAY { [Mvol+1] [mn] H5T_NATIVE_DOUBLE } "iRbs";

H5T_ARRAY { [Mvol+1] [mn] H5T_NATIVE_DOUBLE } "iZbc";

}
```

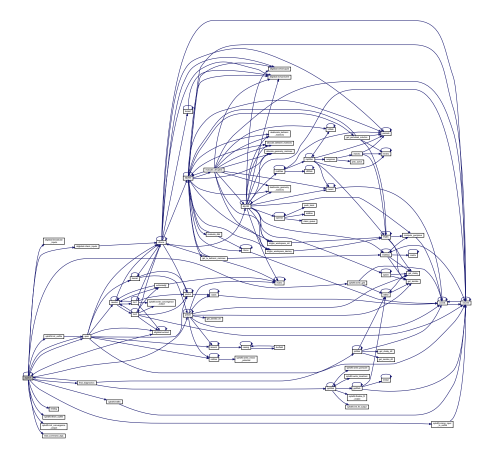
#### restart files

• wrtend() is called to write the restart files.

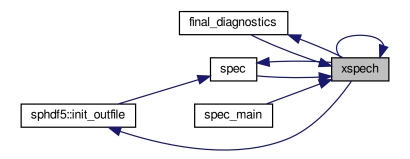
References allglobal::broadcast\_inputs(), allglobal::check\_inputs(), allglobal::cpus, ending(), final\_diagnostics(), sphdf5::finish\_outfile(), sphdf5::hdfint(), sphdf5::init\_convergence\_output(), sphdf5::init\_outfile(), numerical::machprec, sphdf5::mirror\_input\_to\_outfile(), allglobal::mpi\_comm\_spec, allglobal::myid, allglobal::ncpu, fileunits::ounit, preset(), read\_command\_args(), numerical::small, spec(), numerical::vsmall, sphdf5::write\_grid(), allglobal::wrtend(), and xspech().

Referenced by final\_diagnostics(), spec(), spec\_main(), and xspech().

Here is the call graph for this function:



Here is the caller graph for this function:



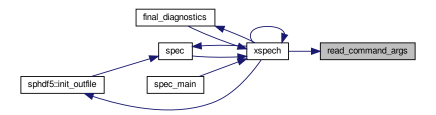
#### 10.42.2.3 read command args() subroutine read\_command\_args

Read command-line arguments; in particular, determine input file (name or extension).

- The input file name, ext, is given as the first command line input, and the input file itself is then ext.sp.
- Alternatively, you can directly specify the input file itself as ext.sp.
- · Additional command line inputs recognized are:
  - help or -h will give help information to user
  - readin will immediately set Wreadin=T; this may be over-ruled when the namelist screenlist is read

References allglobal::cpus, allglobal::mpi\_comm\_spec, allglobal::myid, fileunits::ounit, and inputlist::wreadin. Referenced by xspech().

Here is the caller graph for this function:



## **10.42.2.4 spec()** subroutine spec

This is the main "driver" for the physics part of SPEC.

Picard iterations are performed (if in free-boundary mode) and within each Picard iteration, the fixed-boundary problem is solved (also iteratively). **packing geometrical degrees-of-freedom into vector** 

• If NGdof.gt.0, where NGdof counts the geometrical degrees-of-freedom, i.e. the  $R_{bc}$ ,  $Z_{bs}$ , etc., then packxi() is called to "pack" the geometrical degrees-of-freedom into position (0:NGdof).

#### initialize adiabatic constants

• If Ladiabatic.eq.0, then the "adiabatic constants" in each region,  $P_v$ , are calculated as

$$P_v \equiv p_v V_v^{\gamma},\tag{262}$$

where  $p_v \equiv \texttt{pressure}$  (vvol) , the volume  $V_v$  of each region is computed by volume() , and the adiabatic index  $\gamma \equiv \texttt{gamma}$  .

#### solving force-balance

- If there are geometrical degress of freedom, i.e. if NGdof.gt.0, then
  - Todo If Lminimize.eq.1, call pc00aa() to find minimum of energy functional using quasi-Newton, preconditioned conjugate gradient method, E04DGF
  - If Lfindzero.gt.0, call newton() to find extremum of constrained energy functional using a Newton method, C05PDF.

#### post diagnostics

- The pressure is computed from the adiabatic constants from Eqn. (262), i.e.  $p = P/V^{\gamma}$ .
- The Beltrami/vacuum fields in each region are re-calculated using dforce().
- If Lcheck.eq.5.or. LHevalues.or. LHevectors.or. Lperturbed.eq.1, then the force-gradient matrix is examined using hesian().

#### free-boundary: re-computing normal field

- If Lfreebound.eq.1 and Lfindzero.gt.0 and mfreeits.ne.0, then the magnetic field at the computational boundary produced by the plasma currents is computed using bnorml().
- The "new" magnetic field at the computational boundary produced by the plasma currents is updated using a Picard scheme:

$$\operatorname{Bns}_{i}^{j} = \lambda \operatorname{Bns}_{i}^{j-1} + (1 - \lambda) \operatorname{Bns}_{i}, \tag{263}$$

where j labels free-boundary iterations, the "blending parameter" is  $\lambda \equiv \mathtt{gBnbld}$ , and  $\mathtt{Bns}_i$  is computed by virtual casing. The subscript "\$i\$" labels Fourier harmonics.

• If the new (unblended) normal field is *not* sufficiently close to the old normal field, as quantified by <code>gBntol</code>, then the free-boundary iterations continue. This is quantified by

$$\sum_{i} |\operatorname{Bns}_{i}^{j-1} - \operatorname{Bns}_{i}|/N, \tag{264}$$

where N is the total number of Fourier harmonics.

- There are several choices that are available:
  - if mfreeits=-2: the vacuum magnetic field (really, the normal component of the field produced by the external currents at the computational boundary) required to hold the given equilibrium is written to file. This information is required as input by FOCUS [8] for example. (This option probably needs to revised.)
  - if mfreeits=-1: after the plasma field is computed by virtual casing, the vacuum magnetic field is set to exactly balance the plasma field (again, we are really talking about the normal component at the computational boundary.) This will ensure that the computational boundary itself if a flux surface of the total magnetic field.
  - if mfreeits=0: the plasma field at the computational boundary is not updated; no "free-boundary" iterations take place.

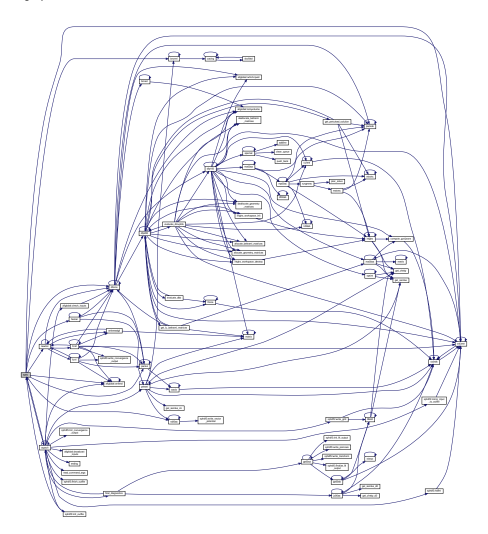
- if mfreeits>0: the plasma field at the computational boundary is updated according to the above blending Eqn. (263), and the free-boundary iterations will continue until either the tolerance condition is met (see <code>gBntol</code> and Eqn. (264)) or the maximum number of free-boundary iterations, namely <code>mfreeits</code>, is reached. For this case, <code>Lzerovac</code> is relevant: if <code>Lzerovac=1</code>, then the vacuum field is set equal to the normal field at every iteration, which results in the computational boundary being a flux surface. (I am not sure if this is identical to setting <code>mfreeits=-1</code>; the logic etc. needs to be revised.)

## output files: vector potential

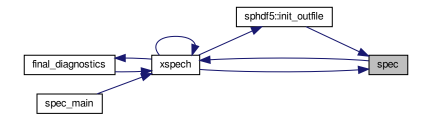
• The vector potential is written to file using ra00aa().

References inputlist::adiabatic, allglobal::ate, allglobal::ate, allglobal::aze, allglobal::aze, allglobal::bbe, allglobal::bbo, allglobal::betramierror, bnorml(), allglobal::cfmn, allglobal::cpus, dforce(), allglobal::dma, allglobal::dmb, allglobal::dmd, allglobal::dmg, allglobal::dpflux, allglobal::dfflux, allglobal::efmn, allglobal::first\_free\_bound, allglobal::forceerr, inputlist::gamma, inputlist::gbnbld, inputlist::gbntol, inputlist::helicity, hesian(), allglobal::ibnc, allglobal::ibns, inputlist::igeometry, allglobal::iie, allglobal::iio, allglobal::im, allglobal::imagneticok, allglobal:iin, allglobal::irbc, allglobal::irbs, inputlist::isurf, allglobal::ivnc, allglobal::ivns, inputlist::ivolume, allglobal::izbc, allglobal::izbs, inputlist::ladiabatic, inputlist::lautoinitbn, inputlist::lcheck, inputlist::lconstraint, allglobal::lcoordinatesingularity, inputlist::lfindzero, inputlist::lfreebound, allglobal::lgdof, inputlist::lhevalues, inputlist::lhevectors, inputlist::lhmatrix, numerical::logtolerance, inputlist::lperturbed, allglobal::lplasmaregion, inputlist::lrad, fileunits::lunit, allglobal::lvacuumregion, inputlist::lzerovac, allglobal::mbpsi, inputlist::mfreeits, allglobal::mn, allglobal::mpi comm spec, inputlist::mu, constants::mu0, allglobal::myid, allglobal::ncpu, newton(), inputlist::nfp, allglobal::nfreeboundaryiterations, allglobal::ngdof, allglobal::notstellsym, inputlist::nppts, inputlist::nptrj, allglobal::ntz, inputlist::nvol, inputlist::odetol, allglobal::ofmn, constants::one, fileunits::ounit, packxi(), inputlist::pflux, inputlist::phiedge, constants::pi2, inputlist::pressure, inputlist::pscale, ra00aa(), inputlist::rbc, inputlist::rbs, allglobal::sfmn, allglobal::solution, inputlist::tflux, inputlist::vcasingtol, constants::version, volume(), numerical::vsmall, allglobal::vvolume, inputlist::wmacros, allglobal::wrtend(), xspech(), allglobal::yesstellsym, inputlist::zbc, inputlist::zbs, and constants::zero. Referenced by sphdf5::init outfile(), and xspech().

Here is the call graph for this function:



Here is the caller graph for this function:



# ${\bf 10.42.2.5} \quad {\bf final\_diagnostics()} \quad {\tt subroutine\ final\_diagnostics} \\ {\tt Final\ diagnostics}.$

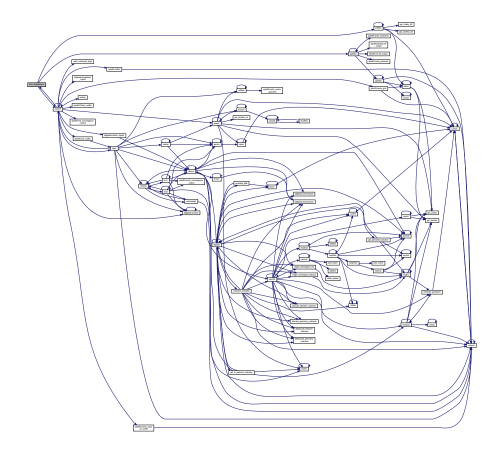
• sc00aa() is called to compute the covariant components of the magnetic field at the interfaces; these are related to the singular currents

- if Lcheck=1, jo00aa() is called to compute the error in the Beltrami equation
- pp00aa() is called to construct the Poincare plot by field-line following.

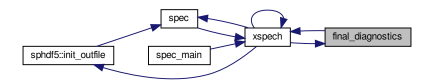
References allglobal::beltramierror, allglobal::btemn, allglobal::btemn, allglobal::bzemn, allglobal::bzemn, allglobal::bzemn, allglobal::bzemn, allglobal::bzemn, allglobal::bzemn, allglobal::bzemn, allglobal::cfmn, inputlist::igeometry, allglobal::imagneticok, allglobal::iquad, inputlist::isurf, inputlist::ivolume, jo00aa(), inputlist::lcheck, allglobal::lcoordinatesingularity, allglobal::lplasmaregion, allglobal::lvacuumregion, allglobal::mn, allglobal::mpi\_comm\_spec, inputlist::mu, allglobal::myid, allglobal::ncpu, inputlist::nppts, inputlist::nptrj, allglobal::ntz, inputlist::nvol, inputlist::odetol, allglobal::ofmn, fileunits::ounit, constants::pi2, pp00aa(), allglobal::sfmn, inputlist::wmacros, xspech(), and constants::zero.

Referenced by xspech().

Here is the call graph for this function:



Here is the caller graph for this function:



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## References

[1] J. D. Hanson. The virtual-casing principle and Helmholtz's theorem. *Plasma Phys. and Contr. Fusion*, 57(11):115006, sep 2015. 21

- [2] S. P. Hirshman and J. Breslau. Explicit spectrally optimized Fourier series for nested magnetic surfaces. *Phys. Plas.*, 5(7):2664–2675, 1998. 36
- [3] S. P. Hirshman and H. K. Meier. Optimized Fourier representations for three-dimensional magnetic surfaces. *Phys. Fluids*, 28(5):1387–1391, 1985. 36
- [4] S. R. Hudson, R. L. Dewar, M. J. Hole, and M. McGann. Non-axisymmetric, multi-region relaxed magnetohydro-dynamic equilibrium solutions. *Plasma Phys. and Contr. Fusion*, 54(1):014005, dec 2011. 14
- [5] S. A. Lazerson. The virtual-casing principle for 3D toroidal systems. *Plasma Phys. and Contr. Fusion*, 54(12):122002, nov 2012. 21
- [6] S. A. Lazerson, S. Sakakibara, and Y. Suzuki. A magnetic diagnostic code for 3D fusion equilibria. *Plasma Phys. and Contr. Fusion*, 55(2):025014, jan 2013. 137
- [7] V. D. Shafranov and L. E. Zakharov. Use of the virtual-casing principle in calculating the containing magnetic field in toroidal plasma systems. *Nucl. Fusion*, 12(5):599–601, sep 1972. 21
- [8] C. Zhu, S. R. Hudson, Y. Song, and Y. Wan. New method to design stellarator coils without the winding surface. *Nucl. Fusion*, 58(1):016008, nov 2017. 234

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