

VMEC

8.52

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Chapter 1

Educational VMEC

This is a heavily stripped-down version of the serial implementation of VMEC 8.52. It is forked from the `v251` branch of the `STELLOPT` repository.

The goal of this project is to have a version of VMEC which only computes the Stellarator MHD equilibrium and nothing more.

The `cmake` build system for stand-alone VMEC is borrowed from `hiddenSymmetries/VMEC2000` and from `ORNL-Fusion/LIBSTELL`.

1.1 Building

This is a fairly standard CMake setup, if you are used to it. Here is how it works:

- Create a directory `build` in the main folder: `mkdir build`
- Go into the `build` directory: `cd build`
- Run CMake: `cmake ..`
- Execute the actual build process: `make` (optional multi-threaded build: `make -j`)
- The VMEC executable `xvmec` is then located in `build/bin` with respect to the main folder.

1.2 Example Execution

- Change into the `test` dir: `cd test`
- Run the `Solov'ev` test case: `../build/bin/xvmec input.solovev`

1.3 External NESTOR

The free-boundary part of VMEC is the Neumann Solver for Toroidal Systems (NESTOR). Its source code is in a separate folder `NESTOR`. The appropriate reference is [https://doi.org/10.1016/0021-9991\(86\)90055-0](https://doi.org/10.1016/0021-9991(86)90055-0).

This version of NESTOR can be run stand-alone. It reads its inputs from a netCDF file and writes its outputs into another netCDF file. The main executable of this stand-alone version of NESTOR is `nestor_main.f90`. The input and output files are read and written in `nestor_io.f90`.

This version of VMEC can be configured to dump the corresponding input and output files, but still run the compiled-in version of NESTOR. This is enabled via the logical flag `ldump_vacuum_ref` in `funct3d.f90`.

Also, an external NESTOR implementation can be called instead of using the compiled-in version of NESTOR. This is enabled via the logical flag `lexternal_nestor` in `funct3d.f90`. The corresponding system call to execute the external NESTOR implementation has to be specified in `nestor_executable` in `funct3d.f90`.

1.4 Angle Constraint

The poloidal angle-like coordinate is a priori not uniquely defined and needs special care. The version of VMEC from the STELLOPT repo had essentially two options for this. They were alternatively compiled in via the preprocessor flag `_HBANGLE`.

1. The Hirshman-Breslau explicit spectrally optimized Fourier series (see <https://doi.org/10.1063/1.872954> for details) and
2. an unknown mixture of several constraints of the $m=1$ Fourier coefficients (the logical `lconm1` is true for this constraint).

By default, the `_HBANGLE` preprocessor flag is not active and thus, the "old" $m=1$ constraint is active.

This version of VMEC has most, if not all, of its preprocessor flags explicitly expanded. It became clear that it is nevertheless useful to have at least a vague idea of what parts of the code are related to the angle constraint. Therefore, those parts of VMEC related to the $m=1$ constraint are marked to start with

```
! #ifndef _hbangle
```

and end with

```
! #end /* ndef _HBANGLE */
```

Chapter 2

Modules Index

2.1 Modules List

Here is a list of all documented modules with brief descriptions:

line_segment	This module contains code to create a profile constructed of line segments. These line segments are assumed to be specified such that $xx(i) < xx(i + 1)$	11
mgrid_mod	Precomputed table of magnetic field due to confinement coils	11
nestor_io	Input and Output for stand-alone NESTOR	13

Chapter 3

Data Type Index

3.1 Data Types List

Here are the data types with brief descriptions:

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---	----

Chapter 4

File Index

4.1 File List

Here is a list of all documented files with brief descriptions:

src/add_fluxes.f90	Add the magnetic fluxes to the tangential derivatives of λ to arrive at the contravariant magnetic field components B^θ and B^ζ	19
src/alias.f90	Fourier transform alias force and also return intermediate output	20
src/allocate_funct3d.f90	Allocate arrays required in <code>funct3d()</code>	21
src/allocate_ns.f90	Allocate arrays depending on the number of flux surfaces <code>ns</code>	21
src/allocate_nunv.f90	Allocate arrays depending on the number of Fourier coefficients <code>nunv</code>	23
src/aspectratio.f90	Compute aspect-ratio (independent of elongation): $A = \langle R \rangle / \sqrt{\langle ab \rangle}$	23
src/bcovar.f90	Compute the covariant components of the magnetic field B_θ, B_ζ	23
src/bextrema.f90	Computes minimum and maximum $ \mathbf{B} $ along ζ between two angle lines ($\theta = 0, \pi$)	25
src/bss.f90	Computes <code>br</code> , <code>bphi</code> , <code>bz</code> , <code>bsubs</code> on half-radial mesh	26
src/calc_fbal.f90	Compute flux-surface averaged radial force balance $\nabla p - \langle \mathbf{j} \times \mathbf{B} \rangle$	27
src/convert.f90	Convert internal mode representation to standard form for output (coefficients of $\cos(\mu-nv)$, $\sin(\mu-nv)$ without internal <code>mscale</code> , <code>nscale</code> norms)	28
src/elongation.f90	Compute Waist thickness and height in $\varphi = 0, \pi$ symmetry planes	39
src/eqfor.f90	Basis physics analysis and evaluation of force balance. This is where most of the contents of the <code>threed1</code> output file is computed	40
src/eqsolve.f90	Iteratively evolve the Fourier coefficients that specify the equilibrium	42
src/evolve.f90	Take a single time step in Fourier space to evolve the Fourier coefficients describing the equilibrium towards force balance	44
src/fileout.f90	Write the output files	45

src/fixaray.f90	Allocate and fill some fixed-size arrays (only depending on Fourier resolution)	48
src/flip_theta.f90	Flip the definition of the poloidal angle in the user-provided initial guess for the LCFS geometry	48
src/forces.f90	Compute the real-space MHD forces	49
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src/fsym_fft.f90	Fourier transforms	51
src/fsym_invfft.f90	Extends function from <code>ntheta2</code> to <code>ntheta3</code> range	53
src/funct3d.f90	Evaluate the three-dimensional MHD energy functional	54
src/functions.f	This module contains functions used by the profiles	55
src/getbsubs.f90	Solves the radial force balance $\mathbf{B} \cdot \mathbf{B}_s = F_s$ for B_s in real space using collocation	57
src/getcurmid.f90	Get current at midplane (?)	58
src/getfsq.f90	Compute total force residual on flux surfaces	59
src/guess_axis.f90	Computes guess for magnetic axis if user guess leads to initial sign change of Jacobian	60
src/heading.f90	Open output files and print banner message at the top	61
src/initialize_radial.f90	Allocates memory for radial arrays and initializes radial profiles	62
src/interp.f90	Interpolate R , Z and λ on full grid	63
src/jacobian.f90	Evaluate the Jacobian of the transform from flux- to cylindrical coordinates	64
src/jxbforce.f90	Program for computing local $\mathbf{K} \times \mathbf{B} = \nabla p$ force balance	65
src/lamcal.f90	Normalization parameters for λ	66
src/line_segment.f	This module contains code to create a profile constructed of line segments	67
src/magnetic_fluxes.f90	Compute toroidal and poloidal magnetic flux profiles	68
src/mercier.f90	Evaluate the Mercier stability criterion	71
src/mgrid_mod.f	Precomputed table of magnetic field due to confinement coils	73
src/open_output_files.f90	Open output files	84
src/parse_extension.f	Parse the first command-line argument into a filename	85
src/precondn.f90	Compute preconditioning matrix elements for R , Z force	86
src/printout.f90	Print iteration progress to screen and <code>threed1</code> output file	87

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Chapter 5

Module Documentation

5.1 line_segment Module Reference

This module contains code to create a profile constructed of line segments. These line segments are assumed to be specified such that $xx(i) < xx(i + 1)$.

Functions/Subroutines

- subroutine, public **line_seg** (x, y, xx, yy, n)
- subroutine, public **line_seg_int** (x, y, xx, yy, n)
- logical function, public **line_seg_test** ()

5.1.1 Detailed Description

This module contains code to create a profile constructed of line segments. These line segments are assumed to be specified such that $xx(i) < xx(i + 1)$.

5.2 mgrid_mod Module Reference

Precomputed table of magnetic field due to confinement coils.

Functions/Subroutines

- subroutine **read_mgrid** (mgrid_file, extcur, nv, nfp, lscreen, ier_flag)
- subroutine **sum_bfield** (bfield, bf_add, cur, n1)
- subroutine **assign_bptrs** (bptr)
- subroutine **free_mgrid** (istat)

Variables

- integer, parameter **nlimset** = 2
- character(len= *), parameter **vn_br0** = 'br'
- character(len= *), parameter **vn_bp0** = 'bp'
- character(len= *), parameter **vn_bz0** = 'bz'
- character(len= *), parameter **vn_ir** = 'ir'
- character(len= *), parameter **vn_jz** = 'jz'
- character(len= *), parameter **vn_kp** = 'kp'
- character(len= *), parameter **vn_nfp** = 'nfp'
- character(len= *), parameter **vn_rmin** = 'rmin'
- character(len= *), parameter **vn_rmax** = 'rmax'
- character(len= *), parameter **vn_zmin** = 'zmin'
- character(len= *), parameter **vn_zmax** = 'zmax'
- character(len= *), parameter **vn_coilgrp** = 'coil_group'
- character(len= *), parameter **vn_nextcur** = 'nextcur'
- character(len= *), parameter **vn_mgmode** = 'mgrid_mode'
- character(len= *), parameter **vn_coilcur** = 'raw_coil_cur'
- character(len= *), parameter **ln_next** = 'External currents'
- integer **nr0b**
- integer **np0b**
- integer **nfper0**
- integer **nz0b**
- integer **nobd**
- integer **nobser**
- integer **nextcur**
- integer **nbfldn**
- integer **nbsets**
- integer **nbcoilsn**
- integer **nbvac**
- integer **nbcoil_max**
- integer **nlim**
- integer **nlim_max**
- integer **nsets**
- integer **nrgrid**
- integer **nzgrid**
- integer, dimension(:), allocatable **needflx**
- integer, dimension(:), allocatable **nbcoils**
- integer, dimension(:), allocatable **limitr**
- integer, dimension(:), allocatable **nsetsn**
- integer, dimension(:,:), allocatable **iconnect**
- integer, dimension(:,:), allocatable **needbfld**
- real(rprec) **rminb**
- real(rprec) **zminb**
- real(rprec) **rmaxb**
- real(rprec) **zmaxb**
- real(rprec) **delrb**
- real(rprec) **delzb**
- real(rprec) **rx1**
- real(rprec) **rx2**
- real(rprec) **zy1**
- real(rprec) **zy2**
- real(rprec) **condif**
- real(rprec), dimension(:,:), allocatable, target **bvac**
- real(rprec), dimension(:,:), pointer **brvac**

- real(rprec), dimension(:,:), pointer **bzvac**
- real(rprec), dimension(:,:), pointer **bpvac**
- real(rprec), dimension(:,:), allocatable **unpsiext**
- real(rprec), dimension(:,:), allocatable **plbflf**
- real(rprec), dimension(:,:), allocatable **rbcoil**
- real(rprec), dimension(:,:), allocatable **zbcoil**
- real(rprec), dimension(:,:), allocatable **abcoil**
- real(rprec), dimension(:,:), allocatable **bcoil**
- real(rprec), dimension(:,:), allocatable **rbcoilsqr**
- real(rprec), dimension(:), allocatable **raw_coil_current**
- real(rprec), dimension(:), allocatable **xobser**
- real(rprec), dimension(:), allocatable **zobser**
- real(rprec), dimension(:), allocatable **xobsqr**
- real(rprec), dimension(:), allocatable **dsiext**
- real(rprec), dimension(:), allocatable **psiext**
- real(rprec), dimension(:), allocatable **plflux**
- real(rprec), dimension(:), allocatable **b_chi**
- character(len=300) **mgrid_path**
- character(len=300) **mgrid_path_old** = " "
- character(len=30), dimension(:), allocatable **curlabel**
- character(len=15), dimension(:), allocatable **dsilabel**
- character(len=15), dimension(:), allocatable **bloopnames**
- character(len=30) **tokid**
- real(rprec), dimension(:,:), allocatable **dbcoil**
- real(rprec), dimension(:,:), allocatable **pfcspec**
- real(rprec), dimension(:,:), allocatable **rlim**
- real(rprec), dimension(:,:), allocatable **zlim**
- real(rprec), dimension(:,:), allocatable **reslim**
- real(rprec), dimension(:,:), allocatable **seplim**
- character(len=1) **mgrid_mode**

5.2.1 Detailed Description

Precomputed table of magnetic field due to confinent coils.

5.3 nestor_io Module Reference

Input and Output for stand-alone NESTOR.

Functions/Subroutines

- subroutine **read_nestor_inputs** (vac_file)
- subroutine **write_nestor_outputs** (vac_file, lasym, ivac, ier_flag)

Variables

- character(len=255) **input_extension**
- character(len=255) **mgrid_file**
- real(dp), dimension(:), allocatable **extcur**
- real(dp), dimension(:), allocatable **raxis**
- real(dp), dimension(:), allocatable **zaxis**
- real(dp), dimension(:), allocatable **xm**
- real(dp), dimension(:), allocatable **xn**
- real(dp), dimension(:), allocatable **rmnc**
- real(dp), dimension(:), allocatable **zmns**
- real(dp), dimension(:), allocatable **rmns**
- real(dp), dimension(:), allocatable **zmnc**
- real(dp), dimension(:), allocatable **wint**
- integer **nfp**
- integer **ntor**
- integer **mpol**
- integer **ntheta**
- integer **nzeta**
- integer **nextcur**
- integer **ier_flag**
- integer **ivac**
- integer **ivacskip**
- integer **mnmax**
- integer **vacuum_calls**
- logical **lasym**
- real(dp) **ctor**
- real(dp) **rbtor**
- real(dp) **signgs**
- integer **mnpd2_nestor**
- real(dp), dimension(:), allocatable **amatsav_nestor**
- real(dp), dimension(:), allocatable **bvecsav_nestor**
- real(dp) **bsubvvac_nestor**
- character(len= *), dimension(1), parameter **mn1dim** = ('mn_mode'/)
- character(len= *), dimension(1), parameter **mnpotdim** = ('mn_mode_pot'/)
- character(len= *), dimension(1), parameter **nzntdim** = ('nznt'/)
- character(len= *), dimension(1), parameter **nzetadim** = ('nzeta'/)
- character(len= *), dimension(1), parameter **nextcurim** = ('nextcur'/)
- character(len= *), dimension(1), parameter **bvecsavdim** = ('mnpd2'/)
- character(len= *), dimension(1), parameter **amatsavdim** = ('mnpd2_times_mnpd2'/)
- character(len= *), dimension(2), parameter **r2dim** = ('mn_mode','radius '/)
- character(len= *), parameter **vn_vacuum_calls** = 'vacuum_calls'
- character(len= *), parameter **vn_ier_flag** = "ier_flag"
- character(len= *), parameter **vn_mgrid** = "mgrid_file"
- character(len= *), parameter **vn_inputext** = "input_extension"
- character(len= *), parameter **vn_ivacskip** = "ivacskip"
- character(len= *), parameter **vn_ivac** = "ivac"
- character(len= *), parameter **vn_nfp** = "nfp"
- character(len= *), parameter **vn_ntor** = "ntor"
- character(len= *), parameter **vn_mpol** = "mpol"
- character(len= *), parameter **vn_nzeta** = "nzeta"
- character(len= *), parameter **vn_ntheta** = "ntheta"
- character(len= *), parameter **vn_mnmax** = "mnmax"
- character(len= *), parameter **vn_pmod** = "xm"
- character(len= *), parameter **vn_tmod** = "xn"

- character(len= *), parameter **vn_rmnc** = "rmnc"
- character(len= *), parameter **vn_zmns** = "zmns"
- character(len= *), parameter **vn_rmns** = "rmns"
- character(len= *), parameter **vn_zmnc** = "zmnc"
- character(len= *), parameter **vn_rbtor** = "rbtor"
- character(len= *), parameter **vn_ctor** = "ctor"
- character(len= *), parameter **vn_lasym** = "lasym"
- character(len= *), parameter **vn_signgs** = "signgs"
- character(len= *), parameter **vn_extcur** = "extcur"
- character(len= *), parameter **vn_raxis_nestor** = "raxis_nestor"
- character(len= *), parameter **vn_zaxis_nestor** = "zaxis_nestor"
- character(len= *), parameter **vn_wint** = "wint"
- character(len= *), parameter **vn_bsqvac** = "bsqvac"
- character(len= *), parameter **vn_mnpd** = "mnpd"
- character(len= *), parameter **vn_xmpot** = "xmpot"
- character(len= *), parameter **vn_xnpot** = "xnpot"
- character(len= *), parameter **vn_potvac** = "potvac"
- character(len= *), parameter **vn_brv** = "brv"
- character(len= *), parameter **vn_bphiv** = "bphiv"
- character(len= *), parameter **vn_bzv** = "bzv"
- character(len= *), parameter **vn_bsubvvac** = "bsubvvac"
- character(len= *), parameter **vn_amatsav** = "amatsav"
- character(len= *), parameter **vn_bvecsav** = "bvecsav"
- character(len= *), parameter **vn_mnpd2** = "mnpd2"
- character(len= *), parameter **vn_r1b** = "r1b"
- character(len= *), parameter **vn_rub** = "rub"
- character(len= *), parameter **vn_rvb** = "rvb"
- character(len= *), parameter **vn_z1b** = "z1b"
- character(len= *), parameter **vn_zub** = "zub"
- character(len= *), parameter **vn_zvb** = "zvb"
- character(len= *), parameter **vn_ruu** = "ruu"
- character(len= *), parameter **vn_ruv** = "ruv"
- character(len= *), parameter **vn_rvv** = "rvv"
- character(len= *), parameter **vn_zuu** = "zuu"
- character(len= *), parameter **vn_zuv** = "zuv"
- character(len= *), parameter **vn_zvv** = "zvv"
- character(len= *), parameter **vn_guu_b** = "guu_b"
- character(len= *), parameter **vn_guv_b** = "guv_b"
- character(len= *), parameter **vn_gvv_b** = "gvv_b"
- character(len= *), parameter **vn_rzb2** = "rzb2"
- character(len= *), parameter **vn_snr** = "snr"
- character(len= *), parameter **vn_snv** = "snv"
- character(len= *), parameter **vn_snz** = "snz"
- character(len= *), parameter **vn_drv** = "drv"
- character(len= *), parameter **vn_auu** = "auu"
- character(len= *), parameter **vn_auv** = "auv"
- character(len= *), parameter **vn_avv** = "avv"
- character(len= *), parameter **vn_rcosuv** = "rcosuv"
- character(len= *), parameter **vn_rsinuv** = "rsinuv"
- character(len= *), parameter **vn_brاد** = "brاد"
- character(len= *), parameter **vn_bphi** = "bphi"
- character(len= *), parameter **vn_bz** = "bz"
- character(len= *), parameter **vn_bexu** = "bexu"
- character(len= *), parameter **vn_bexv** = "bexv"
- character(len= *), parameter **vn_bexn** = "bexn"

- character(len= *), parameter **vn_bexni** = "bexni"
- character(len= *), parameter **vn_grpmn** = "grpmn"
- character(len= *), parameter **vn_adp** = "adp"
- character(len= *), parameter **vn_adm** = "adm"
- character(len= *), parameter **vn_cma** = "cma"
- character(len= *), parameter **vn_sqrtc** = "sqrtc"
- character(len= *), parameter **vn_sqrta** = "sqrta"
- character(len= *), parameter **vn_delt1u** = "delt1u"
- character(len= *), parameter **vn_azp1u** = "azp1u"
- character(len= *), parameter **vn_azm1u** = "azm1u"
- character(len= *), parameter **vn_cma11u** = "cma11u"
- character(len= *), parameter **vn_r1p** = "r1p"
- character(len= *), parameter **vn_r1m** = "r1m"
- character(len= *), parameter **vn_r0p** = "r0p"
- character(len= *), parameter **vn_r0m** = "r0m"
- character(len= *), parameter **vn_ra1p** = "ra1p"
- character(len= *), parameter **vn_ra1m** = "ra1m"
- character(len= *), parameter **vn_sqad1u** = "sqad1u"
- character(len= *), parameter **vn_sqad2u** = "sqad2u"
- character(len= *), parameter **vn_all_tlp** = "all_tlp"
- character(len= *), parameter **vn_all_tlm** = "all_tlm"
- character(len= *), parameter **vn_all_slp** = "all_slp"
- character(len= *), parameter **vn_all_slm** = "all_slm"
- character(len= *), parameter **vn_m_map** = "m_map"
- character(len= *), parameter **vn_n_map** = "n_map"
- character(len= *), parameter **vn_green** = "green"
- character(len= *), parameter **vn_greenp** = "greenp"
- character(len= *), parameter **vn_tanu** = "tanu"
- character(len= *), parameter **vn_tanv** = "tanv"
- character(len= *), parameter **vn_gstore** = "gstore"
- character(len= *), parameter **vn_grpmn_m_map** = "grpmn_m_map"
- character(len= *), parameter **vn_grpmn_n_map** = "grpmn_n_map"
- character(len= *), parameter **vn_imirr** = "imirr"
- character(len= *), parameter **vn_amatrix** = "amatrix"
- character(len= *), parameter **vn_potu** = "potu"
- character(len= *), parameter **vn_potv** = "potv"
- character(len= *), parameter **vn_bsubu** = "bsubu"
- character(len= *), parameter **vn_bsubv** = "bsubv"

5.3.1 Detailed Description

Input and Output for stand-alone NESTOR.

Chapter 6

Data Type Documentation

6.1 read_wout_mod::read_wout_file Interface Reference

Public Member Functions

- subroutine **readw_and_open** (file_or_extension, ierr, iopen)

6.1.1 Detailed Description

Definition at line 236 of file read_wout_mod.f.

Chapter 7

File Documentation

7.1 src/add_fluxes.f90 File Reference

Add the magnetic fluxes to the tangential derivatives of λ to arrive at the contravariant magnetic field components B^θ and B^ζ .

Functions/Subroutines

- subroutine [add_fluxes](#) (overg, bsupu, bsupv)

Add the magnetic fluxes to the tangential derivatives of λ to arrive at the contravariant magnetic field components B^θ and B^ζ .

7.1.1 Detailed Description

Add the magnetic fluxes to the tangential derivatives of λ to arrive at the contravariant magnetic field components B^θ and B^ζ .

7.1.2 Function/Subroutine Documentation

7.1.2.1 add_fluxes()

```
subroutine add_fluxes (  
    real(rprec), dimension(nrzt), intent(in) overg,  
    real(rprec), dimension(nrzt), intent(inout) bsupu,  
    real(rprec), dimension(nrzt), intent(inout) bsupv )
```

Add the magnetic fluxes to the tangential derivatives of λ to arrive at the contravariant magnetic field components B^θ and B^ζ .

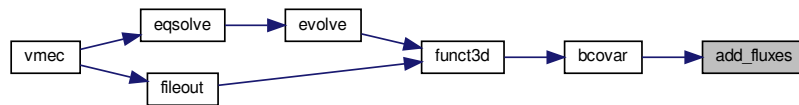
Parameters

<i>overg</i>	$1/\sqrt{g}$
<i>bsupu</i>	B^θ
<i>bsupv</i>	B^ζ

Definition at line 11 of file add_fluxes.f90.

Referenced by bcovar().

Here is the caller graph for this function:



7.2 src/alias.f90 File Reference

Fourier transform alias force and also return intermediate output.

Functions/Subroutines

- subroutine [alias](#) (gcons, ztemp, gcs, gsc, gcc, gss)
Fourier transform alias force from ztemp to gcons and also return intermediate output in g(c,s)(c,s)

7.2.1 Detailed Description

Fourier transform alias force and also return intermediate output.

7.2.2 Function/Subroutine Documentation

7.2.2.1 alias()

```

subroutine alias (
    real(rprec), dimension(ns*nzeta,ntheta3), intent(out) gcons,
    real(rprec), dimension(ns*nzeta,ntheta3), intent(in) ztemp,
    real(rprec), dimension(ns,0:ntor,0:mpoll), intent(inout) gcs,
    real(rprec), dimension(ns,0:ntor,0:mpoll), intent(inout) gsc,
    real(rprec), dimension(ns,0:ntor,0:mpoll), intent(inout) gcc,
    real(rprec), dimension(ns,0:ntor,0:mpoll), intent(inout) gss )

```

Fourier transform alias force from ztemp to gcons and also return intermediate output in g(c,s)(c,s)

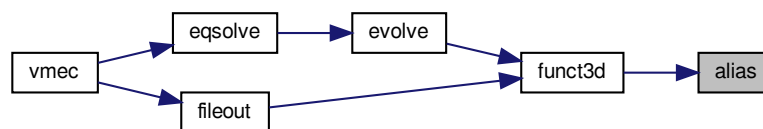
Parameters

<i>gcons</i>	
<i>ztemp</i>	
<i>gcs</i>	
<i>gsc</i>	
<i>gcc</i>	
<i>gss</i>	

Definition at line 12 of file alias.f90.

Referenced by `func3d()`.

Here is the caller graph for this function:



7.3 src/allocate_func3d.f90 File Reference

allocate arrays required in `func3d()`

Functions/Subroutines

- subroutine `allocate_func3d`
allocate arrays required in `func3d()`

7.3.1 Detailed Description

allocate arrays required in `func3d()`

7.4 src/allocate_ns.f90 File Reference

allocate arrays depending on the number of flux surfaces `ns`

Functions/Subroutines

- subroutine `allocate_ns` (`linterp`, `neqs_old`)
allocate arrays depending on the number of flux surfaces `ns`

7.4.1 Detailed Description

allocate arrays depending on the number of flux surfaces `ns`

7.4.2 Function/Subroutine Documentation

7.4.2.1 `allocate_ns()`

```
subroutine allocate_ns (
    logical, intent(in) linterp,
    integer, intent(in) neqs_old )
```

allocate arrays depending on the number of flux surfaces `ns`

Parameters

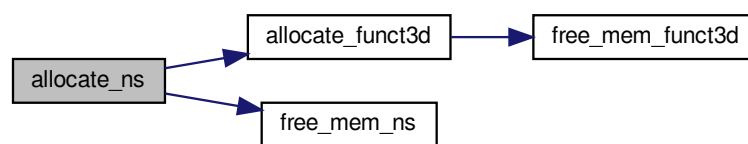
<i>linterp</i>	interpolate from coars to finer mesh?
<i>neqs_old</i>	previous number of degrees-of-freedom, i.e., Fourier coefficients for R , Z and λ

Definition at line 8 of file `allocate_ns.f90`.

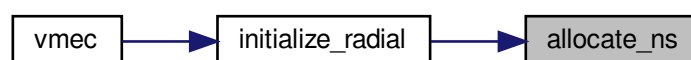
References `allocate_func3d()`, and `free_mem_ns()`.

Referenced by `initialize_radial()`.

Here is the call graph for this function:



Here is the caller graph for this function:



7.5 src/allocate_nunv.f90 File Reference

allocate arrays depending on the number of Fourier coefficients `nunv`

Functions/Subroutines

- subroutine `allocate_nunv`
allocate arrays depending on the number of Fourier coefficients `nunv`

7.5.1 Detailed Description

allocate arrays depending on the number of Fourier coefficients `nunv`

7.6 src/aspectratio.f90 File Reference

compute aspect-ratio (independent of elongation): $A = \langle R \rangle / \sqrt{\langle ab \rangle}$

Functions/Subroutines

- real(rprec) function `aspectratio` ()
compute aspect-ratio (independent of elongation): $A = \langle R \rangle / \sqrt{\langle ab \rangle}$ where $\pi \langle a \rangle^2 = \text{Area}$ (toroidally averaged) and $2\pi \langle R \rangle \text{Area} = \text{Volume}$

7.6.1 Detailed Description

compute aspect-ratio (independent of elongation): $A = \langle R \rangle / \sqrt{\langle ab \rangle}$

7.7 src/bcovar.f90 File Reference

Compute the covariant components of the magnetic field B_θ, B_ζ .

Functions/Subroutines

- subroutine `bcovar` (lu, lv)
Compute the covariant components of the magnetic field B_θ, B_ζ .

7.7.1 Detailed Description

Compute the covariant components of the magnetic field B_θ, B_ζ .

7.7.2 Function/Subroutine Documentation

7.7.2.1 bcovar()

```
subroutine bcovar (
    real(rprec), dimension(nrzt,0:1), intent(inout) lu,
    real(rprec), dimension(nrzt,0:1), intent(inout) lv )
```

Compute the covariant components of the magnetic field B_θ, B_ζ .

Parameters

lu	$\partial\lambda/\partial\theta$
lv	$\partial\lambda/\partial\zeta$

R12 from RP in force

Norm, unpreconditioned R,Z forces

Norm for preconditioned R,Z forces

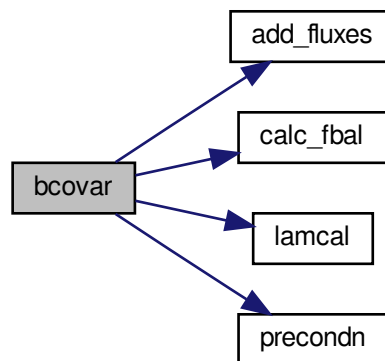
Norm for unpreconditioned Lambda force

Definition at line 8 of file bcovar.f90.

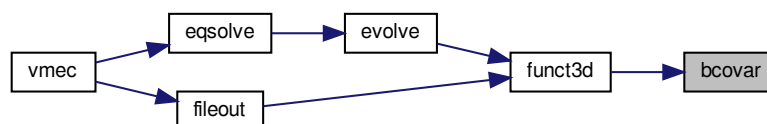
References `add_fluxes()`, `calc_fbal()`, `lamcal()`, and `precondn()`.

Referenced by `funct3d()`.

Here is the call graph for this function:



Here is the caller graph for this function:



7.8 src/bextrema.f90 File Reference

Computes minimum and maximum $|\mathbf{B}|$ along ζ between two angle lines ($\theta = 0, \pi$).

Functions/Subroutines

- subroutine [bextrema](#) (modb, bmin, bmax, nzeta, ntheta)
Computes minimum and maximum $|\mathbf{B}|$ along ζ between two angle lines ($\theta = 0, \pi$).

7.8.1 Detailed Description

Computes minimum and maximum $|\mathbf{B}|$ along ζ between two angle lines ($\theta = 0, \pi$).

7.8.2 Function/Subroutine Documentation

7.8.2.1 bextrema()

```
subroutine bextrema (
    real(rprec), dimension(nzeta,ntheta), intent(in) modb,
    real(rprec), dimension(ntheta), intent(out) bmin,
    real(rprec), dimension(ntheta), intent(out) bmax,
    integer, intent(in) nzeta,
    integer, intent(in) ntheta )
```

Computes minimum and maximum $|\mathbf{B}|$ along ζ between two angle lines ($\theta = 0, \pi$).

Parameters

<i>modb</i>	magnitude of magnetic field $ \mathbf{B} $
<i>bmin</i>	minimum value of $ \mathbf{B} $
<i>bmax</i>	maximum value of $ \mathbf{B} $
<i>nzeta</i>	number of grid points in toroidal direction
<i>ntheta</i>	number of grid points in poloidal direction

Definition at line 11 of file bextrema.f90.

Referenced by [eqfor\(\)](#).

Here is the caller graph for this function:



7.9 src/bss.f90 File Reference

Computes br, bphi, bz, bsubs on half-radial mesh.

Functions/Subroutines

- subroutine [bss](#) (r12, rs, zs, ru12, zu12, bsubs, bsupu, bsupv, br, bphi, bz)
Computes br, bphi, bz, bsubs on half-radial mesh.

7.9.1 Detailed Description

Computes br, bphi, bz, bsubs on half-radial mesh.

7.9.2 Function/Subroutine Documentation

7.9.2.1 bss()

```

subroutine bss (
    real(rprec), dimension(nrzt), intent(in) r12,
    real(rprec), dimension(nrzt), intent(in) rs,
    real(rprec), dimension(nrzt), intent(in) zs,
    real(rprec), dimension(nrzt), intent(in) ru12,
    real(rprec), dimension(nrzt), intent(in) zu12,
    real(rprec), dimension(nrzt), intent(out) bsubs,
    real(rprec), dimension(nrzt), intent(in) bsupu,
    real(rprec), dimension(nrzt), intent(in) bsupv,
    real(rprec), dimension(nrzt), intent(out) br,
    real(rprec), dimension(nrzt), intent(out) bphi,
    real(rprec), dimension(nrzt), intent(out) bz )

```

Computes br, bphi, bz, bsubs on half-radial mesh.

Parameters

<i>r12</i>	R^2
<i>rs</i>	$\partial R / \partial s$
<i>zs</i>	$\partial Z / \partial s$
<i>ru12</i>	$(\partial R / \partial \theta)^2$
<i>zu12</i>	$(\partial Z / \partial \theta)^2$
<i>bsubs</i>	covariant component of magnetic field B_s
<i>bsupu</i>	contravariant component of magnetic field B^θ
<i>bsupv</i>	contravariant component of magnetic field B^ζ
<i>br</i>	cylindrical component of magnetic field B^R
<i>bphi</i>	cylindrical component of magnetic field B^φ
<i>bz</i>	cylindrical component of magnetic field B^Z

Definition at line 17 of file bss.f90.

Referenced by eqfor().

Here is the caller graph for this function:



7.10 src/calc_fbal.f90 File Reference

Compute flux-surface averaged radial force balance $\nabla p - \langle \mathbf{j} \times \mathbf{B} \rangle$.

Functions/Subroutines

- subroutine [calc_fbal](#) (bsubu, bsubv)
Compute flux-surface averaged radial force balance $\nabla p - \langle \mathbf{j} \times \mathbf{B} \rangle$.

7.10.1 Detailed Description

Compute flux-surface averaged radial force balance $\nabla p - \langle \mathbf{j} \times \mathbf{B} \rangle$.

7.10.2 Function/Subroutine Documentation

7.10.2.1 calc_fbal()

```
subroutine calc_fbal (
    real(dp), dimension(1:nrzt), intent(in) bsubu,
    real(dp), dimension(1:nrzt), intent(in) bsubv )
```

Compute flux-surface averaged radial force balance $\nabla p - \langle \mathbf{j} \times \mathbf{B} \rangle$.

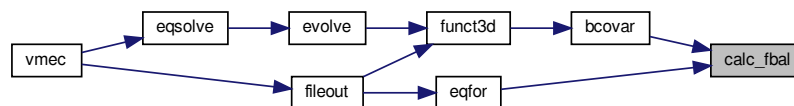
Parameters

<i>bsubu</i>	covariant component of magnetic field B_θ
<i>bsubv</i>	covariant component of magnetic field B_ζ

Definition at line 8 of file calc_fbal.f90.

Referenced by bcovar(), and eqfor().

Here is the caller graph for this function:



7.11 src/convert.f90 File Reference

Convert internal mode representation to standard form for output (coefficients of $\cos(\mu-nv)$, $\sin(\mu-nv)$ without internal `mscale`, `nscale` norms).

Functions/Subroutines

- subroutine [convert](#) (rmnc, zmns, lmns, rmns, zmnc, lmnc, rzl_array, js)

Convert internal mode representation to standard form for output (coefficients of $\cos(\mu-nv)$, $\sin(\mu-nv)$ without internal `mscale`, `nscale` norms).

7.11.1 Detailed Description

Convert internal mode representation to standard form for output (coefficients of $\cos(\mu-nv)$, $\sin(\mu-nv)$ without internal `mscale`, `nscale` norms).

7.11.2 Function/Subroutine Documentation

7.11.2.1 convert()

```

subroutine convert (
    real(rprec), dimension(mnmax), intent(out) rmnc,
    real(rprec), dimension(mnmax), intent(out) zmns,
    real(rprec), dimension(mnmax), intent(out) lmns,
    real(rprec), dimension(mnmax), intent(out) rmns,
    real(rprec), dimension(mnmax), intent(out) zmnc,
    real(rprec), dimension(mnmax), intent(out) lmnc,
    real(rprec), dimension(ns, 0:ntor, 0:mpoll, 3*ntmax), intent(in) rzl_array,
    integer, intent(in) js )

```

Convert internal mode representation to standard form for output (coefficients of $\cos(\mu-nv)$, $\sin(\mu-nv)$ without internal `mscale`, `nscale` norms).

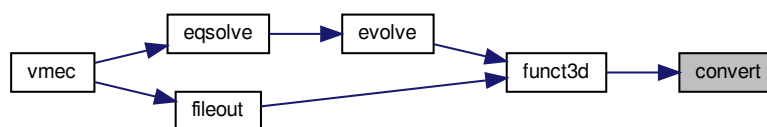
Parameters

<i>rmnc</i>	stellarator-symmetric Fourier coefficients of R
<i>zmns</i>	stellarator-symmetric Fourier coefficients of Z
<i>lmns</i>	stellarator-symmetric Fourier coefficients of λ
<i>rmns</i>	non-stellarator-symmetric Fourier coefficients of R
<i>zmnc</i>	non-stellarator-symmetric Fourier coefficients of Z
<i>lmnc</i>	non-stellarator-symmetric Fourier coefficients of λ
<i>rzl_array</i>	state vector (all Fourier coefficients) of VMEC
<i>js</i>	index of flux surface at which to do the conversion

Definition at line 16 of file `convert.f90`.

Referenced by `funct3d()`.

Here is the caller graph for this function:



7.12 src/data/fbal.f90 File Reference

Variables

- `real(dp), dimension(:), allocatable fbal::rzu_fac`
- `real(dp), dimension(:), allocatable fbal::rru_fac`
- `real(dp), dimension(:), allocatable fbal::frcc_fac`
- `real(dp), dimension(:), allocatable fbal::fzsc_fac`

7.13 src/data/realspace.f90 File Reference

Variables

- `real(rprec), dimension(:, :), allocatable realspace::r1`
- `real(rprec), dimension(:, :), allocatable realspace::ru`
- `real(rprec), dimension(:, :), allocatable realspace::rv`
- `real(rprec), dimension(:, :), allocatable, target realspace::z1`
- `real(rprec), dimension(:, :), allocatable realspace::zu`
- `real(rprec), dimension(:, :), allocatable realspace::zv`
- `real(rprec), dimension(:, :), allocatable realspace::rcon`
- `real(rprec), dimension(:, :), allocatable realspace::zcon`
- `real(rprec), dimension(:), allocatable realspace::guu`
- `real(rprec), dimension(:), allocatable realspace::guv`
- `real(rprec), dimension(:), allocatable realspace::gvv`
- `real(rprec), dimension(:), allocatable realspace::ru0`
- `real(rprec), dimension(:), allocatable realspace::zu0`
- `real(rprec), dimension(:), allocatable realspace::gcon`
- `real(rprec), dimension(:), allocatable realspace::rcon0`
- `real(rprec), dimension(:), allocatable realspace::zcon0`
- `real(rprec), dimension(:), allocatable realspace::phip`
radial derivative of $\phi/(2\pi)$ on half-grid
- `real(rprec), dimension(:), allocatable realspace::chip`
radial derivative of $\chi/(2\pi)$ on half-grid
- `real(rprec), dimension(:), allocatable realspace::shalf`
 \sqrt{s} , two-dimensional array on half-grid
- `real(rprec), dimension(:), allocatable realspace::sqrts`
 \sqrt{s} , two-dimensional array on full-grid
- `real(rprec), dimension(:), allocatable realspace::wint`
two-dimensional array for normalizing angle integrations
- `real(rprec), dimension(:, :), allocatable, target realspace::extra1`
- `real(rprec), dimension(:, :), allocatable, target realspace::extra2`
- `real(rprec), dimension(:, :), allocatable, target realspace::extra3`
- `real(rprec), dimension(:, :), allocatable, target realspace::extra4`

7.14 src/data/stel_constants.f90 File Reference

Variables

- `real(dp), parameter stel_constants::pi = 3.14159265358979323846264338328_dp`
- `real(dp), parameter stel_constants::pio2 = pi/2`
- `real(dp), parameter stel_constants::twopi = 2*pi`
- `real(dp), parameter stel_constants::sqrt2 = 1.41421356237309504880168872_dp`
- `real(dp), parameter stel_constants::degree = twopi / 360`
- `real(dp), parameter stel_constants::one = 1`
- `real(dp), parameter stel_constants::zero = 0`
- `real(dp), parameter stel_constants::mu0 = 2 * twopi * 1.0e-7_dp`

7.15 src/data/stel_kinds.f90 File Reference

Variables

- integer, parameter **stel_kinds::rprec** = SELECTED_REAL_KIND(12, 100)
- integer, parameter **stel_kinds::iprec** = SELECTED_INT_KIND(8)
- integer, parameter **stel_kinds::cprec** = KIND((1.0_rprec, 1.0_rprec))
- integer, parameter **stel_kinds::dp** = rprec

7.16 src/data/vforces.f90 File Reference

Variables

- real(rprec), dimension(:), allocatable, target **vforces::armn**
- real(rprec), dimension(:), allocatable, target **vforces::azmn**
- real(rprec), dimension(:), allocatable, target **vforces::brmn**
- real(rprec), dimension(:), allocatable, target **vforces::bzm**
- real(rprec), dimension(:), allocatable, target **vforces::blmn**
- real(rprec), dimension(:), allocatable, target **vforces::crmn**
- real(rprec), dimension(:), allocatable, target **vforces::czmn**
- real(rprec), dimension(:), allocatable, target **vforces::clmn**
- real(rprec), dimension(:), pointer **vforces::armn_e**
- real(rprec), dimension(:), pointer **vforces::armn_o**
- real(rprec), dimension(:), pointer **vforces::azmn_e**
- real(rprec), dimension(:), pointer **vforces::azmn_o**
- real(rprec), dimension(:), pointer **vforces::brmn_e**
- real(rprec), dimension(:), pointer **vforces::brmn_o**
- real(rprec), dimension(:), pointer **vforces::bzm_e**
- real(rprec), dimension(:), pointer **vforces::bzm_o**
- real(rprec), dimension(:), pointer **vforces::blmn_e**
- real(rprec), dimension(:), pointer **vforces::blmn_o**
- real(rprec), dimension(:), pointer **vforces::crmn_e**
- real(rprec), dimension(:), pointer **vforces::crmn_o**
- real(rprec), dimension(:), pointer **vforces::czmn_e**
- real(rprec), dimension(:), pointer **vforces::czmn_o**
- real(rprec), dimension(:), pointer **vforces::clmn_e**
- real(rprec), dimension(:), pointer **vforces::clmn_o**

7.17 src/data/vmec_dim.f90 File Reference

Variables

- integer **vmec_dim::mpol1**
- integer **vmec_dim::ntor1**
- integer **vmec_dim::mnmax**
- integer **vmec_dim::ntheta1**
- integer **vmec_dim::ntheta2**
- integer **vmec_dim::ntheta3**
- integer **vmec_dim::nznt**
- integer **vmec_dim::nrzt**
- integer **vmec_dim::mns**
- integer **vmec_dim::mnsize**
- integer **vmec_dim::mnmax_nyq**
- integer **vmec_dim::ns**
- integer **vmec_dim::ns1**
- integer **vmec_dim::ns_maxval**

7.18 src/data/vmec_input.f90 File Reference

Functions/Subroutines

- subroutine **vmec_input::read_indata_namelist** (iunit, istat)
- subroutine **vmec_input::write_indata_namelist** (iunit, istat)

Variables

- integer, parameter **vmec_input::mpol_default** = 6
- integer, parameter **vmec_input::ntor_default** = 0
- integer, parameter **vmec_input::ns_default** = 31
- integer, parameter **vmec_input::niter_default** = 100
- real(rprec), parameter **vmec_input::ftol_default** = 1.E-10_dp
- integer **vmec_input::nfp**
- integer **vmec_input::ncurr**
- integer **vmec_input::nstep**
- integer **vmec_input::nvacskip**
- integer **vmec_input::mpol**
- integer **vmec_input::ntor**
- integer **vmec_input::ntheta**
- integer **vmec_input::nzeta**
- integer **vmec_input::mfilter_fbdy**
- integer **vmec_input::nfilter_fbdy**
- integer, dimension(100) **vmec_input::ns_array**
- integer, dimension(100) **vmec_input::niter_array**
- real(rprec), dimension(100) **vmec_input::ftol_array**
- real(rprec), dimension(-ntord:ntord, 0:mpol1d) **vmec_input::rbc**
- real(rprec), dimension(-ntord:ntord, 0:mpol1d) **vmec_input::zbs**
- real(rprec), dimension(-ntord:ntord, 0:mpol1d) **vmec_input::rbs**
- real(rprec), dimension(-ntord:ntord, 0:mpol1d) **vmec_input::zbc**
- real(rprec) **vmec_input::curtor**
- real(rprec) **vmec_input::delt**
- real(rprec) **vmec_input::tcon0**
- real(rprec) **vmec_input::gamma**
- real(rprec) **vmec_input::bloat**
- real(rprec) **vmec_input::pres_scale**
- real(rprec) **vmec_input::spres_ped**
value of s beyond which pressure profile is flat (pedestal)
- real(rprec) **vmec_input::phiedge**
value of real toroidal flux at plasma edge ($s=1$)
- real(rprec), dimension(0:20) **vmec_input::am**
array of coefficients in phi-series for mass (NWT/m^{**2})
- real(rprec), dimension(0:20) **vmec_input::ai**
array of coefficients in phi-series for iota ($ncurr=0$)
- real(rprec), dimension(0:20) **vmec_input::ac**
array of coefficients in phi-series for the quantity $d(lcurv)/ds = \text{toroidal current density} * Vprime$, so $lcurv(s) = ltor(s)$ (used for $ncurr=1$)
- real(rprec), dimension(1:20) **vmec_input::aphi**
- character(len=20) **vmec_input::pcurr_type**
- character(len=20) **vmec_input::piota_type**
- character(len=20) **vmec_input::pmass_type**

- real(rprec), dimension(ndatafmax) **vmec_input::am_aux_s**
- real(rprec), dimension(ndatafmax) **vmec_input::am_aux_f**
- real(rprec), dimension(ndatafmax) **vmec_input::ai_aux_s**
- real(rprec), dimension(ndatafmax) **vmec_input::ai_aux_f**
- real(rprec), dimension(ndatafmax) **vmec_input::ac_aux_s**
- real(rprec), dimension(ndatafmax) **vmec_input::ac_aux_f**
- real(rprec), dimension(0:ntord) **vmec_input::raxis_cc**
- real(rprec), dimension(0:ntord) **vmec_input::raxis_cs**
- real(rprec), dimension(0:ntord) **vmec_input::zaxis_cc**
- real(rprec), dimension(0:ntord) **vmec_input::zaxis_cs**
- real(rprec), dimension(nigroup) **vmec_input::extcur**
- logical **vmec_input::lfreeb**
- logical **vmec_input::lasym**
- logical **vmec_input::lbsubs**
- character(len=200) **vmec_input::mgrid_file**
- character(len=100) **vmec_input::input_extension**

7.19 src/data/vmec_io.f90 File Reference

Variables

- real(rprec) **vmec_io::volavgb**
- real(rprec) **vmec_io::ionlarmor**
- real(rprec) **vmec_io::aminor_p**
- real(rprec) **vmec_io::rmajor_p**
- real(rprec) **vmec_io::betatot**
- real(rprec) **vmec_io::betapol**
- real(rprec) **vmec_io::betator**
- real(rprec) **vmec_io::betaxis**
- real(rprec) **vmec_io::b0**
- real(rprec) **vmec_io::volume_p**
- real(rprec) **vmec_io::cross_area_p**
- real(rprec) **vmec_io::surf_area_p**
- real(rprec) **vmec_io::circum_p**
- real(rprec) **vmec_io::kappa_p**
- real(rprec) **vmec_io::rmax_surf**
- real(rprec) **vmec_io::rmin_surf**
- real(rprec) **vmec_io::zmax_surf**

7.20 src/data/vmec_main.f90 File Reference

Variables

- real(rprec), dimension(:,:), allocatable **vmec_main::ard**
- real(rprec), dimension(:,:), allocatable **vmec_main::arm**
- real(rprec), dimension(:,:), allocatable **vmec_main::brd**
- real(rprec), dimension(:,:), allocatable **vmec_main::brm**
- real(rprec), dimension(:,:), allocatable **vmec_main::azd**
- real(rprec), dimension(:,:), allocatable **vmec_main::azm**
- real(rprec), dimension(:,:), allocatable **vmec_main::bzd**

- `real(rprec), dimension(:, :,), allocatable vmec_main::bzm`
- `real(rprec), dimension(:, :,), allocatable vmec_main::bmin`
- `real(rprec), dimension(:, :,), allocatable vmec_main::bmax`
- `real(rprec), dimension(:,), allocatable vmec_main::crd`
- `real(rprec), dimension(:,), allocatable vmec_main::iotaf`
- `real(rprec), dimension(:,), allocatable vmec_main::phipf`
- `real(rprec), dimension(:,), allocatable vmec_main::chipf`
- `real(rprec), dimension(:,), allocatable vmec_main::phi`
- `real(rprec), dimension(:,), allocatable vmec_main::beta_vol`
- `real(rprec), dimension(:,), allocatable vmec_main::jcuru`
- `real(rprec), dimension(:,), allocatable vmec_main::jcurv`
- `real(rprec), dimension(:,), allocatable vmec_main::jdotb`
- `real(rprec), dimension(:,), allocatable vmec_main::buco`
- `real(rprec), dimension(:,), allocatable vmec_main::bvco`
- `real(rprec), dimension(:,), allocatable vmec_main::bdotgradv`
- `real(rprec), dimension(:,), allocatable vmec_main::equif`
- `real(rprec), dimension(:,), allocatable vmec_main::specw`
- `real(rprec), dimension(:,), allocatable vmec_main::tcon`
- `real(rprec), dimension(:,), allocatable vmec_main::psi`
- `real(rprec), dimension(:,), allocatable vmec_main::yellip`
- `real(rprec), dimension(:,), allocatable vmec_main::yinden`
- `real(rprec), dimension(:,), allocatable vmec_main::ytrian`
- `real(rprec), dimension(:,), allocatable vmec_main::yshift`
- `real(rprec), dimension(:,), allocatable vmec_main::ygeo`
- `real(rprec), dimension(:,), allocatable vmec_main::overr`
- `real(rprec), dimension(:,), allocatable vmec_main::sm`
- `real(rprec), dimension(:,), allocatable vmec_main::sp`
- `real(rprec), dimension(:,), allocatable vmec_main::pres`
- `real(rprec), dimension(:,), allocatable vmec_main::vp`
- `real(rprec), dimension(:,), allocatable vmec_main::jpar2`
- `real(rprec), dimension(:,), allocatable vmec_main::jperp2`
- `real(rprec), dimension(:,), allocatable vmec_main::bdotb`
- `real(rprec), dimension(:,), allocatable vmec_main::blam`
- `real(rprec), dimension(:,), allocatable vmec_main::clam`
- `real(rprec), dimension(:,), allocatable vmec_main::dlam`
- `real(rprec), dimension(:,), allocatable vmec_main::vpphi`
- `real(rprec), dimension(:,), allocatable vmec_main::presgrad`
- `real(rprec), dimension(:,), allocatable vmec_main::bdamp`
- `real(rprec), dimension(:,), allocatable vmec_main::bucof`
- `real(rprec), dimension(:,), allocatable vmec_main::bvcof`
- `real(rprec), dimension(:,), allocatable vmec_main::chi`
- `real(rprec), dimension(:,), allocatable vmec_main::presf`
*pressure profile on full-grid, mass/phi**gamma*
- `real(rprec), dimension(:,), allocatable vmec_main::chips`
poloidal flux (same as chip), one-dimensional array
- `real(rprec), dimension(:,), allocatable vmec_main::phips`
toroidal flux (same as phip), one-dimensional array
- `real(rprec), dimension(:,), allocatable vmec_main::iotas`
rotational transform , on half radial mesh
- `real(rprec), dimension(:,), allocatable vmec_main::icurv`
(-)toroidal current inside flux surface (vanishes like s)
- `real(rprec), dimension(:,), allocatable vmec_main::mass`
mass profile on half-grid
- `real(rprec), dimension(:, :, :,), allocatable vmec_main::faclam`

- real(rprec), dimension(:,:,:), allocatable **vmec_main::faclam0**
- real(rprec), dimension(:,:), allocatable **vmec_main::bsqsav**
- real(rprec), dimension(:), allocatable **vmec_main::bledge**
- real(rprec), dimension(:), allocatable **vmec_main::bpedge**
- real(rprec), dimension(:), allocatable **vmec_main::bzedge**
- real(rprec), dimension(:), allocatable **vmec_main::xcl0**
- real(rprec), dimension(0:mpol1d, 3) **vmec_main::xmpq**
- real(rprec), dimension(0:mpol1d) **vmec_main::faccon**
- real(rprec) **vmec_main::hs**
radial mesh size increment
- real(rprec) **vmec_main::currv**
- real(rprec) **vmec_main::aspect**
- real(rprec) **vmec_main::ohs**
- real(rprec) **vmec_main::voli**
- real(rprec) **vmec_main::r00**
- real(rprec) **vmec_main::r0scale**
- real(rprec) **vmec_main::z00**
- real(rprec) **vmec_main::fsqsum0**
- real(rprec) **vmec_main::fnorm**
- real(rprec) **vmec_main::fsqr** =1
- real(rprec) **vmec_main::fsqz** =1
- real(rprec) **vmec_main::fsql** =1
- real(rprec) **vmec_main::fnorm1**
- real(rprec) **vmec_main::fnorml**
- real(rprec) **vmec_main::fsqr1**
- real(rprec) **vmec_main::fsqz1**
- real(rprec) **vmec_main::fsql1**
- real(rprec) **vmec_main::fsq**
- real(rprec) **vmec_main::fedge**
- real(rprec) **vmec_main::wb**
- real(rprec) **vmec_main::wp**
- real(rprec) **vmec_main::router**
- real(rprec) **vmec_main::rinner**
- real(rprec) **vmec_main::ftolv**
- real(rprec) **vmec_main::otav**
time-step algorithm
- real(rprec), dimension(ndamp) **vmec_main::otau**
- real(rprec), dimension(:,:,:), allocatable, target **vmec_main::rmn_bdy**
- real(rprec), dimension(:,:,:), allocatable, target **vmec_main::zmn_bdy**
- real(rprec), dimension(:), allocatable **vmec_main::bsub0**
- real(rprec), dimension(:), allocatable **vmec_main::dbsq**
- real(rprec), dimension(:), allocatable **vmec_main::rbsq**
- real(rprec) **vmec_main::rbtor**
- real(rprec) **vmec_main::rbtor0**
- real(rprec) **vmec_main::ctor**
- real(rprec) **vmec_main::delbsq**
- real(rprec) **vmec_main::res0**
- real(rprec) **vmec_main::delt0r**
- real(rprec), dimension(ndatafmax) **vmec_main::spfa**
- real(rprec), dimension(ndatafmax) **vmec_main::spfa2**
- real(rprec), dimension(ndatafmax) **vmec_main::hp**
- real(rprec), dimension(ndatafmax) **vmec_main::sifa**
- real(rprec), dimension(ndatafmax) **vmec_main::sifa2**
- real(rprec), dimension(ndatafmax) **vmec_main::hi**

- logical **vmec_main::lthreed**
- logical **vmec_main::lconm1**
- logical **vmec_main::lflip**
from init_geometry
- integer, dimension(:), allocatable **vmec_main::ireflect**
two-dimensional array for computing 2pi-v angle
- integer **vmec_main::multi_ns_grid**
- integer **vmec_main::itfsq**
- integer **vmec_main::ndatap**
- integer **vmec_main::ndatai**
- integer **vmec_main::niter**
max iterations for current multi-grid iteration
- integer **vmec_main::neqs**
total number of equations to evolve (size of xc)
- integer **vmec_main::irzloff**
offset in xc array between R,Z,L components
- integer **vmec_main::iequi**
counter used to call -EQFOR- at end of run
- integer **vmec_main::ijacob**
counter for number of times jacobian changes sign
- integer **vmec_main::irst**
"counter" monitoring sign of jacobian; resets R, Z, and Lambda when jacobian changes sign and decreases time step
- integer **vmec_main::iter1**
number of iterations at which the currently active evolution was branched off from
- integer **vmec_main::iter2**
total number of iterations
- integer **vmec_main::ivac**
counts number of free-boundary iterations
- integer **vmec_main::vacuum_calls** = 0

7.21 src/data/vmec_params.f90 File Reference

Variables

- integer, parameter **vmec_params::meven** = 0
parity selection label for even poloidal modes of R and Z
- integer, parameter **vmec_params::modd** = 1
parity selection label for odd poloidal modes of R and Z
- integer, parameter **vmec_params::ndamp** = 10
number of iterations over which damping is averaged
- integer, parameter **vmec_params::ns4** = 25
- integer, dimension(0:mpold), parameter **vmec_params::jmin1** = (/ 1,1,(2,ink=2,mpold) /)
starting js(m) values where R,Z are non-zero
- integer, dimension(0:mpold), parameter **vmec_params::jmin2** = (/ 1,2,(2,ink=2,mpold) /)
starting js(m) values for which R,Z are evolved
- integer, dimension(0:mpold), parameter **vmec_params::jlam** = (/ 2,2,(2,ink=2,mpold) /)
starting js(m) values for which Lambda is evolved
- integer, parameter **vmec_params::norm_term_flag** = 0
- integer, parameter **vmec_params::bad_jacobian_flag** = 1
- integer, parameter **vmec_params::jac75_flag** = 4

- integer, parameter **vmec_params::input_error_flag** = 5
- integer, parameter **vmec_params::phiedge_error_flag** = 7
- integer, parameter **vmec_params::ns_error_flag** = 8
- integer, parameter **vmec_params::misc_error_flag** = 9
- integer, parameter **vmec_params::successful_term_flag** = 11
- integer, parameter **vmec_params::restart_flag** = 1
- integer, parameter **vmec_params::readin_flag** = 2
- integer, parameter **vmec_params::timestep_flag** = 4
- integer, parameter **vmec_params::output_flag** = 8
- integer, parameter **vmec_params::cleanup_flag** = 16
- integer, parameter **vmec_params::reset_jacdt_flag** = 32
- real(rprec), parameter **vmec_params::pdamp** = 0.05_dp
- character(len= *), parameter **vmec_params::version_** = '8.52'
- integer **vmec_params::ntmax**
number of contributing Fourier basis function (can be 1, 2 or 4); assigned in read_indata()
- integer **vmec_params::rcc**
- integer **vmec_params::rss**
- integer **vmec_params::rsc**
- integer **vmec_params::rcs**
- integer **vmec_params::zsc**
- integer **vmec_params::zcs**
- integer **vmec_params::zcc**
- integer **vmec_params::zss**
- integer **vmec_params::mnyq**
- integer **vmec_params::nnyq**
- integer, dimension(:), allocatable **vmec_params::uminus**
- real(rprec), dimension(:), allocatable **vmec_params::mscale**
*array for norming theta-trig functions (internal use only) so that the discrete SUM[cos(mu)*cos(m'u)] = .5 delta(m,m')*
- real(rprec), dimension(:), allocatable **vmec_params::nscale**
array for norming zeta -trig functions (internal use only)
- real(rprec) **vmec_params::signgs**
sign of Jacobian : must be =1 (right-handed) or =-1 (left-handed)
- real(rprec) **vmec_params::lamscale** =1
- integer, parameter **vmec_params::m0** =0
from totzsp
- integer, parameter **vmec_params::m1** =1
from totzsp
- integer, parameter **vmec_params::n0** =0
from totzsp

7.22 src/data/vmec_persistent.f90 File Reference

Variables

- integer, dimension(:), allocatable **vmec_persistent::ixm**
- integer, dimension(:), allocatable **vmec_persistent::jmin3**
- real(rprec), dimension(:, :), allocatable **vmec_persistent::cosmu**
- real(rprec), dimension(:, :), allocatable **vmec_persistent::sinmu**
- real(rprec), dimension(:, :), allocatable **vmec_persistent::cosmum**
- real(rprec), dimension(:, :), allocatable **vmec_persistent::sinmum**
- real(rprec), dimension(:, :), allocatable **vmec_persistent::cosmumi**

- `real(rprec), dimension(:, :), allocatable vmec_persistent::sinmumi`
- `real(rprec), dimension(:, :), allocatable vmec_persistent::cosnv`
- `real(rprec), dimension(:, :), allocatable vmec_persistent::sinnv`
- `real(rprec), dimension(:, :), allocatable vmec_persistent::cosnvn`
- `real(rprec), dimension(:, :), allocatable vmec_persistent::sinnavn`
- `real(rprec), dimension(:, :), allocatable vmec_persistent::cosmui`
- `real(rprec), dimension(:, :), allocatable vmec_persistent::sinmui`
- `real(rprec), dimension(:, :), allocatable vmec_persistent::cosmui3`
- `real(rprec), dimension(:, :), allocatable vmec_persistent::cosmumi3`
- `real(rprec), dimension(:), allocatable, target vmec_persistent::xm`
- `real(rprec), dimension(:), allocatable, target vmec_persistent::xn`
- `real(rprec), dimension(:), allocatable, target vmec_persistent::xm_nyq`
- `real(rprec), dimension(:), allocatable, target vmec_persistent::xn_nyq`
- `real(rprec), dimension(:), allocatable vmec_persistent::cos01`
- `real(rprec), dimension(:), allocatable vmec_persistent::sin01`

7.23 src/data/vmercier.f90 File Reference

Variables

- `real(rprec), dimension(nsd) vmmercier::dshear`
- `real(rprec), dimension(nsd) vmmercier::dwell`
- `real(rprec), dimension(nsd) vmmercier::dcurr`
- `real(rprec), dimension(nsd) vmmercier::dmerc`
- `real(rprec), dimension(nsd) vmmercier::dgeod`

7.24 src/data/vparams.f90 File Reference

Variables

- integer, parameter `vparams::nsd` = 10001
maximum number of radial nodes
- integer, parameter `vparams::mpold` = 101
maximum number of poloidal harmonics (in r,z,lam fourier series)
- integer, parameter `vparams::ntord` = 101
maximum number of toroidal harmonics
- integer, parameter `vparams::ndatafmax` = 101
- integer, parameter `vparams::nstore_seq` = 100
- integer, parameter `vparams::mpol1d` = mpold - 1
- integer, parameter `vparams::ntor1d` = ntord + 1
- integer, parameter `vparams::nthreed0` = 9
- integer, parameter `vparams::indata0` = nthreed0 + 2
- integer, parameter `vparams::nwout0` = nthreed0 + 3
- integer, parameter `vparams::jxbout0` = nthreed0 + 4
- integer, parameter `vparams::nfort18` = 18
- integer, parameter `vparams::nmercier0` = 52
- integer `vparams::nthreed`
- `real(rprec), parameter vparams::c1pm2` = 1.e-2_dp
- `real(rprec), parameter vparams::cp15` = 0.15_dp
- `real(rprec), parameter vparams::cp25` = 0.25_dp

- real(rprec), parameter **vparams::cp5** = 0.50_dp
- real(rprec), parameter **vparams::c1pm8** = 1.0e-8_dp
- real(rprec), parameter **vparams::cbig** = 0.9e30_dp
- real(rprec), parameter **vparams::c2p0** = 2
- real(rprec), parameter **vparams::c3p0** = 3
- real(rprec), parameter **vparams::cp05** = 0.05_dp
- real(rprec), parameter **vparams::c1pm13** = 1.0e-13_dp
- real(rprec), parameter **vparams::osqrt2** = 0.707106781186547462_dp

7.25 src/data/vsvd0.f90 File Reference

Variables

- integer, parameter **vsvd0::nigroup** = 100
number of external current groups

7.26 src/data/xstuff.f90 File Reference

Variables

- real(rprec), dimension(:), allocatable **xstuff::gc**
stacked array of R, Z, Lambda Spectral force coefficients (see above for stack order)
- real(rprec), dimension(:), allocatable, target **xstuff::xc**
stacked array of scaled R, Z, Lambda Fourier coefficients (see above for stack order)
- real(rprec), dimension(:), allocatable **xstuff::xcdot**
"velocity": change of Fourier coefficients per time step
- real(rprec), dimension(:), allocatable **xstuff::xsave**
- real(rprec), dimension(:), allocatable **xstuff::xstore**
backup copy of last-known-good xc
- real(rprec), dimension(:), allocatable **xstuff::scalxc**

7.27 src/elongation.f90 File Reference

Compute Waist thickness and height in $\varphi = 0, \pi$ symmetry planes.

Functions/Subroutines

- subroutine **elongation** (r1, z1, waist, height)
Compute Waist thickness and height in $\varphi = 0, \pi$ symmetry planes.

7.27.1 Detailed Description

Compute Waist thickness and height in $\varphi = 0, \pi$ symmetry planes.

7.27.2 Function/Subroutine Documentation

7.27.2.1 elongation()

```
subroutine elongation (
    real(rprec), dimension(ns,nzeta,ntheta3,0:1), intent(in) r1,
    real(rprec), dimension(ns,nzeta,ntheta3,0:1), intent(in) z1,
    real(rprec), dimension(2), intent(out) waist,
    real(rprec), dimension(2), intent(out) height )
```

Compute Waist thickness and height in $\varphi = 0, \pi$ symmetry planes.

Parameters

<i>r1</i>	<i>R</i>
<i>z1</i>	<i>Z</i>
<i>waist</i>	
<i>height</i>	

Definition at line 10 of file elongation.f90.

Referenced by eqfor().

Here is the caller graph for this function:



7.28 src/eqfor.f90 File Reference

Basis physics analysis and evaluation of force balance. This is where most of the contents of the `threed1` output file is computed.

Functions/Subroutines

- subroutine [eqfor](#) (br, bz, bsubu, bsubv, tau, rzl_array, ier_flag)

Basis physics analysis and evaluation of force balance. This is where most of the contents of the `threed1` output file is computed.

7.28.1 Detailed Description

Basis physics analysis and evaluation of force balance. This is where most of the contents of the `threed1` output file is computed.

7.28.2 Function/Subroutine Documentation

7.28.2.1 eqfor()

```
subroutine eqfor (
    real(rprec), dimension(nrzt), intent(out) br,
    real(rprec), dimension(nrzt), intent(out) bz,
    real(rprec), dimension(ns,nznt,0:1), intent(in) bsubu,
    real(rprec), dimension(ns,nznt,0:1), intent(in) bsubv,
    real(rprec), dimension(nrzt), intent(out) tau,
    real(rprec), dimension(ns,0:ntor,0:mpoll,3*ntmax), intent(in), target rzl_array,
    integer ier_flag )
```

Basis physics analysis and evaluation of force balance. This is where most of the contents of the `threed1` output file is computed.

Parameters

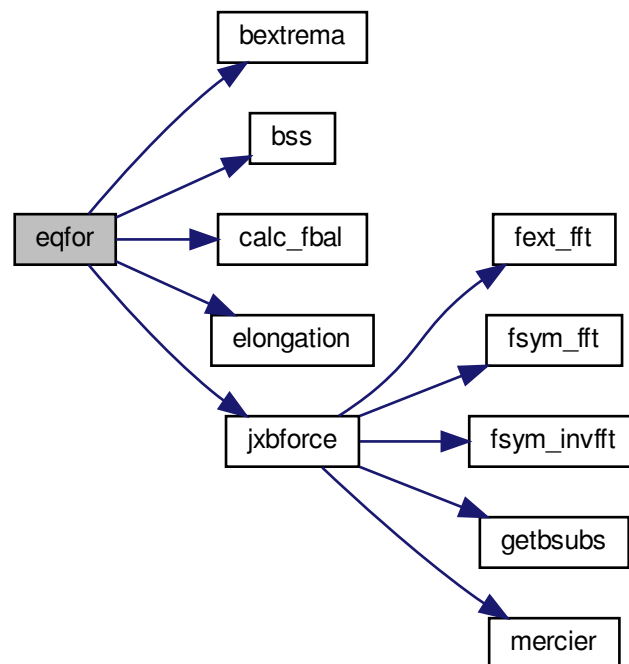
<i>br</i>	cylindrical component of magnetic field B^R
<i>bz</i>	cylindrical component of magnetic field B^Z
<i>bsubu</i>	covariant component of magnetic field B_θ
<i>bsubv</i>	covariant component of magnetic field B_ζ
<i>tau</i>	Jacobian $\sqrt{g} = R\tau$
<i>rzl_array</i>	state vector (all Fourier coefficients) of VMEC
<i>ier_flag</i>	error flag

Definition at line 15 of file eqfor.f90.

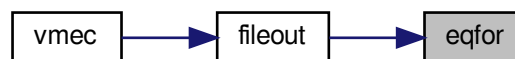
References `bextrema()`, `bss()`, `calc_fbal()`, `elongation()`, and `jxbforce()`.

Referenced by `fileout()`.

Here is the call graph for this function:



Here is the caller graph for this function:



7.29 src/eqsolve.f90 File Reference

Iteratively evolve the Fourier coefficients that specify the equilibrium.

Functions/Subroutines

- subroutine [eqsolve](#) (ier_flag)

Iteratively evolve the Fourier coefficients that specify the equilibrium.

7.29.1 Detailed Description

Iteratively evolve the Fourier coefficients that specify the equilibrium.

7.29.2 Function/Subroutine Documentation

7.29.2.1 eqsolve()

```
subroutine eqsolve (
    integer, intent(inout) ier_flag )
```

Iteratively evolve the Fourier coefficients that specify the equilibrium.

Parameters

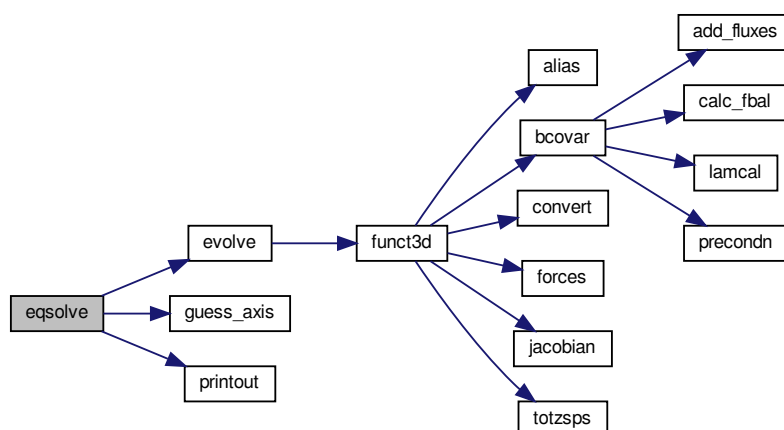
<i>ier_flag</i>	error flag
-----------------	------------

Definition at line 7 of file eqsolve.f90.

References `evolve()`, `guess_axis()`, and `printout()`.

Referenced by `vmec()`.

Here is the call graph for this function:



Here is the caller graph for this function:



7.30 src/evolve.f90 File Reference

Take a single time step in Fourier space to evolve the Fourier coefficients describing the equilibrium towards force balance.

Functions/Subroutines

- subroutine [evolve](#) (time_step, ier_flag, liter_flag)

Take a single time step in Fourier space to evolve the Fourier coefficients describing the equilibrium towards force balance.

7.30.1 Detailed Description

Take a single time step in Fourier space to evolve the Fourier coefficients describing the equilibrium towards force balance.

7.30.2 Function/Subroutine Documentation

7.30.2.1 evolve()

```

subroutine evolve (
    real(rprec), intent(in) time_step,
    integer, intent(inout) ier_flag,
    logical, intent(inout) liter_flag )
  
```

Take a single time step in Fourier space to evolve the Fourier coefficients describing the equilibrium towards force balance.

Parameters

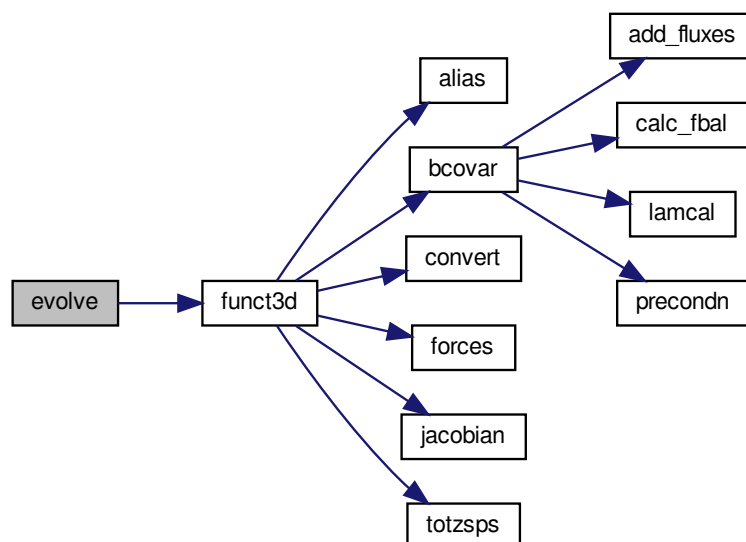
<i>time_step</i>	step length in parameter space to take
<i>ier_flag</i>	error flag
<i>liter_flag</i>	keep running?

Definition at line 11 of file evolve.f90.

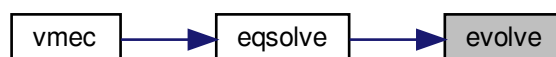
References `funct3d()`.

Referenced by `eqsolve()`.

Here is the call graph for this function:



Here is the caller graph for this function:



7.31 src/fileout.f90 File Reference

Write the output files.

Functions/Subroutines

- subroutine `fileout` (`ier_flag`)
Write the output files.

7.31.1 Detailed Description

Write the output files.

7.31.2 Function/Subroutine Documentation

7.31.2.1 fileout()

```
subroutine fileout (  
    integer, intent(inout) ier_flag )
```

Write the output files.

Parameters

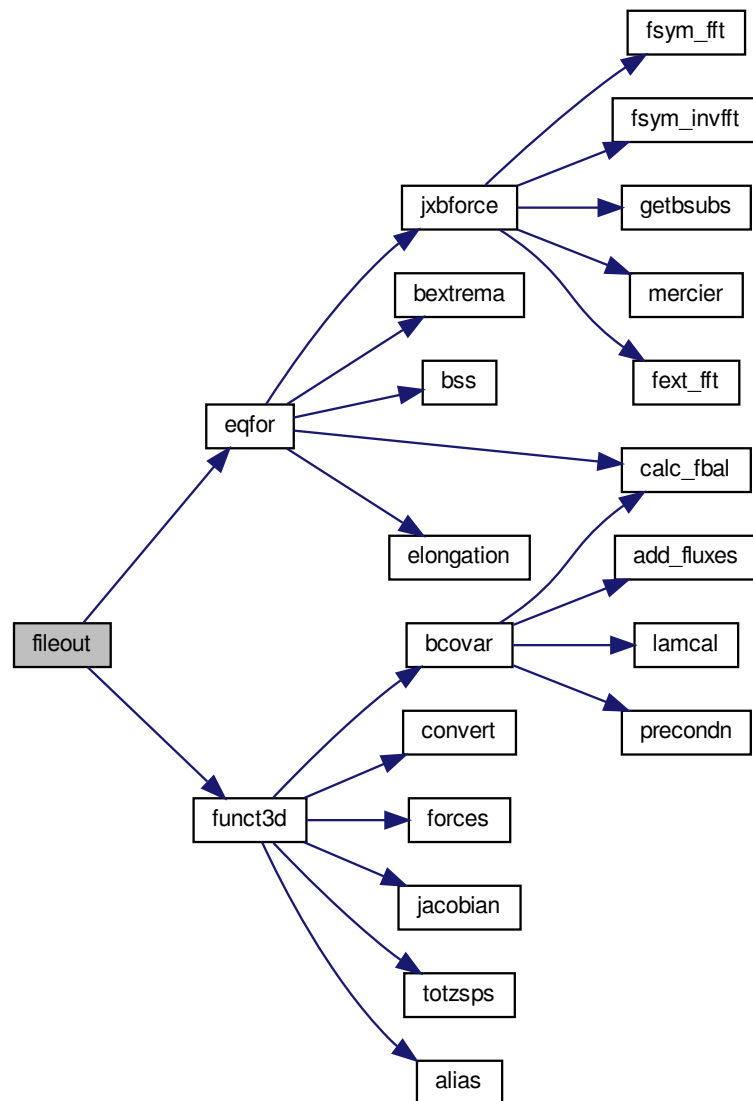
<i>ier_flag</i>	error flag
-----------------	------------

Definition at line 7 of file fileout.f90.

References `eqfor()`, and `funct3d()`.

Referenced by `vmec()`.

Here is the call graph for this function:



Here is the caller graph for this function:



7.32 src/fixaray.f90 File Reference

allocate and fill some fixed-size arrays (only depending on Fourier resolution).

Functions/Subroutines

- subroutine [fixaray](#)
allocate and fill some fixed-size arrays (only depending on Fourier resolution).

7.32.1 Detailed Description

allocate and fill some fixed-size arrays (only depending on Fourier resolution).

7.33 src/flip_theta.f90 File Reference

Flip the definition of the poloidal angle in the user-provided initial guess for the LCFS geometry.

Functions/Subroutines

- subroutine [flip_theta](#) (rmn, zmn, lmn)
Flip the definition of the poloidal angle in the user-provided initial guess for the LCFS geometry.

7.33.1 Detailed Description

Flip the definition of the poloidal angle in the user-provided initial guess for the LCFS geometry.

7.33.2 Function/Subroutine Documentation

7.33.2.1 flip_theta()

```
subroutine flip_theta (
    real(rprec), dimension(0:ntor,0:mpoll,ntmax), intent(inout) rmn,
    real(rprec), dimension(0:ntor,0:mpoll,ntmax), intent(inout) zmn,
    real(rprec), dimension(0:ntor,0:mpoll,ntmax), intent(inout), optional lmn )
```

Flip the definition of the poloidal angle in the user-provided initial guess for the LCFS geometry.

Parameters

	<i>rmn</i>	Fourier coefficients for R
	<i>zmn</i>	Fourier coefficients for Z
	<i>lmn</i>	Fourier coefficients for λ
<i>in, out</i>	<i>lmn</i>	never used: can also flip lambda...

Definition at line 9 of file flip_theta.f90.

7.34 src/forces.f90 File Reference

Compute the real-space MHD forces.

Functions/Subroutines

- subroutine [forces](#)
Compute the real-space MHD forces.

7.34.1 Detailed Description

Compute the real-space MHD forces.

7.35 src/free_mem_func3d.f90 File Reference

Free memory required by [func3d\(\)](#)

Functions/Subroutines

- subroutine [free_mem_func3d](#)
Free memory required by [func3d\(\)](#)

7.35.1 Detailed Description

Free memory required by [func3d\(\)](#)

7.36 src/free_mem_ns.f90 File Reference

Free memory depending on the number of flux surfaces *ns*.

Functions/Subroutines

- subroutine [free_mem_ns](#)
*Free memory depending on the number of flux surfaces *ns*.*

7.36.1 Detailed Description

Free memory depending on the number of flux surfaces *ns*.

7.37 src/free_mem_nunv.f90 File Reference

Free arrays depending on the number of Fourier coefficients `nunv`.

Functions/Subroutines

- subroutine `free_mem_nunv`
Free arrays depending on the number of Fourier coefficients `nunv`.

7.37.1 Detailed Description

Free arrays depending on the number of Fourier coefficients `nunv`.

7.38 src/freeb_data.f90 File Reference

Write out edge values of fields.

Functions/Subroutines

- subroutine `freeb_data` (`rmnc`, `zmns`, `rmns`, `zmnc`, `bmodmn`, `bmodmn1`)
Write out edge values of fields.

7.38.1 Detailed Description

Write out edge values of fields.

7.38.2 Function/Subroutine Documentation

7.38.2.1 freeb_data()

```
subroutine freeb_data (
    real(rprec), dimension(mnmax) rmnc,
    real(rprec), dimension(mnmax) zmns,
    real(rprec), dimension(mnmax) rmns,
    real(rprec), dimension(mnmax) zmnc,
    real(rprec), dimension(mnmax) bmodmn,
    real(rprec), dimension(mnmax) bmodmn1 )
```

Write out edge values of fields.

Parameters

<i>rmnc</i>	stellarator-symmetric Fourier coefficients of R
<i>zmns</i>	stellarator-symmetric Fourier coefficients of Z
<i>rmns</i>	non-stellarator-symmetric Fourier coefficients of R
<i>zmnc</i>	non-stellarator-symmetric Fourier coefficients of Z
<i>bmodmn</i>	stellarator-symmetric Fourier coefficients of $ \mathbf{B} $
<i>bmodmn1</i>	non-stellarator-symmetric Fourier coefficients of $ \mathbf{B} $

Definition at line 12 of file freeb_data.f90.

7.39 src/fsym_fft.f90 File Reference

Fourier transforms.

Functions/Subroutines

- subroutine `fext_fft` (bout, bs_s, bs_a)
Extends B_s from $n\theta_{a2}$ interval to full $n\theta_{a3}$ interval in angle θ .
- subroutine `fsym_fft` (bs, bu, bv, bs_s, bu_s, bv_s, bs_a, bu_a, bv_a)
Contract bs,bu,bv from full nu interval to half- u interval so cos, sin integrals can be performed on half- u interval.

7.39.1 Detailed Description

Fourier transforms.

7.39.2 Function/Subroutine Documentation

7.39.2.1 `fext_fft()`

```
subroutine fext_fft (
    real(rprec), dimension(nzeta,ntheta3), intent(out) bout,
    real(rprec), dimension(nzeta,ntheta2), intent(in) bs_s,
    real(rprec), dimension(nzeta,ntheta2), intent(in) bs_a )
```

Extends B_s from $n\theta_{a2}$ interval to full $n\theta_{a3}$ interval in angle θ .

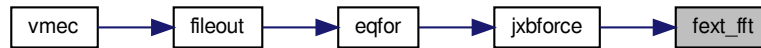
Parameters

<i>bout</i>	output B_s
<i>bs_s</i>	symmetric part of B_s
<i>bs_a</i>	anti-symmetric part of B_s

Definition at line 9 of file fsym_fft.f90.

Referenced by jxbforce().

Here is the caller graph for this function:



7.39.2.2 fsym_fft()

```

subroutine fsym_fft (
    real(rprec), dimension(nzeta,ntheta3), intent(in) bs,
    real(rprec), dimension(nzeta,ntheta3,0:1), intent(in) bu,
    real(rprec), dimension(nzeta,ntheta3,0:1), intent(in) bv,
    real(rprec), dimension(nzeta,ntheta2) bs_s,
    real(rprec), dimension(nzeta,ntheta2,0:1), intent(out) bu_s,
    real(rprec), dimension(nzeta,ntheta2,0:1), intent(out) bv_s,
    real(rprec), dimension(nzeta,ntheta2) bs_a,
    real(rprec), dimension(nzeta,ntheta2,0:1), intent(out) bu_a,
    real(rprec), dimension(nzeta,ntheta2,0:1), intent(out) bv_a )

```

Contract bs,bu,bv from full ν_u interval to half-u interval so cos, sin integrals can be performed on half-u interval.

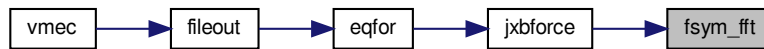
Parameters

<i>bs</i>	output B_s
<i>bu</i>	output B_θ
<i>bv</i>	output B_{zeta}
<i>bs</i> ↔ <i>_s</i>	symmetric part of B_s
<i>bu</i> ↔ <i>_s</i>	symmetric part of B_θ
<i>bv</i> ↔ <i>_s</i>	symmetric part of B_ζ
<i>bs</i> ↔ <i>_a</i>	anti-symmetric part of B_s
<i>bu</i> ↔ <i>_a</i>	anti-symmetric part of B_θ
<i>bv</i> ↔ <i>_a</i>	anti-symmetric part of B_ζ

Definition at line 47 of file fsym_fft.f90.

Referenced by jxbforce().

Here is the caller graph for this function:



7.40 src/fsym_invfft.f90 File Reference

Extends function from `ntheta2` to `ntheta3` range.

Functions/Subroutines

- subroutine `fsym_invfft` (`bsubs_u`, `bsubs_v`)
Extends function from `ntheta2` to `ntheta3` range.

7.40.1 Detailed Description

Extends function from `ntheta2` to `ntheta3` range.

7.40.2 Function/Subroutine Documentation

7.40.2.1 `fsym_invfft()`

```

subroutine fsym_invfft (
    real(rprec), dimension(ns*nzeta,ntheta3,0:1), intent(inout) bsubs_u,
    real(rprec), dimension(ns*nzeta,ntheta3,0:1), intent(inout) bsubs_v )

```

Extends function from `ntheta2` to `ntheta3` range.

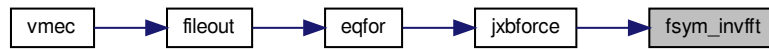
Parameters

<code>bsubs_u</code>	tangential derivative of covariant magnetic field component $\partial B_s / \partial \theta$
<code>bsubs_v</code>	tangential derivative of covariant magnetic field component $\partial B_s / \partial \zeta$

Definition at line 8 of file `fsym_invfft.f90`.

Referenced by `jxbforce()`.

Here is the caller graph for this function:



7.41 src/funct3d.f90 File Reference

Evaluate the three-dimensional MHD energy functional.

Functions/Subroutines

- subroutine `funct3d` (`ier_flag`)
Evaluate the three-dimensional MHD energy functional.

7.41.1 Detailed Description

Evaluate the three-dimensional MHD energy functional.

7.41.2 Function/Subroutine Documentation

7.41.2.1 `funct3d()`

```

subroutine funct3d (
    integer, intent(inout) ier_flag )

```

Evaluate the three-dimensional MHD energy functional.

Parameters

<code>ier_flag</code>	error flag
-----------------------	------------

use system call to stand-alone NESTOR for vacuum computation

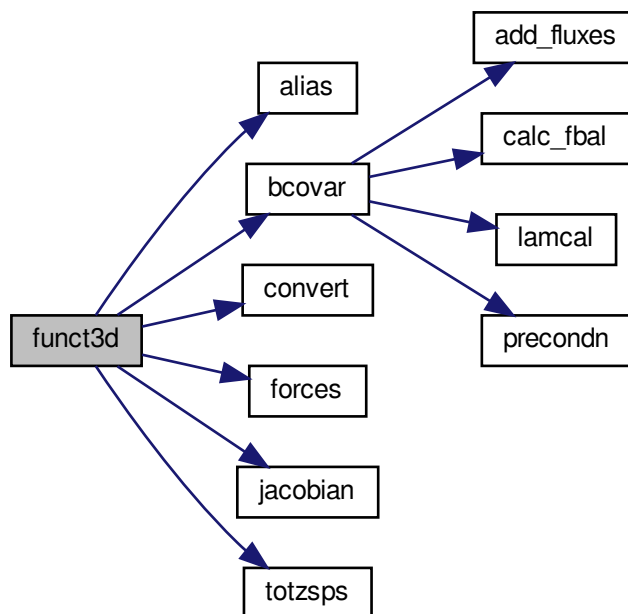
dump reference input for and output of NESTOR when using internal NESTOR

Definition at line 7 of file `funct3d.f90`.

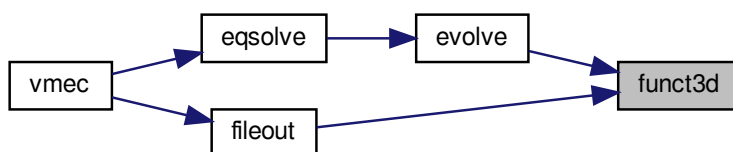
References `alias()`, `bcovar()`, `convert()`, `forces()`, `jacobian()`, and `totzsps()`.

Referenced by evolve(), and fileout().

Here is the call graph for this function:



Here is the caller graph for this function:



7.42 src/functions.f File Reference

This module contains functions used by the profiles.

Functions/Subroutines

- real(rprec) function, public `functions::two_power` (x, b)
*Profile function for the two_power profile. $b(0) * (1 - x^{b(1)})^{b(2)}$.*
- real(rprec) function, public `functions::two_power_gs` (x, b)
*Profile function for the two_power_gs profile. $two_power(x) * (1 + \sum [b(i) * \exp(-(x - b(i + 1))/b(i + 2))^2])$.*
- logical function `functions::function_test` ()
Main test function.

7.42.1 Detailed Description

This module contains functions used by the profiles.

7.42.2 Function/Subroutine Documentation

7.42.2.1 function_test()

logical function `functions::function_test`

Main test function.

Test `two_power` function for x = 0, b = {1,10,2} is 1

Test `two_power` function for x = 1, b = {1,10,2} is 0

Test `two_power` function for x = 0.5, b = {1,1,1} is 0.5

Test `two_power` function for x = 0.5, b = {1,1,2} is 0.25

Test `two_power_gs` function for x = 0.4, b = {1,1,1,0,0,1} is `two_power(x,b)`

Test `two_power_gs` function for x = 0.8, b = {1,1,0,1,0.8,0.1} is 2

Definition at line 51 of file `functions.f`.

7.42.2.2 two_power()

```
real(rprec) function, public functions::two_power (
    real(rprec), intent(in) x,
    real(rprec), dimension(0:20), intent(in) b )
```

Profile function for the `two_power` profile. $b(0) * (1 - x^{b(1)})^{b(2)}$.

Parameters

x	evaluation location
b	parameter vector

Definition at line 20 of file functions.f.

7.42.2.3 two_power_gs()

```
real(rprec) function, public functions::two_power_gs (
    real(rprec), intent(in) x,
    real(rprec), dimension(0:20), intent(in) b )
```

Profile function for the `two_power_gs` profile. $\text{two_power}(x) * (1 + \sum [b(i) * \exp(-(x - b(i + 1))/b(i + 2))^2])$.

Parameters

x	evaluation location
b	parameter vector

Definition at line 34 of file functions.f.

7.43 src/getbsubs.f90 File Reference

Solves the radial force balance $\mathbf{B} \cdot B_s = F_s$ for B_s in real space using collocation.

Functions/Subroutines

- subroutine [getbsubs](#) (bsubsmn, frho, bsupu, bsupv, mmax, nmax, info)
Solves the radial force balance $\mathbf{B} \cdot B_s = F_s$ for B_s in real space using collocation.

7.43.1 Detailed Description

Solves the radial force balance $\mathbf{B} \cdot B_s = F_s$ for B_s in real space using collocation.

7.43.2 Function/Subroutine Documentation

7.43.2.1 getbsubs()

```

subroutine getbsubs (
    real(rprec), dimension(0:mmax, -nmax:nmax, 0:1), intent(out) bsubsmn,
    real(rprec), dimension(nzeta, ntheta3), intent(in) frho,
    real(rprec), dimension(nzeta, ntheta3), intent(in) bsupu,
    real(rprec), dimension(nzeta, ntheta3), intent(in) bsupv,
    integer, intent(in) mmax,
    integer, intent(in) nmax,
    integer, intent(out) info )

```

Solves the radial force balance $\mathbf{B} \cdot B_s = F_s$ for B_s in real space using collocation.

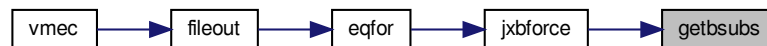
Parameters

<i>bsubsmn</i>	Fourier coefficients of B_s
<i>frho</i>	Fourier coefficients of radial Force component
<i>bsupu</i>	contravariant component of magnetic field B^θ
<i>bsupv</i>	contravariant component of magnetic field B^ζ
<i>mmax</i>	maximum poloidal mode number
<i>nmax</i>	maximum toroidal mode number
<i>info</i>	error flag

Definition at line 13 of file getbsubs.f90.

Referenced by jxbforce().

Here is the caller graph for this function:



7.44 src/getcurmid.f90 File Reference

Get current at midplane (?)

Functions/Subroutines

- subroutine [getcurmid](#) (curmid, izeta, gsqrt, r12)
Get current at midplane (?)

7.44.1 Detailed Description

Get current at midplane (?)

7.44.2 Function/Subroutine Documentation

7.44.2.1 getcurmid()

```
subroutine getcurmid (
    real(rprec), dimension(2*ns) curmid,
    real(rprec), dimension(ns,nzeta,*) izeta,
    real(rprec), dimension(ns,nzeta,*) gsqrt,
    real(rprec), dimension(ns,nzeta,*) r12 )
```

Get current at midplane (?)

Parameters

<i>curmid</i>	current at midplane (?)
<i>izeta</i>	index in toroidal direction
<i>gsqrt</i>	Jacobian
<i>r12</i>	R^2

Definition at line 10 of file getcurmid.f90.

7.45 src/getfsq.f90 File Reference

Compute total force residual on flux surfaces.

Functions/Subroutines

- subroutine [getfsq](#) (gcr, gcx, gnormr, gnormz, gnorn, medge)
Compute total force residual on flux surfaces.

7.45.1 Detailed Description

Compute total force residual on flux surfaces.

7.45.2 Function/Subroutine Documentation

7.45.2.1 getfsq()

```
subroutine getfsq (
    real(rprec), dimension(ns,mnsize*ntmax), intent(in) gcr,
    real(rprec), dimension(ns,mnsize*ntmax), intent(in) gcx,
    real(rprec), intent(out) gnormr,
    real(rprec), intent(out) gnormz,
    real(rprec), intent(in) gnorn,
    integer, intent(in) medge )
```

Compute total force residual on flux surfaces.

Parameters

<i>gcr</i>	<i>R</i> -component of force
<i>gcz</i>	<i>Z</i> -component of force
<i>gnormr</i>	normalized total force residual in <i>R</i>
<i>gnormz</i>	normalized total force residual in <i>Z</i>
<i>gnorm</i>	normalization factor for forces
<i>medge</i>	=0: exclude contribution from LCFS; =1: include LCFS contribution

Definition at line 12 of file getfsq.f90.

7.46 src/guess_axis.f90 File Reference

Computes guess for magnetic axis if user guess leads to initial sign change of Jacobian.

Functions/Subroutines

- subroutine [guess_axis](#) (*r1*, *z1*, *ru0*, *zu0*)
Computes guess for magnetic axis if user guess leads to initial sign change of Jacobian.

7.46.1 Detailed Description

Computes guess for magnetic axis if user guess leads to initial sign change of Jacobian.

7.46.2 Function/Subroutine Documentation

7.46.2.1 guess_axis()

```
subroutine guess_axis (
    real (rprec), dimension (ns,nzeta,ntheta3,0:1), intent (in) r1,
    real (rprec), dimension (ns,nzeta,ntheta3,0:1), intent (in) z1,
    real (rprec), dimension (ns,nzeta,ntheta3), intent (in) ru0,
    real (rprec), dimension (ns,nzeta,ntheta3), intent (in) zu0 )
```

Computes guess for magnetic axis if user guess leads to initial sign change of Jacobian.

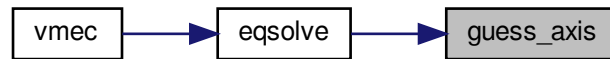
Parameters

<i>r1</i>	<i>R</i>
<i>z1</i>	<i>Z</i>
<i>ru0</i>	$\partial R / \partial \theta$
<i>zu0</i>	$\partial Z / \partial \theta$

Definition at line 10 of file guess_axis.f90.

Referenced by eqsolve().

Here is the caller graph for this function:



7.47 src/heading.f90 File Reference

Open output files and print banner message at the top.

Functions/Subroutines

- subroutine [heading](#) (extension)
Open output files and print banner message at the top.

7.47.1 Detailed Description

Open output files and print banner message at the top.

7.47.2 Function/Subroutine Documentation

7.47.2.1 heading()

```

subroutine heading (
    character(len=*), intent(in) extension )

```

Open output files and print banner message at the top.

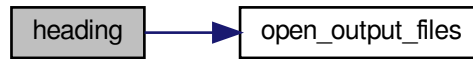
Parameters

<i>extension</i>	input file "extension": part after 'input.'.
------------------	--

Definition at line 7 of file heading.f90.

References `open_output_files()`.

Here is the call graph for this function:



7.48 src/initialize_radial.f90 File Reference

Allocates memory for radial arrays and initializes radial profiles.

Functions/Subroutines

- subroutine [initialize_radial](#) (`nsval`, `ns_old`, `delt0`)
Allocates memory for radial arrays and initializes radial profiles.

7.48.1 Detailed Description

Allocates memory for radial arrays and initializes radial profiles.

7.48.2 Function/Subroutine Documentation

7.48.2.1 initialize_radial()

```

subroutine initialize_radial (
    integer, intent(in) nsval,
    integer, intent(inout) ns_old,
    real(rprec), intent(out) delt0 )
  
```

Allocates memory for radial arrays and initializes radial profiles.

Parameters

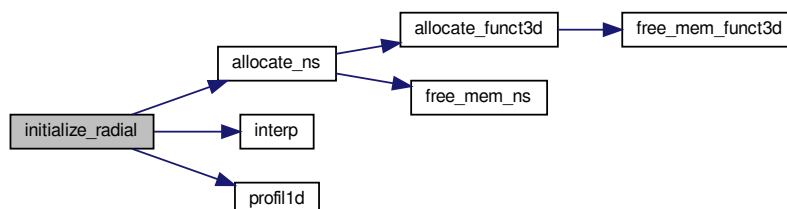
<i>nsval</i>	new number of flux surfaces
<i>ns_old</i>	old number of flux surfaces (from previous multi-grid iteration)
<i>delt0</i>	time step to be used in the new multi-grid iteration

Definition at line 9 of file initialize_radial.f90.

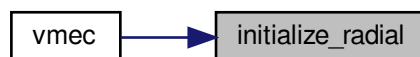
References `allocate_ns()`, `interp()`, and `profil1d()`.

Referenced by `vmec()`.

Here is the call graph for this function:



Here is the caller graph for this function:



7.49 src/interp.f90 File Reference

Interpolate R , Z and λ on full grid.

Functions/Subroutines

- subroutine `interp` (xnew, xold, scalxc, nsnew, nsold)
Interpolate R , Z and λ on full grid.

7.49.1 Detailed Description

Interpolate R , Z and λ on full grid.

7.49.2 Function/Subroutine Documentation

7.49.2.1 interp()

```

subroutine interp (
    real(rprec), dimension(nsnew,mnsize,3*ntmax), intent(out) xnew,
    real(rprec), dimension(nsold,mnsize,3*ntmax), intent(inout) xold,
    real(rprec), dimension(nsnew,mnsize,3*ntmax), intent(in) scalxc,
    integer, intent(in) nsnew,
    integer, intent(in) nsold )

```

Interpolate R , Z and λ on full grid.

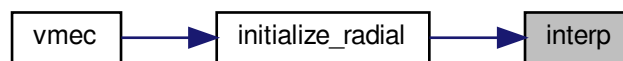
Parameters

<i>xnew</i>	interpolated state vector (nsnew surfaces)
<i>xold</i>	interpolation basis: old state vector (nsold surfaces)
<i>scalxc</i>	scaling factors to normalize the new state vector to
<i>nsnew</i>	new number of flux surfaces
<i>nsold</i>	old number of flux surfaces

Definition at line 11 of file interp.f90.

Referenced by initialize_radial().

Here is the caller graph for this function:



7.50 src/jacobian.f90 File Reference

Evaluate the Jacobian of the transform from flux- to cylindrical coordinates.

Functions/Subroutines

- subroutine [jacobian](#)
Evaluate the Jacobian of the transform from flux- to cylindrical coordinates.

7.50.1 Detailed Description

Evaluate the Jacobian of the transform from flux- to cylindrical coordinates.

7.51 src/jxbforce.f90 File Reference

Program for computing local $\mathbf{K} \times \mathbf{B} = \nabla p$ force balance.

Functions/Subroutines

- subroutine [jxbforce](#) (bsupu, bsupv, bsubu, bsubv, bsubsh, bsubsu, bsubsv, gsqrt, bsq, itheta, izeta, brho, ier_flag)

Program for computing local $\mathbf{K} \times \mathbf{B} = \nabla p$ force balance.

7.51.1 Detailed Description

Program for computing local $\mathbf{K} \times \mathbf{B} = \nabla p$ force balance.

7.51.2 Function/Subroutine Documentation

7.51.2.1 jxbforce()

```
subroutine jxbforce (
    real(rprec), dimension(ns,nznt), intent(in) bsupu,
    real(rprec), dimension(ns,nznt), intent(in) bsupv,
    real(rprec), dimension(ns,nznt,0:1), intent(inout), target bsubu,
    real(rprec), dimension(ns,nznt,0:1), intent(inout), target bsubv,
    real(rprec), dimension(ns,nznt), intent(in) bsubsh,
    real(rprec), dimension(ns,nznt,0:1) bsubsu,
    real(rprec), dimension(ns,nznt,0:1) bsubsv,
    real(rprec), dimension(ns,nznt), intent(in) gsqrt,
    real(rprec), dimension(ns,nznt), intent(in) bsq,
    real(rprec), dimension(ns,nznt), intent(out) itheta,
    real(rprec), dimension(ns,nznt), intent(out) izeta,
    real(rprec), dimension(ns,nznt), intent(out) brho,
    integer, intent(in) ier_flag )
```

Program for computing local $\mathbf{K} \times \mathbf{B} = \nabla p$ force balance.

Parameters

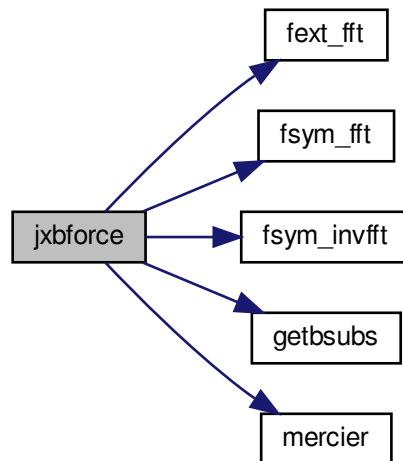
<i>bsupu</i>	contravariant component of magnetic field B^θ
<i>bsupv</i>	contravariant component of magnetic field B^ζ
<i>bsubu</i>	covariant component of magnetic field B_θ
<i>bsubv</i>	covariant component of magnetic field B_ζ
<i>bsubsh</i>	covariant component of magnetic field B_s (on half grid?)
<i>bsubsu</i>	tangential derivate of covariant component of magnetic field $\partial B_s / \partial \theta$ (?)
<i>bsubsv</i>	tangential derivate of covariant component of magnetic field $\partial B_s / \partial \zeta$ (?)
<i>gsqrt</i>	Jacobian \sqrt{g}
<i>bsq</i>	modulus of magnetic field $ \mathbf{B} ^2$
<i>itheta</i>	index in poloidal direction
<i>izeta</i>	index in toroidal direction
<i>brho</i>	radial component of magnetic field B_ρ (?)
<i>ier_flag</i>	error flag

Definition at line 19 of file jxbforce.f90.

References `fext_fft()`, `fsym_fft()`, `fsym_invfft()`, `getbsubs()`, and `mercier()`.

Referenced by `eqfor()`.

Here is the call graph for this function:



Here is the caller graph for this function:



7.52 src/lamcal.f90 File Reference

Normalization parameters for λ .

Functions/Subroutines

- subroutine `lamcal` (`overg`, `guu`, `guv`, `gvv`)
Normalization parameters for λ .

7.52.1 Detailed Description

Normalization parameters for λ .

7.52.2 Function/Subroutine Documentation

7.52.2.1 lamcal()

```
subroutine lamcal (
    real(rprec), dimension(ns,nznt), intent(in) overg,
    real(rprec), dimension(ns,nznt), intent(in) guu,
    real(rprec), dimension(ns,nznt), intent(in) guv,
    real(rprec), dimension(ns,nznt), intent(in) gvv )
```

Normalization parameters for λ .

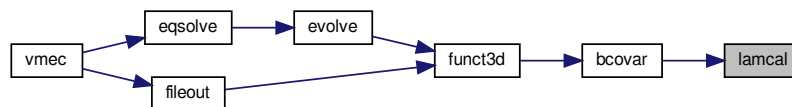
Parameters

<i>overg</i>	inverse of Jacobian $1/\sqrt{g}$
<i>guu</i>	metric element $g_{\theta\theta}$
<i>guv</i>	metric element $g_{\theta\zeta}$
<i>gvv</i>	metric element $g_{\zeta\zeta}$

Definition at line 10 of file lamcal.f90.

Referenced by bcovar().

Here is the caller graph for this function:



7.53 src/line_segment.f File Reference

This module contains code to create a profile constructed of line segments.

Modules

- module [line_segment](#)

This module contains code to create a profile constructed of line segments. These line segments are assumed to be specified such that $xx(i) < xx(i + 1)$.

Functions/Subroutines

- subroutine, public `line_segment::line_seg` (x, y, xx, yy, n)
- subroutine, public `line_segment::line_seg_int` (x, y, xx, yy, n)
- logical function, public `line_segment::line_seg_test` ()

7.53.1 Detailed Description

This module contains code to create a profile constructed of line segments.

7.54 src/magnetic_fluxes.f90 File Reference

Compute toroidal and poloidal magnetic flux profiles.

Functions/Subroutines

- real(rprec) function `torflux_deriv` (x)
Compute the radial derivative of the enclosed toroidal magnetic flux.
- real(rprec) function `polflux_deriv` (x)
Compute the radial derivative of the enclosed poloidal magnetic flux.
- real(rprec) function `torflux` (x)
Compute the enclosed toroidal magnetic flux.
- real(rprec) function `polflux` (x)
Compute the enclosed poloidal magnetic flux.

7.54.1 Detailed Description

Compute toroidal and poloidal magnetic flux profiles.

7.54.2 Function/Subroutine Documentation

7.54.2.1 polflux()

```
real(rprec) function polflux (  
    real(rprec), intent(in) x )
```

Compute the enclosed poloidal magnetic flux.

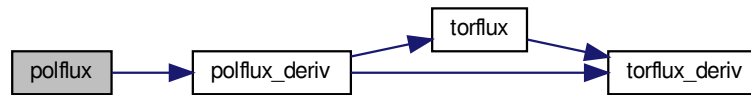
Parameters

	x	evaluation location
in	x	radial flux variable (=TOROIDAL FLUX ONLY IF APHI=1)

Definition at line 75 of file magnetic_fluxes.f90.

References polflux_deriv().

Here is the call graph for this function:



7.54.2.2 polflux_deriv()

```
real(rprec) function polflux_deriv (
    real(rprec), intent(in) x )
```

Compute the radial derivative of the enclosed poloidal magnetic flux.

Parameters

	<i>x</i>	evaluation location
<i>in</i>	<i>x</i>	radial flux variable (=TOROIDAL FLUX ONLY IF APhi=1)

Returns

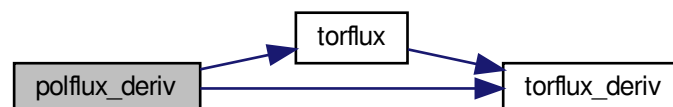
$\text{polflux_deriv} == d(\chi)/dx = \text{iota}(\text{TF}(x)) * \text{torflux_deriv}(x)$

Definition at line 28 of file magnetic_fluxes.f90.

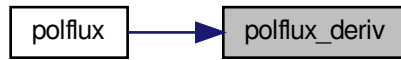
References torflux(), and torflux_deriv().

Referenced by polflux().

Here is the call graph for this function:



Here is the caller graph for this function:



7.54.2.3 torflux()

```

real(rprec) function torflux (
    real(rprec), intent(in) x )
  
```

Compute the enclosed toroidal magnetic flux.

Parameters

	<i>x</i>	evaluation location
<i>in</i>	<i>x</i>	radial flux variable (=TOROIDAL FLUX ONLY IF APhi=1)

Definition at line 51 of file magnetic_fluxes.f90.

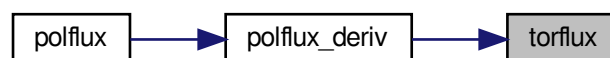
References torflux_deriv().

Referenced by polflux_deriv().

Here is the call graph for this function:



Here is the caller graph for this function:



7.54.2.4 torflux_deriv()

```
real(rprec) function torflux_deriv (
    real(rprec), intent(in) x )
```

Compute the radial derivative of the enclosed toroidal magnetic flux.

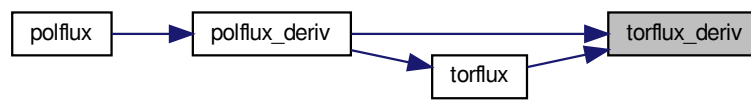
Parameters

	<i>x</i>	evaluation location
<i>in</i>	<i>x</i>	radial flux variable (=TOROIDAL FLUX ONLY IF APhi=1)

Definition at line 7 of file magnetic_fluxes.f90.

Referenced by polflux_deriv(), and torflux().

Here is the caller graph for this function:



7.55 src/mercier.f90 File Reference

Evaluate the Mercier stability criterion.

Functions/Subroutines

- subroutine [mercier](#) (gsqrt, bsq, bdotj, iotas, wint, r1, rt, rz, zt, zz, bsubu, vp, phips, pres, ns, nznt)
Evaluate the Mercier stability criterion.

7.55.1 Detailed Description

Evaluate the Mercier stability criterion.

7.55.2 Function/Subroutine Documentation

7.55.2.1 mercier()

```

subroutine mercier (
    real(rprec), dimension(ns,nznt), intent(in) gsqrt,
    real(rprec), dimension(ns,nznt), intent(in) bsq,
    real(rprec), dimension(ns,nznt), intent(inout) bdotj,
    real(rprec), dimension(ns), intent(in) iotas,
    real(rprec), dimension(ns*nznt), intent(in) wint,
    real(rprec), dimension(ns,nznt,0:1), intent(in) r1,
    real(rprec), dimension(ns,nznt,0:1), intent(in) rt,
    real(rprec), dimension(ns,nznt,0:1), intent(in) rz,
    real(rprec), dimension(ns,nznt,0:1), intent(in) zt,
    real(rprec), dimension(ns,nznt,0:1), intent(in) zz,
    real(rprec), dimension(ns*nznt), intent(in) bsubu,
    real(rprec), dimension(ns), intent(in) vp,
    real(rprec), dimension(ns), intent(in) phips,
    real(rprec), dimension(ns), intent(in) pres,
    integer, intent(in) ns,
    integer, intent(in) nznt )

```

Evaluate the Mercier stability criterion.

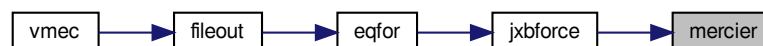
Parameters

<i>gsqrt</i>	Jacobian \sqrt{g}
<i>bsq</i>	modulus of magnetic field $ \mathbf{B} $
<i>bdotj</i>	parallel current density $\mathbf{B} \cdot \mathbf{j}$
<i>iotas</i>	rotational transform profile
<i>wint</i>	normalization constant for flux-surface integrals
<i>r1</i>	R
<i>rt</i>	$\partial R / \partial \theta$
<i>rz</i>	$\partial R / \partial \zeta$
<i>zt</i>	$\partial Z / \partial \theta$
<i>zz</i>	$\partial Z / \partial \zeta$
<i>bsubu</i>	contravariant component of magnetic field B^ζ
<i>vp</i>	radial profile of specific volume $\partial V / \partial s$
<i>phips</i>	radial derivative of enclosed toroidal magnetic flux
<i>pres</i>	pressure profile
<i>ns</i>	number of flux surfaces
<i>nznt</i>	number of grid points per flux surface

Definition at line 22 of file mercier.f90.

Referenced by jxbforce().

Here is the caller graph for this function:



7.56 src/mgrid_mod.f File Reference

Precomputed table of magnetic field due to confinement coils.

Modules

- module [mgrid_mod](#)
Precomputed table of magnetic field due to confinement coils.

Functions/Subroutines

- subroutine **mgrid_mod::read_mgrid** (mgrid_file, extcur, nv, nfp, lscreen, ier_flag)
- subroutine **mgrid_mod::sum_bfield** (bfield, bf_add, cur, n1)
- subroutine **mgrid_mod::assign_bptrs** (bptr)
- subroutine **mgrid_mod::free_mgrid** (istat)

Variables

- integer, parameter **mgrid_mod::nlimset** = 2
- character(len= *), parameter **mgrid_mod::vn_br0** = 'br'
- character(len= *), parameter **mgrid_mod::vn_bp0** = 'bp'
- character(len= *), parameter **mgrid_mod::vn_bz0** = 'bz'
- character(len= *), parameter **mgrid_mod::vn_ir** = 'ir'
- character(len= *), parameter **mgrid_mod::vn_jz** = 'jz'
- character(len= *), parameter **mgrid_mod::vn_kp** = 'kp'
- character(len= *), parameter **mgrid_mod::vn_nfp** = 'nfp'
- character(len= *), parameter **mgrid_mod::vn_rmin** = 'rmin'
- character(len= *), parameter **mgrid_mod::vn_rmax** = 'rmax'
- character(len= *), parameter **mgrid_mod::vn_zmin** = 'zmin'
- character(len= *), parameter **mgrid_mod::vn_zmax** = 'zmax'
- character(len= *), parameter **mgrid_mod::vn_coilgrp** = 'coil_group'
- character(len= *), parameter **mgrid_mod::vn_nextcur** = 'nextcur'
- character(len= *), parameter **mgrid_mod::vn_mgmode** = 'mgrid_mode'
- character(len= *), parameter **mgrid_mod::vn_coilcur** = 'raw_coil_cur'
- character(len= *), parameter **mgrid_mod::ln_next** = 'External currents'
- integer **mgrid_mod::nr0b**
- integer **mgrid_mod::np0b**
- integer **mgrid_mod::nfper0**
- integer **mgrid_mod::nz0b**
- integer **mgrid_mod::nobd**
- integer **mgrid_mod::nobser**
- integer **mgrid_mod::nextcur**
- integer **mgrid_mod::nbfldn**
- integer **mgrid_mod::nbsets**
- integer **mgrid_mod::nbcoilsn**
- integer **mgrid_mod::nbvac**
- integer **mgrid_mod::nbcoil_max**
- integer **mgrid_mod::nlim**
- integer **mgrid_mod::nlim_max**
- integer **mgrid_mod::nsets**
- integer **mgrid_mod::nrgrid**

- integer **mgrid_mod::nzgrid**
- integer, dimension(:), allocatable **mgrid_mod::needflx**
- integer, dimension(:), allocatable **mgrid_mod::nbcoils**
- integer, dimension(:), allocatable **mgrid_mod::limitr**
- integer, dimension(:), allocatable **mgrid_mod::nsetsn**
- integer, dimension(:,:), allocatable **mgrid_mod::iconnect**
- integer, dimension(:,:), allocatable **mgrid_mod::needbfld**
- real(rprec) **mgrid_mod::rminb**
- real(rprec) **mgrid_mod::zminb**
- real(rprec) **mgrid_mod::rmaxb**
- real(rprec) **mgrid_mod::zmaxb**
- real(rprec) **mgrid_mod::delrb**
- real(rprec) **mgrid_mod::delzb**
- real(rprec) **mgrid_mod::rx1**
- real(rprec) **mgrid_mod::rx2**
- real(rprec) **mgrid_mod::zy1**
- real(rprec) **mgrid_mod::zy2**
- real(rprec) **mgrid_mod::condif**
- real(rprec), dimension(:,:), allocatable, target **mgrid_mod::bvac**
- real(rprec), dimension(:,,:), pointer **mgrid_mod::brvac**
- real(rprec), dimension(:,,:), pointer **mgrid_mod::bzvac**
- real(rprec), dimension(:,,:), pointer **mgrid_mod::bpvac**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::unpsiext**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::plbfld**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::rbcoil**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::zbcoil**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::abcoil**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::bcoil**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::rbcoilsqr**
- real(rprec), dimension(:), allocatable **mgrid_mod::raw_coil_current**
- real(rprec), dimension(:), allocatable **mgrid_mod::xobser**
- real(rprec), dimension(:), allocatable **mgrid_mod::zobser**
- real(rprec), dimension(:), allocatable **mgrid_mod::xobsqr**
- real(rprec), dimension(:), allocatable **mgrid_mod::dsiext**
- real(rprec), dimension(:), allocatable **mgrid_mod::psiext**
- real(rprec), dimension(:), allocatable **mgrid_mod::plflux**
- real(rprec), dimension(:), allocatable **mgrid_mod::b_chi**
- character(len=300) **mgrid_mod::mgrid_path**
- character(len=300) **mgrid_mod::mgrid_path_old** = " "
- character(len=30), dimension(:), allocatable **mgrid_mod::curlabel**
- character(len=15), dimension(:), allocatable **mgrid_mod::dsilabel**
- character(len=15), dimension(:), allocatable **mgrid_mod::bloopnames**
- character(len=30) **mgrid_mod::tokid**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::dbcoil**
- real(rprec), dimension(:,,:), allocatable **mgrid_mod::pfcspec**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::rlim**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::zlim**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::reslim**
- real(rprec), dimension(:,:), allocatable **mgrid_mod::seplim**
- character(len=1) **mgrid_mod::mgrid_mode**

7.56.1 Detailed Description

Precomputed table of magnetic field due to confinent coils.

7.57 src/NESTOR/analysum.f90 File Reference

Functions/Subroutines

- subroutine **analysum** (grpmn, bvec, sl, tl, m, n, l, ivacskip, lasym, m_map, n_map, grpmn_m_map, grpmn↔
_n_map)

7.58 src/NESTOR/analysum2.f90 File Reference

Functions/Subroutines

- subroutine **analysum2** (grpmn, bvec, m, n, l, ivacskip, lasym, m_map, n_map, grpmn_m_map, grpmn_n↔
map)

7.59 src/NESTOR/analyt.f90 File Reference

Functions/Subroutines

- subroutine **analyt** (grpmn, bvec, ivacskip, lasym, m_map, n_map, grpmn_m_map, grpmn_n_map)

7.60 src/NESTOR/becoil.f90 File Reference

Functions/Subroutines

- subroutine **becoil** (rad, zee, brvac, bpvac, bzvac)

7.61 src/NESTOR/belicu.f90 File Reference

Functions/Subroutines

- subroutine **belicu** (torcur, bx, by, bz, cos1, sin1, rp, zp)

7.62 src/NESTOR/bextern.f90 File Reference

Functions/Subroutines

- subroutine **bextern** (plascur, wint)

7.63 src/NESTOR/data/nestor_io.f90 File Reference

Input and Output for stand-alone NESTOR.

Modules

- module `nestor_io`

Input and Output for stand-alone NESTOR.

Functions/Subroutines

- subroutine `nestor_io::read_nestor_inputs` (vac_file)
- subroutine `nestor_io::write_nestor_outputs` (vac_file, lasym, ivac, ier_flag)
- subroutine `write_nestor_inputs` (vac_file, vacuum_calls, ier_flag, mgrid_file, input_extension, ivacskip, ivac, nfp, ntor, mpol, nzeta, ntheta, mnmax, xm, xn, rmnc, zmns, rmns, zmnc, rbtor, ctor, lasym, signgs, extcur_↔ nestor, raxis_nestor, zaxis_nestor, wint, nznt, amatsav, bvecsav, mnpd2, bsubvvac)
- subroutine `read_nestor_outputs` (vac_file, ier_flag, ivac)

Variables

- character(len=255) `nestor_io::input_extension`
- character(len=255) `nestor_io::mgrid_file`
- real(dp), dimension(:), allocatable `nestor_io::extcur`
- real(dp), dimension(:), allocatable `nestor_io::raxis`
- real(dp), dimension(:), allocatable `nestor_io::zaxis`
- real(dp), dimension(:), allocatable `nestor_io::xm`
- real(dp), dimension(:), allocatable `nestor_io::xn`
- real(dp), dimension(:), allocatable `nestor_io::rmnc`
- real(dp), dimension(:), allocatable `nestor_io::zmns`
- real(dp), dimension(:), allocatable `nestor_io::rmns`
- real(dp), dimension(:), allocatable `nestor_io::zmnc`
- real(dp), dimension(:), allocatable `nestor_io::wint`
- integer `nestor_io::nfp`
- integer `nestor_io::ntor`
- integer `nestor_io::mpol`
- integer `nestor_io::ntheta`
- integer `nestor_io::nzeta`
- integer `nestor_io::nextcur`
- integer `nestor_io::ier_flag`
- integer `nestor_io::ivac`
- integer `nestor_io::ivacskip`
- integer `nestor_io::mnmax`
- integer `nestor_io::vacuum_calls`
- logical `nestor_io::lasym`
- real(dp) `nestor_io::ctor`
- real(dp) `nestor_io::rbtor`
- real(dp) `nestor_io::signgs`
- integer `nestor_io::mnpd2_nestor`
- real(dp), dimension(:), allocatable `nestor_io::amatsav_nestor`
- real(dp), dimension(:), allocatable `nestor_io::bvecsav_nestor`
- real(dp) `nestor_io::bsubvvac_nestor`
- character(len= *), dimension(1), parameter `nestor_io::mn1dim` = ('mn_mode'/)
- character(len= *), dimension(1), parameter `nestor_io::mnpotdim` = ('mn_mode_pot'/)
- character(len= *), dimension(1), parameter `nestor_io::nzntdim` = ('nznt'/)
- character(len= *), dimension(1), parameter `nestor_io::nzetadim` = ('nzeta'/)
- character(len= *), dimension(1), parameter `nestor_io::nextcurim` = ('nextcur'/)
- character(len= *), dimension(1), parameter `nestor_io::bvecsavdim` = ('mnpd2'/)

- character(len= *), dimension(1), parameter **nestor_io::amatsavdim** = (/mnpd2_times_mnpd2/)
- character(len= *), dimension(2), parameter **nestor_io::r2dim** = (/mn_mode',radius '/)
- character(len= *), parameter **nestor_io::vn_vacuum_calls** = 'vacuum_calls'
- character(len= *), parameter **nestor_io::vn_ier_flag** = "ier_flag"
- character(len= *), parameter **nestor_io::vn_mgrid** = "mgrid_file"
- character(len= *), parameter **nestor_io::vn_inputext** = "input_extension"
- character(len= *), parameter **nestor_io::vn_ivacskip** = "ivacskip"
- character(len= *), parameter **nestor_io::vn_ivac** = "ivac"
- character(len= *), parameter **nestor_io::vn_nfp** = "nfp"
- character(len= *), parameter **nestor_io::vn_ntor** = "ntor"
- character(len= *), parameter **nestor_io::vn_mpol** = "mpol"
- character(len= *), parameter **nestor_io::vn_nzeta** = "nzeta"
- character(len= *), parameter **nestor_io::vn_ntheta** = "ntheta"
- character(len= *), parameter **nestor_io::vn_mnmax** = "mnmax"
- character(len= *), parameter **nestor_io::vn_pmod** = "xm"
- character(len= *), parameter **nestor_io::vn_tmod** = "xn"
- character(len= *), parameter **nestor_io::vn_rmnc** = "rmnc"
- character(len= *), parameter **nestor_io::vn_zmns** = "zmns"
- character(len= *), parameter **nestor_io::vn_rmns** = "rmns"
- character(len= *), parameter **nestor_io::vn_zmnc** = "zmnc"
- character(len= *), parameter **nestor_io::vn_rbtor** = "rbtor"
- character(len= *), parameter **nestor_io::vn_ctor** = "ctor"
- character(len= *), parameter **nestor_io::vn_lasym** = "lasym"
- character(len= *), parameter **nestor_io::vn_signgs** = "signgs"
- character(len= *), parameter **nestor_io::vn_extcur** = "extcur"
- character(len= *), parameter **nestor_io::vn_raxis_nestor** = "raxis_nestor"
- character(len= *), parameter **nestor_io::vn_zaxis_nestor** = "zaxis_nestor"
- character(len= *), parameter **nestor_io::vn_wint** = "wint"
- character(len= *), parameter **nestor_io::vn_bsqvac** = "bsqvac"
- character(len= *), parameter **nestor_io::vn_mnpd** = "mnpd"
- character(len= *), parameter **nestor_io::vn_xmpot** = "xmpot"
- character(len= *), parameter **nestor_io::vn_xnpot** = "xnpot"
- character(len= *), parameter **nestor_io::vn_potvac** = "potvac"
- character(len= *), parameter **nestor_io::vn_brν** = "brν"
- character(len= *), parameter **nestor_io::vn_bphiv** = "bphiv"
- character(len= *), parameter **nestor_io::vn_bzv** = "bzv"
- character(len= *), parameter **nestor_io::vn_bsubvvac** = "bsubvvac"
- character(len= *), parameter **nestor_io::vn_amatsav** = "amatsav"
- character(len= *), parameter **nestor_io::vn_bvecsav** = "bvecsav"
- character(len= *), parameter **nestor_io::vn_mnpd2** = "mnpd2"
- character(len= *), parameter **nestor_io::vn_r1b** = "r1b"
- character(len= *), parameter **nestor_io::vn_rub** = "rub"
- character(len= *), parameter **nestor_io::vn_rvb** = "rvb"
- character(len= *), parameter **nestor_io::vn_z1b** = "z1b"
- character(len= *), parameter **nestor_io::vn_zub** = "zub"
- character(len= *), parameter **nestor_io::vn_zvb** = "zvb"
- character(len= *), parameter **nestor_io::vn_ruu** = "ruu"
- character(len= *), parameter **nestor_io::vn_ruv** = "ruv"
- character(len= *), parameter **nestor_io::vn_rvv** = "rvv"
- character(len= *), parameter **nestor_io::vn_zuu** = "zuu"
- character(len= *), parameter **nestor_io::vn_zuv** = "zuv"
- character(len= *), parameter **nestor_io::vn_zvv** = "zvv"
- character(len= *), parameter **nestor_io::vn_guu_b** = "guu_b"
- character(len= *), parameter **nestor_io::vn_guv_b** = "guv_b"
- character(len= *), parameter **nestor_io::vn_gvv_b** = "gvv_b"

- character(len= *), parameter **nestor_io::vn_rzb2** = "rzb2"
- character(len= *), parameter **nestor_io::vn_snr** = "snr"
- character(len= *), parameter **nestor_io::vn_snv** = "snv"
- character(len= *), parameter **nestor_io::vn_snz** = "snz"
- character(len= *), parameter **nestor_io::vn_drv** = "drv"
- character(len= *), parameter **nestor_io::vn_auu** = "auu"
- character(len= *), parameter **nestor_io::vn_auv** = "auv"
- character(len= *), parameter **nestor_io::vn_avv** = "avv"
- character(len= *), parameter **nestor_io::vn_rcosuv** = "rcosuv"
- character(len= *), parameter **nestor_io::vn_rsinuv** = "rsinuv"
- character(len= *), parameter **nestor_io::vn_brad** = "brad"
- character(len= *), parameter **nestor_io::vn_bphi** = "bphi"
- character(len= *), parameter **nestor_io::vn_bz** = "bz"
- character(len= *), parameter **nestor_io::vn_bexu** = "bexu"
- character(len= *), parameter **nestor_io::vn_bexv** = "bexv"
- character(len= *), parameter **nestor_io::vn_bexn** = "bexn"
- character(len= *), parameter **nestor_io::vn_bexni** = "bexni"
- character(len= *), parameter **nestor_io::vn_grpmn** = "grpmn"
- character(len= *), parameter **nestor_io::vn_adp** = "adp"
- character(len= *), parameter **nestor_io::vn_adm** = "adm"
- character(len= *), parameter **nestor_io::vn_cma** = "cma"
- character(len= *), parameter **nestor_io::vn_sqrtc** = "sqrtc"
- character(len= *), parameter **nestor_io::vn_sqrta** = "sqrta"
- character(len= *), parameter **nestor_io::vn_delt1u** = "delt1u"
- character(len= *), parameter **nestor_io::vn_azp1u** = "azp1u"
- character(len= *), parameter **nestor_io::vn_azm1u** = "azm1u"
- character(len= *), parameter **nestor_io::vn_cma11u** = "cma11u"
- character(len= *), parameter **nestor_io::vn_r1p** = "r1p"
- character(len= *), parameter **nestor_io::vn_r1m** = "r1m"
- character(len= *), parameter **nestor_io::vn_r0p** = "r0p"
- character(len= *), parameter **nestor_io::vn_r0m** = "r0m"
- character(len= *), parameter **nestor_io::vn_ra1p** = "ra1p"
- character(len= *), parameter **nestor_io::vn_ra1m** = "ra1m"
- character(len= *), parameter **nestor_io::vn_sqad1u** = "sqad1u"
- character(len= *), parameter **nestor_io::vn_sqad2u** = "sqad2u"
- character(len= *), parameter **nestor_io::vn_all_tlp** = "all_tlp"
- character(len= *), parameter **nestor_io::vn_all_tlm** = "all_tlm"
- character(len= *), parameter **nestor_io::vn_all_slp** = "all_slp"
- character(len= *), parameter **nestor_io::vn_all_slm** = "all_slm"
- character(len= *), parameter **nestor_io::vn_m_map** = "m_map"
- character(len= *), parameter **nestor_io::vn_n_map** = "n_map"
- character(len= *), parameter **nestor_io::vn_green** = "green"
- character(len= *), parameter **nestor_io::vn_greenp** = "greenp"
- character(len= *), parameter **nestor_io::vn_tanu** = "tanu"
- character(len= *), parameter **nestor_io::vn_tanv** = "tanv"
- character(len= *), parameter **nestor_io::vn_gstore** = "gstore"
- character(len= *), parameter **nestor_io::vn_grpmn_m_map** = "grpmn_m_map"
- character(len= *), parameter **nestor_io::vn_grpmn_n_map** = "grpmn_n_map"
- character(len= *), parameter **nestor_io::vn_imirr** = "imirr"
- character(len= *), parameter **nestor_io::vn_amatrix** = "amatrix"
- character(len= *), parameter **nestor_io::vn_potu** = "potu"
- character(len= *), parameter **nestor_io::vn_potv** = "potv"
- character(len= *), parameter **nestor_io::vn_bsubu** = "bsubu"
- character(len= *), parameter **nestor_io::vn_bsubv** = "bsubv"

7.63.1 Detailed Description

Input and Output for stand-alone NESTOR.

7.64 src/NESTOR/data/vac_persistent.f90 File Reference

Variables

- integer, dimension(:), allocatable **vac_persistent::imirr**
- real(rprec), dimension(:), allocatable **vac_persistent::sinper**
- real(rprec), dimension(:), allocatable **vac_persistent::cosper**
- real(rprec), dimension(:), allocatable **vac_persistent::sinuv**
- real(rprec), dimension(:), allocatable **vac_persistent::cosuv**
- real(rprec), dimension(:), allocatable **vac_persistent::tanu**
- real(rprec), dimension(:), allocatable **vac_persistent::tanv**
- real(rprec), dimension(:), allocatable **vac_persistent::tanu_1d**
- real(rprec), dimension(:), allocatable **vac_persistent::tanv_1d**
- real(rprec), dimension(:), allocatable **vac_persistent::xmpot**
- real(rprec), dimension(:), allocatable **vac_persistent::xnpot**
- real(rprec), dimension(:), allocatable **vac_persistent::csign**
- real(rprec), dimension(:,), allocatable **vac_persistent::sinu**
- real(rprec), dimension(:,), allocatable **vac_persistent::cosu**
- real(rprec), dimension(:,), allocatable **vac_persistent::sinv**
- real(rprec), dimension(:,), allocatable **vac_persistent::cosv**
- real(rprec), dimension(:,), allocatable **vac_persistent::sinui**
- real(rprec), dimension(:,), allocatable **vac_persistent::cosui**
- real(rprec), dimension(:,), allocatable **vac_persistent::sinu1**
- real(rprec), dimension(:,), allocatable **vac_persistent::cosu1**
- real(rprec), dimension(:,), allocatable **vac_persistent::sinv1**
- real(rprec), dimension(:,), allocatable **vac_persistent::cosv1**
- real(rprec), dimension(:,,:), allocatable **vac_persistent::cmns**
- real(rprec), dimension(:), allocatable **vac_persistent::bsubu_sur**
- real(rprec), dimension(:), allocatable **vac_persistent::bsubv_sur**
- real(rprec), dimension(:), allocatable **vac_persistent::bsupu_sur**
- real(rprec), dimension(:), allocatable **vac_persistent::bsupv_sur**

7.65 src/NESTOR/data/vacmod.f90 File Reference

Functions/Subroutines

- subroutine **vacmod::allocate_nestor**
- subroutine **vacmod::free_mem_nestor**

Variables

- real(rprec), parameter **vacmod::p5** = cp5
- real(rprec), parameter **vacmod::two** = c2p0
- real(rprec) **vacmod::bsubvvac**
- real(rprec) **vacmod::pi2**
- real(rprec) **vacmod::pi3**
- real(rprec) **vacmod::pi4**
- real(rprec) **vacmod::alp**
- real(rprec) **vacmod::alu**
- real(rprec) **vacmod::alv**
- real(rprec) **vacmod::alvp**
- real(rprec) **vacmod::onp**
- real(rprec) **vacmod::onp2**
- logical **vacmod::precal_done**
- real(rprec), dimension(:), allocatable, target **vacmod::potvac**
- real(rprec), dimension(:), allocatable **vacmod::m_map_wrt**
- real(rprec), dimension(:), allocatable **vacmod::n_map_wrt**
- real(rprec), dimension(:), allocatable **vacmod::bvecsav**
- real(rprec), dimension(:), allocatable **vacmod::amatsav**
- real(rprec), dimension(:), allocatable **vacmod::bexni**
- real(rprec), dimension(:), allocatable **vacmod::brv**
- real(rprec), dimension(:), allocatable **vacmod::bphiv**
- real(rprec), dimension(:), allocatable **vacmod::bzv**
- real(rprec), dimension(:), allocatable **vacmod::bsqvac**
- real(rprec), dimension(:), allocatable **vacmod::r1b**
- real(rprec), dimension(:), allocatable **vacmod::rub**
- real(rprec), dimension(:), allocatable **vacmod::rvb**
- real(rprec), dimension(:), allocatable **vacmod::z1b**
- real(rprec), dimension(:), allocatable **vacmod::zub**
- real(rprec), dimension(:), allocatable **vacmod::zvb**
- real(rprec), dimension(:), allocatable **vacmod::bexu**
- real(rprec), dimension(:), allocatable **vacmod::bexv**
- real(rprec), dimension(:), allocatable **vacmod::bexn**
- real(rprec), dimension(:), allocatable **vacmod::auu**
- real(rprec), dimension(:), allocatable **vacmod::auv**
- real(rprec), dimension(:), allocatable **vacmod::avv**
- real(rprec), dimension(:), allocatable **vacmod::snr**
- real(rprec), dimension(:), allocatable **vacmod::snv**
- real(rprec), dimension(:), allocatable **vacmod::snz**
- real(rprec), dimension(:), allocatable **vacmod::drv**
- real(rprec), dimension(:), allocatable **vacmod::guu_b**
- real(rprec), dimension(:), allocatable **vacmod::guv_b**
- real(rprec), dimension(:), allocatable **vacmod::gvv_b**
- real(rprec), dimension(:), allocatable **vacmod::rzb2**
- real(rprec), dimension(:), allocatable **vacmod::rcosuv**
- real(rprec), dimension(:), allocatable **vacmod::rsinuv**
- real(rprec), dimension(:), allocatable **vacmod::raxis_nestor**
- real(rprec), dimension(:), allocatable **vacmod::zaxis_nestor**
- real(rprec), dimension(:), allocatable **vacmod::bsubu**
- real(rprec), dimension(:), allocatable **vacmod::bsubv**
- real(rprec), dimension(:), allocatable **vacmod::potu**
- real(rprec), dimension(:), allocatable **vacmod::potv**
- real(rprec), dimension(:), allocatable **vacmod::amatrix**
- real(rprec), dimension(:), allocatable **vacmod::ruu**

- real(rprec), dimension(:), allocatable **vacmod::ruv**
- real(rprec), dimension(:), allocatable **vacmod::rvv**
- real(rprec), dimension(:), allocatable **vacmod::zuu**
- real(rprec), dimension(:), allocatable **vacmod::zuv**
- real(rprec), dimension(:), allocatable **vacmod::zvv**
- real(rprec), dimension(:), allocatable **vacmod::brad**
- real(rprec), dimension(:), allocatable **vacmod::bphi**
- real(rprec), dimension(:), allocatable **vacmod::bz**
- real(rprec), dimension(:,), allocatable **vacmod::xpts**
- real(rprec), dimension(:), allocatable **vacmod::grpmn**
- real(rprec), dimension(:), allocatable **vacmod::grpmn_m_map_wrt**
- real(rprec), dimension(:), allocatable **vacmod::grpmn_n_map_wrt**
- real(rprec), dimension(:), allocatable **vacmod::gstore**
- real(rprec), dimension(:,), allocatable **vacmod::green**
- real(rprec), dimension(:,), allocatable **vacmod::greenp**
- real(rprec), dimension(:), allocatable **vacmod::r0p**
- real(rprec), dimension(:), allocatable **vacmod::r1p**
- real(rprec), dimension(:), allocatable **vacmod::r0m**
- real(rprec), dimension(:), allocatable **vacmod::r1m**
- real(rprec), dimension(:), allocatable **vacmod::sqrtc**
- real(rprec), dimension(:), allocatable **vacmod::sqrta**
- real(rprec), dimension(:), allocatable **vacmod::t1p2**
- real(rprec), dimension(:), allocatable **vacmod::t1p1**
- real(rprec), dimension(:), allocatable **vacmod::t1p**
- real(rprec), dimension(:), allocatable **vacmod::t1m2**
- real(rprec), dimension(:), allocatable **vacmod::t1m1**
- real(rprec), dimension(:), allocatable **vacmod::t1m**
- real(rprec), dimension(:), allocatable **vacmod::adp**
- real(rprec), dimension(:), allocatable **vacmod::adm**
- real(rprec), dimension(:), allocatable **vacmod::cma**
- real(rprec), dimension(:), allocatable **vacmod::ra1p**
- real(rprec), dimension(:), allocatable **vacmod::ra1m**
- real(rprec), dimension(:), allocatable **vacmod::slm**
- real(rprec), dimension(:), allocatable **vacmod::slp**
- real(rprec), dimension(:), allocatable **vacmod::tlpm**
- real(rprec), dimension(:), allocatable **vacmod::slpm**
- real(rprec), dimension(:), allocatable **vacmod::delt1u**
- real(rprec), dimension(:), allocatable **vacmod::azp1u**
- real(rprec), dimension(:), allocatable **vacmod::azm1u**
- real(rprec), dimension(:), allocatable **vacmod::cma11u**
- real(rprec), dimension(:), allocatable **vacmod::sqad1u**
- real(rprec), dimension(:), allocatable **vacmod::sqad2u**
- real(rprec), dimension(:,), allocatable **vacmod::all_t1p**
- real(rprec), dimension(:,), allocatable **vacmod::all_t1m**
- real(rprec), dimension(:,), allocatable **vacmod::all_slp**
- real(rprec), dimension(:,), allocatable **vacmod::all_slm**
- real(rprec), dimension(:), allocatable **vacmod::gsave**
- real(rprec), dimension(:), allocatable **vacmod::ga1**
- real(rprec), dimension(:), allocatable **vacmod::ga2**
- real(rprec), dimension(:), allocatable **vacmod::dsave**
- real(rprec), dimension(:,,:), allocatable **vacmod::g1**
- real(rprec), dimension(:,,:), allocatable **vacmod::g2**
- real(rprec), dimension(:,,:), allocatable **vacmod::bcos**
- real(rprec), dimension(:,,:), allocatable **vacmod::bsin**
- real(rprec), dimension(:,,:), allocatable **vacmod::source**
- real(rprec), dimension(:,,:), allocatable **vacmod::actemp**
- real(rprec), dimension(:,,:), allocatable **vacmod::astemp**

7.66 src/NESTOR/data/vacmod0.f90 File Reference

Functions/Subroutines

- subroutine **vacmod0::set_nestor_sizes** (nfp, ntor, mpol, nzeta, ntheta, lasym)

Variables

- integer **vacmod0::mf**
- integer **vacmod0::nf**
- integer **vacmod0::nu**
- integer **vacmod0::nv**
- integer **vacmod0::mf1**
- integer **vacmod0::nf1**
- integer **vacmod0::mnpd**
- integer **vacmod0::mnpd2**
- integer **vacmod0::nuv**
- integer **vacmod0::nu2**
- integer **vacmod0::nu3**
- integer **vacmod0::nuv2**
- integer **vacmod0::nfper**
- integer **vacmod0::nvper**
- integer **vacmod0::nuv_tan**
- integer **vacmod0::nvp**
- integer **vacmod0::ndim**

7.67 src/NESTOR/fouri.f90 File Reference

Functions/Subroutines

- subroutine **fouri** (grpmn, gsource, amatrix, amatsq, bvec, wint, lasym)

7.67.1 Function/Subroutine Documentation

7.67.1.1 **fouri()**

```
subroutine fouri (
    real(rprec), dimension(mnpd,nv,nu3,ndim), intent(in) grpmn,
    real(rprec), dimension(nuv), intent(in) gsource,
    real(rprec), dimension(mnpd,mnpd,ndim**2), intent(out) amatrix,
    real(rprec), dimension(mnpd2,mnpd2), intent(out) amatsq,
    real(rprec), dimension(0:mf,-nf:nf,ndim), intent(inout) bvec,
    real(rprec), dimension(nuv2), intent(in) wint,
    logical, intent(in) lasym )
```

interior (int_ext=-1), exterior (int_ext=+1) neumann problem

Definition at line 2 of file fouri.f90.

7.68 src/NESTOR/fourp.f90 File Reference

Functions/Subroutines

- subroutine **fourp** (grpmn, grp)

7.69 src/NESTOR/greenf.f90 File Reference

Functions/Subroutines

- subroutine **greenf** (delgr, delgrp, ip)

7.70 src/NESTOR/nestor_main.f90 File Reference

Main program of stand-alone version of NESTOR.

Functions/Subroutines

- program **nestor**
Main program of stand-alone version of NESTOR.

7.70.1 Detailed Description

Main program of stand-alone version of NESTOR.

7.71 src/NESTOR/precalf.f90 File Reference

Functions/Subroutines

- subroutine **precalf**

7.72 src/NESTOR/scalpot.f90 File Reference

Functions/Subroutines

- subroutine **scalpot** (bvec, amatrix, wint, ivacskip, lasym, m_map, n_map)

7.73 src/NESTOR/surface.f90 File Reference

Functions/Subroutines

- subroutine **surface** (rc, rs, zs, zc, xm, xn, mnmax, lasym, signs)

7.74 src/NESTOR/vacuum.f90 File Reference

Functions/Subroutines

- subroutine **vacuum** (rmnc, rmns, zmns, zmnc, xm, xn, plascur, rbtor, wint, ivac_skip, ivac, mnmax, ier_flag, lasym, signsg, raxis, zaxis)

7.75 src/open_output_files.f90 File Reference

Open output files.

Functions/Subroutines

- subroutine [open_output_files](#) (extension, lfirst)
Open output files.

7.75.1 Detailed Description

Open output files.

7.75.2 Function/Subroutine Documentation

7.75.2.1 open_output_files()

```
subroutine open_output_files (
    character(len=*) extension,
    logical lfirst )
```

Open output files.

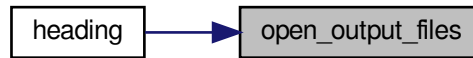
Parameters

<i>extension</i>	input file "extension": part after 'input.'.
<i>lfirst</i>	flag to indicate if this is the first call to this routine or not

Definition at line 8 of file open_output_files.f90.

Referenced by heading().

Here is the caller graph for this function:



7.76 src/parse_extension.f File Reference

Parse the first command-line argument into a filename.

Functions/Subroutines

- subroutine [parse_extension](#) (file_to_parse, file_or_extension, Inc)
Parse the first command-line argument into a filename.

7.76.1 Detailed Description

Parse the first command-line argument into a filename.

7.76.2 Function/Subroutine Documentation

7.76.2.1 parse_extension()

```

subroutine parse_extension (
    character(len=*) , intent(inout) file_to_parse,
    character(len=*) , intent(in)   file_or_extension,
    logical, intent(out) Inc )
  
```

Parse the first command-line argument into a filename.

Parameters

<i>file_to_parse</i>	actual filename to read the input for VMEC from
<i>file_or_extension</i>	first command-line parameter given to VMEC
<i>Inc</i>	flag to indicate that a netCDF file is given

Definition at line 9 of file parse_extension.f.

7.77 src/precondn.f90 File Reference

Compute preconditioning matrix elements for R , Z force.

Functions/Subroutines

- subroutine [precondn](#) (*lu1*, *bsq*, *gsqrt*, *r12*, *xs*, *xu12*, *xue*, *xuo*, *xodd*, *axm*, *axd*, *bxm*, *bx**d*, *cx*, *eqfactor*, *trigmult*)
Compute preconditioning matrix elements for R , Z force.

7.77.1 Detailed Description

Compute preconditioning matrix elements for R , Z force.

7.77.2 Function/Subroutine Documentation

7.77.2.1 [precondn\(\)](#)

```
subroutine precondn (
    real(rprec), dimension(nrzt), intent(in) lu1,
    real(rprec), dimension(nrzt), intent(in) bsq,
    real(rprec), dimension(nrzt), intent(in) gsqrt,
    real(rprec), dimension(nrzt), intent(in) r12,
    real(rprec), dimension(nrzt), intent(in) xs,
    real(rprec), dimension(nrzt), intent(in) xu12,
    real(rprec), dimension(nrzt), intent(in) xue,
    real(rprec), dimension(nrzt), intent(in) xuo,
    real(rprec), dimension(nrzt), intent(in) xodd,
    real(rprec), dimension(ns+1,2), intent(out) axm,
    real(rprec), dimension(ns+1,2), intent(out) axd,
    real(rprec), dimension(ns+1,2), intent(out) bxm,
    real(rprec), dimension(ns+1,2), intent(out) bxd,
    real(rprec), dimension(ns+1), intent(out) cx,
    real(rprec), dimension(ns), intent(out) eqfactor,
    real(rprec), dimension(nznt), intent(in) trigmult )
```

Compute preconditioning matrix elements for R , Z force.

Parameters

<i>lu1</i>	
<i>bsq</i>	
<i>gsqrt</i>	
<i>r12</i>	
<i>xs</i>	
<i>xu12</i>	
<i>xue</i>	
<i>xuo</i>	
<i>xodd</i>	

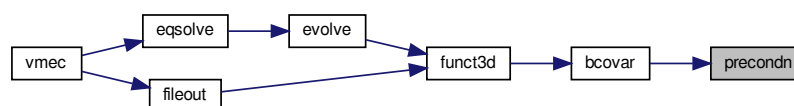
Parameters

<i>axm</i>	
<i>axd</i>	
<i>bxm</i>	
<i>bxm</i>	
<i>cx</i>	
<i>eqfactor</i>	
<i>trigmult</i>	

Definition at line 22 of file precondition.f90.

Referenced by bcovar().

Here is the caller graph for this function:



7.78 src/printout.f90 File Reference

Print iteration progress to screen and `threed1` output file.

Functions/Subroutines

- subroutine `printout` (*i0*, *delt0*, *w0*)
Print iteration progress to screen and threed1 output file.

7.78.1 Detailed Description

Print iteration progress to screen and `threed1` output file.

7.78.2 Function/Subroutine Documentation

7.78.2.1 printout()

```

subroutine printout (
    integer i0,
    real(rprec) delt0,
    real(rprec) w0 )

```

Print iteration progress to screen and `threed1` output file.

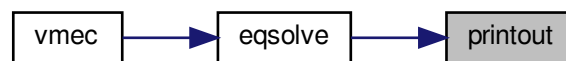
Parameters

<i>i0</i>	current iteration number
<i>delt0</i>	current time step
<i>w0</i>	current MHD energy

Definition at line 9 of file printout.f90.

Referenced by eqsolve().

Here is the caller graph for this function:



7.79 src/profil1d.f90 File Reference

Compute phip and iota profiles on full grid.

Functions/Subroutines

- subroutine [profil1d](#) (xc, xcdot, lreset)
Compute phip and iota profiles on full grid.

7.79.1 Detailed Description

Compute phip and iota profiles on full grid.

7.79.2 Function/Subroutine Documentation

7.79.2.1 profil1d()

```

subroutine profil1d (
    real(rprec), dimension(neqs), intent(out) xc,
    real(rprec), dimension(neqs), intent(out) xcdot,
    logical, intent(in) lreset )

```

Compute phip and iota profiles on full grid.

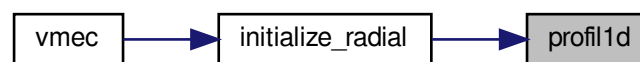
Parameters

<i>xc</i>	state vector of VMEC, i.e., all Fourier coefficients of R , Z and λ
<i>xcdot</i>	velocity vector in Fourier space
<i>lreset</i>	xc will be zeroes if this is true

Definition at line 9 of file profil1d.f90.

Referenced by `initialize_radial()`.

Here is the caller graph for this function:



7.80 src/profil3d.f90 File Reference

Functions/Subroutines

- subroutine **profil3d** (rmn, zmn, lreset)

7.81 src/profile_functions.f File Reference

Functions/Subroutines

- real(rprec) function **pcurr** (xx)
- real(rprec) function **piota** (x)
- real(rprec) function **pmass** (xx)

7.82 src/read_indata.f90 File Reference

Functions/Subroutines

- subroutine **read_indata** (in_file, iunit, ier_flag)

7.83 src/read_wout_mod.f File Reference

Data Types

- interface [read_wout_mod::read_wout_file](#)

Functions/Subroutines

- subroutine **read_wout_mod::readw_and_open** (file_or_extension, ierr, iopen)
- subroutine **read_wout_mod::compute_currents** (ierror)
- subroutine **read_wout_mod::read_wout_deallocate**
- subroutine **read_wout_mod::tosuvspace** (s_in, u_in, v_in, gsqr, bsupu, bsupv, jsupu, jsupv, lam)
- subroutine **read_wout_mod::loadrzi**

Variables

- character(len= *), parameter **read_wout_mod::vn_version** = 'version_'
- character(len= *), parameter **read_wout_mod::vn_extension** = 'input_extension'
- character(len= *), parameter **read_wout_mod::vn_mgrid** = 'mgrid_file'
- character(len= *), parameter **read_wout_mod::vn_magen** = 'wb'
- character(len= *), parameter **read_wout_mod::vn_therm** = 'wp'
- character(len= *), parameter **read_wout_mod::vn_gam** = 'gamma'
- character(len= *), parameter **read_wout_mod::vn_maxr** = 'rmax_surf'
- character(len= *), parameter **read_wout_mod::vn_minr** = 'rmin_surf'
- character(len= *), parameter **read_wout_mod::vn_maxz** = 'zmax_surf'
- character(len= *), parameter **read_wout_mod::vn_fp** = 'nfp'
- character(len= *), parameter **read_wout_mod::vn_radnod** = 'ns'
- character(len= *), parameter **read_wout_mod::vn_polmod** = 'mpol'
- character(len= *), parameter **read_wout_mod::vn_tormod** = 'ntor'
- character(len= *), parameter **read_wout_mod::vn_maxmod** = 'mnmax'
- character(len= *), parameter **read_wout_mod::vn_maxit** = 'niter'
- character(len= *), parameter **read_wout_mod::vn_actit** = 'itfsq'
- character(len= *), parameter **read_wout_mod::vn_asym** = 'lasym'
- character(len= *), parameter **read_wout_mod::vn_free** = 'lfreeb'
- character(len= *), parameter **read_wout_mod::vn_error** = 'ier_flag'
- character(len= *), parameter **read_wout_mod::vn_aspect** = 'aspect'
- character(len= *), parameter **read_wout_mod::vn_maxmod_nyq** = 'mnmax_nyq'
- character(len= *), parameter **read_wout_mod::vn_beta** = 'betatotal'
- character(len= *), parameter **read_wout_mod::vn_pbeta** = 'betapol'
- character(len= *), parameter **read_wout_mod::vn_tbeta** = 'betator'
- character(len= *), parameter **read_wout_mod::vn_abeta** = 'betaxis'
- character(len= *), parameter **read_wout_mod::vn_b0** = 'b0'
- character(len= *), parameter **read_wout_mod::vn_rbt0** = 'rbtor0'
- character(len= *), parameter **read_wout_mod::vn_rbt1** = 'rbtor'
- character(len= *), parameter **read_wout_mod::vn_sgs** = 'signs'
- character(len= *), parameter **read_wout_mod::vn_lar** = 'lonLarmor'
- character(len= *), parameter **read_wout_mod::vn_modb** = 'volavgB'
- character(len= *), parameter **read_wout_mod::vn_ctor** = 'ctor'
- character(len= *), parameter **read_wout_mod::vn_amin** = 'Aminor_p'
- character(len= *), parameter **read_wout_mod::vn_rmaj** = 'Rmajor_p'
- character(len= *), parameter **read_wout_mod::vn_vol** = 'volume_p'
- character(len= *), parameter **read_wout_mod::vn_am** = 'am'
- character(len= *), parameter **read_wout_mod::vn_ai** = 'ai'
- character(len= *), parameter **read_wout_mod::vn_ac** = 'ac'
- character(len= *), parameter **read_wout_mod::vn_ah** = 'hot particle fraction'
- character(len= *), parameter **read_wout_mod::vn_atuname** = 'T-perp/T-par'
- character(len= *), parameter **read_wout_mod::vn_pmass_type** = 'pmass_type'
- character(len= *), parameter **read_wout_mod::vn_piota_type** = 'piota_type'
- character(len= *), parameter **read_wout_mod::vn_pcurr_type** = 'pcurr_type'
- character(len= *), parameter **read_wout_mod::vn_am_aux_s** = 'am_aux_s'

- character(len= *), parameter **read_wout_mod::vn_am_aux_f** = 'am_aux_f'
- character(len= *), parameter **read_wout_mod::vn_ai_aux_s** = 'ai_aux_s'
- character(len= *), parameter **read_wout_mod::vn_ai_aux_f** = 'ai_aux_f'
- character(len= *), parameter **read_wout_mod::vn_ac_aux_s** = 'ac_aux_s'
- character(len= *), parameter **read_wout_mod::vn_ac_aux_f** = 'ac_aux_f'
- character(len= *), parameter **read_wout_mod::vn_mse** = 'imse'
- character(len= *), parameter **read_wout_mod::vn_thom** = 'itse'
- character(len= *), parameter **read_wout_mod::vn_pmod** = 'xm'
- character(len= *), parameter **read_wout_mod::vn_tmod** = 'xn'
- character(len= *), parameter **read_wout_mod::vn_pmod_nyq** = 'xm_nyq'
- character(len= *), parameter **read_wout_mod::vn_tmod_nyq** = 'xn_nyq'
- character(len= *), parameter **read_wout_mod::vn_racc** = 'raxis_cc'
- character(len= *), parameter **read_wout_mod::vn_zacs** = 'zaxis_cs'
- character(len= *), parameter **read_wout_mod::vn_racs** = 'raxis_cs'
- character(len= *), parameter **read_wout_mod::vn_zacc** = 'zaxis_cc'
- character(len= *), parameter **read_wout_mod::vn_iotaf** = 'iotaf'
- character(len= *), parameter **read_wout_mod::vn_qfact** = 'q-factor'
- character(len= *), parameter **read_wout_mod::vn_chi** = 'chi'
- character(len= *), parameter **read_wout_mod::vn_chipf** = 'chipf'
- character(len= *), parameter **read_wout_mod::vn_presf** = 'presf'
- character(len= *), parameter **read_wout_mod::vn_phi** = 'phi'
- character(len= *), parameter **read_wout_mod::vn_phipf** = 'phipf'
- character(len= *), parameter **read_wout_mod::vn_jcuru** = 'jcuru'
- character(len= *), parameter **read_wout_mod::vn_jcurv** = 'jcurv'
- character(len= *), parameter **read_wout_mod::vn_iotah** = 'iotas'
- character(len= *), parameter **read_wout_mod::vn_mass** = 'mass'
- character(len= *), parameter **read_wout_mod::vn_presh** = 'pres'
- character(len= *), parameter **read_wout_mod::vn_betah** = 'beta_vol'
- character(len= *), parameter **read_wout_mod::vn_buco** = 'buco'
- character(len= *), parameter **read_wout_mod::vn_bvco** = 'bvco'
- character(len= *), parameter **read_wout_mod::vn_vp** = 'vp'
- character(len= *), parameter **read_wout_mod::vn_specw** = 'specw'
- character(len= *), parameter **read_wout_mod::vn_phip** = 'phips'
- character(len= *), parameter **read_wout_mod::vn_jdotb** = 'jdotb'
- character(len= *), parameter **read_wout_mod::vn_overr** = 'over_r'
- character(len= *), parameter **read_wout_mod::vn_bgrv** = 'bdotgradv'
- character(len= *), parameter **read_wout_mod::vn_merc** = 'DMerc'
- character(len= *), parameter **read_wout_mod::vn_mshear** = 'DShear'
- character(len= *), parameter **read_wout_mod::vn_mwell** = 'DWell'
- character(len= *), parameter **read_wout_mod::vn_mcurr** = 'DCurr'
- character(len= *), parameter **read_wout_mod::vn_mgeo** = 'DGeod'
- character(len= *), parameter **read_wout_mod::vn_equif** = 'equip'
- character(len= *), parameter **read_wout_mod::vn_fsq** = 'fsqt'
- character(len= *), parameter **read_wout_mod::vn_wdot** = 'wdot'
- character(len= *), parameter **read_wout_mod::vn_ftolv** = 'ftolv'
- character(len= *), parameter **read_wout_mod::vn_fsqli** = 'fsqli'
- character(len= *), parameter **read_wout_mod::vn_fsqr** = 'fsqr'
- character(len= *), parameter **read_wout_mod::vn_fsqz** = 'fsqz'
- character(len= *), parameter **read_wout_mod::vn_extcur** = 'extcur'
- character(len= *), parameter **read_wout_mod::vn_curlab** = 'curlabel'
- character(len= *), parameter **read_wout_mod::vn_rmnc** = 'rmnc'
- character(len= *), parameter **read_wout_mod::vn_zmns** = 'zmns'
- character(len= *), parameter **read_wout_mod::vn_lmns** = 'lmns'
- character(len= *), parameter **read_wout_mod::vn_gmnc** = 'gmnc'
- character(len= *), parameter **read_wout_mod::vn_bmnc** = 'bmnc'

- character(len= *), parameter **read_wout_mod::vn_bsubumnc** = 'bsubumnc'
- character(len= *), parameter **read_wout_mod::vn_bsubvmnc** = 'bsubvmnc'
- character(len= *), parameter **read_wout_mod::vn_bsubsmns** = 'bsubsmns'
- character(len= *), parameter **read_wout_mod::vn_bsupumnc** = 'bsupumnc'
- character(len= *), parameter **read_wout_mod::vn_bsupvmnc** = 'bsupvmnc'
- character(len= *), parameter **read_wout_mod::vn_rmns** = 'rmns'
- character(len= *), parameter **read_wout_mod::vn_zmnc** = 'zmnc'
- character(len= *), parameter **read_wout_mod::vn_lmnc** = 'lmnc'
- character(len= *), parameter **read_wout_mod::vn_gmns** = 'gmns'
- character(len= *), parameter **read_wout_mod::vn_bmns** = 'bmns'
- character(len= *), parameter **read_wout_mod::vn_bsubumns** = 'bsubumns'
- character(len= *), parameter **read_wout_mod::vn_bsubvmns** = 'bsubvmns'
- character(len= *), parameter **read_wout_mod::vn_bsubsmnc** = 'bsubsmnc'
- character(len= *), parameter **read_wout_mod::vn_bsupumns** = 'bsupumns'
- character(len= *), parameter **read_wout_mod::vn_bsupvmns** = 'bsupvmns'
- character(len= *), parameter **read_wout_mod::vn_bsubumnc_sur** = 'bsubumnc_sur'
- character(len= *), parameter **read_wout_mod::vn_bsubvmnc_sur** = 'bsubvmnc_sur'
- character(len= *), parameter **read_wout_mod::vn_bsupumnc_sur** = 'bsupumnc_sur'
- character(len= *), parameter **read_wout_mod::vn_bsupvmnc_sur** = 'bsupvmnc_sur'
- character(len= *), parameter **read_wout_mod::vn_bsubumns_sur** = 'bsubumns_sur'
- character(len= *), parameter **read_wout_mod::vn_bsubvmns_sur** = 'bsubvmns_sur'
- character(len= *), parameter **read_wout_mod::vn_bsupumns_sur** = 'bsupumns_sur'
- character(len= *), parameter **read_wout_mod::vn_bsupvmns_sur** = 'bsupvmns_sur'
- character(len= *), parameter **read_wout_mod::vn_rbc** = 'rbc'
- character(len= *), parameter **read_wout_mod::vn_zbs** = 'zbs'
- character(len= *), parameter **read_wout_mod::vn_rbs** = 'rbs'
- character(len= *), parameter **read_wout_mod::vn_zbc** = 'zbc'
- character(len= *), parameter **read_wout_mod::vn_potvac** = 'potvac'
- character(len= *), parameter **read_wout_mod::ln_version** = 'VMEC Version'
- character(len= *), parameter **read_wout_mod::ln_extension** = 'Input file extension'
- character(len= *), parameter **read_wout_mod::ln_mgrid** = 'MGRID file'
- character(len= *), parameter **read_wout_mod::ln_magen** = 'Magnetic Energy'
- character(len= *), parameter **read_wout_mod::ln_therm** = 'Thermal Energy'
- character(len= *), parameter **read_wout_mod::ln_gam** = 'Gamma'
- character(len= *), parameter **read_wout_mod::ln_maxr** = 'Maximum R'
- character(len= *), parameter **read_wout_mod::ln_minr** = 'Minimum R'
- character(len= *), parameter **read_wout_mod::ln_maxz** = 'Maximum Z'
- character(len= *), parameter **read_wout_mod::ln_fp** = 'Field Periods'
- character(len= *), parameter **read_wout_mod::ln_radnod** = 'Radial nodes'
- character(len= *), parameter **read_wout_mod::ln_polmod** = 'Poloidal modes'
- character(len= *), parameter **read_wout_mod::ln_tormod** = 'Toroidal modes'
- character(len= *), parameter **read_wout_mod::ln_maxmod** = 'Fourier modes'
- character(len= *), parameter **read_wout_mod::ln_maxmod_nyq** = 'Fourier modes (Nyquist)'
- character(len= *), parameter **read_wout_mod::ln_maxit** = 'Max iterations'
- character(len= *), parameter **read_wout_mod::ln_actit** = 'Actual iterations'
- character(len= *), parameter **read_wout_mod::ln_asym** = 'Asymmetry'
- character(len= *), parameter **read_wout_mod::ln_recon** = 'Reconstruction'
- character(len= *), parameter **read_wout_mod::ln_free** = 'Free boundary'
- character(len= *), parameter **read_wout_mod::ln_error** = 'Error flag'
- character(len= *), parameter **read_wout_mod::ln_aspect** = 'Aspect ratio'
- character(len= *), parameter **read_wout_mod::ln_beta** = 'Total beta'
- character(len= *), parameter **read_wout_mod::ln_pbeta** = 'Poloidal beta'
- character(len= *), parameter **read_wout_mod::ln_tbeta** = 'Toroidal beta'
- character(len= *), parameter **read_wout_mod::ln_abeta** = 'Beta axis'
- character(len= *), parameter **read_wout_mod::ln_b0** = 'RB-t over R axis'

- character(len= *), parameter **read_wout_mod::ln_rbt0** = 'RB-t axis'
- character(len= *), parameter **read_wout_mod::ln_rbt1** = 'RB-t edge'
- character(len= *), parameter **read_wout_mod::ln_sgs** = 'Sign [jacobian](#)'
- character(len= *), parameter **read_wout_mod::ln_lar** = 'Ion Larmor radius'
- character(len= *), parameter **read_wout_mod::ln_modb** = 'avg mod B'
- character(len= *), parameter **read_wout_mod::ln_ctor** = 'Toroidal current'
- character(len= *), parameter **read_wout_mod::ln_amin** = 'minor radius'
- character(len= *), parameter **read_wout_mod::ln_rmaj** = 'major radius'
- character(len= *), parameter **read_wout_mod::ln_vol** = 'Plasma volume'
- character(len= *), parameter **read_wout_mod::ln_mse** = 'Number of MSE points'
- character(len= *), parameter **read_wout_mod::ln_thom** = 'Number of Thompson scattering points'
- character(len= *), parameter **read_wout_mod::ln_am** = 'Specification parameters for mass(s)'
- character(len= *), parameter **read_wout_mod::ln_ac** = 'Specification parameters for <J>(s)'
- character(len= *), parameter **read_wout_mod::ln_ai** = 'Specification parameters for iota(s)'
- character(len= *), parameter **read_wout_mod::ln_pmass_type** = 'Profile type specifier for mass(s)'
- character(len= *), parameter **read_wout_mod::ln_pcurr_type** = 'Profile type specifier for <J>(s)'
- character(len= *), parameter **read_wout_mod::ln_piota_type** = 'Profile type specifier for iota(s)'
- character(len= *), parameter **read_wout_mod::ln_am_aux_s** = 'Auxiliary-s parameters for mass(s)'
- character(len= *), parameter **read_wout_mod::ln_am_aux_f** = 'Auxiliary-f parameters for mass(s)'
- character(len= *), parameter **read_wout_mod::ln_ac_aux_s** = 'Auxiliary-s parameters for <J>(s)'
- character(len= *), parameter **read_wout_mod::ln_ac_aux_f** = 'Auxiliary-f parameters for <J>(s)'
- character(len= *), parameter **read_wout_mod::ln_ai_aux_s** = 'Auxiliary-s parameters for iota(s)'
- character(len= *), parameter **read_wout_mod::ln_ai_aux_f** = 'Auxiliary-f parameters for iota(s)'
- character(len= *), parameter **read_wout_mod::ln_pmod** = 'Poloidal mode numbers'
- character(len= *), parameter **read_wout_mod::ln_tmod** = 'Toroidal mode numbers'
- character(len= *), parameter **read_wout_mod::ln_pmod_nyq** = 'Poloidal mode numbers (Nyquist)'
- character(len= *), parameter **read_wout_mod::ln_tmod_nyq** = 'Toroidal mode numbers (Nyquist)'
- character(len= *), parameter **read_wout_mod::ln_racc** = 'raxis (cosnv)'
- character(len= *), parameter **read_wout_mod::ln_racs** = 'raxis (sinnv)'
- character(len= *), parameter **read_wout_mod::ln_zacs** = 'zaxis (sinnv)'
- character(len= *), parameter **read_wout_mod::ln_zacc** = 'zaxis (cosnv)'
- character(len= *), parameter **read_wout_mod::ln_iotaf** = 'iota on full mesh'
- character(len= *), parameter **read_wout_mod::ln_qfact** = 'q-factor on full mesh'
- character(len= *), parameter **read_wout_mod::ln_presf** = 'pressure on full mesh'
- character(len= *), parameter **read_wout_mod::ln_phi** = 'Toroidal flux on full mesh'
- character(len= *), parameter **read_wout_mod::ln_phipf** = 'd(phi)/ds: Toroidal flux deriv on full mesh'
- character(len= *), parameter **read_wout_mod::ln_chi** = 'Poloidal flux on full mesh'
- character(len= *), parameter **read_wout_mod::ln_chipf** = 'd(chi)/ds: Poroidal flux deriv on full mesh'
- character(len= *), parameter **read_wout_mod::ln_jcuru** = 'j dot gradu full'
- character(len= *), parameter **read_wout_mod::ln_jcurv** = 'j dot gradv full'
- character(len= *), parameter **read_wout_mod::ln_iotah** = 'iota half'
- character(len= *), parameter **read_wout_mod::ln_mass** = 'mass half'
- character(len= *), parameter **read_wout_mod::ln_presh** = 'pressure half'
- character(len= *), parameter **read_wout_mod::ln_betah** = 'beta half'
- character(len= *), parameter **read_wout_mod::ln_buco** = 'bsubu half'
- character(len= *), parameter **read_wout_mod::ln_bvco** = 'bsubv half'
- character(len= *), parameter **read_wout_mod::ln_vp** = 'volume deriv half'
- character(len= *), parameter **read_wout_mod::ln_specw** = 'Spectral width half'
- character(len= *), parameter **read_wout_mod::ln_phip** = 'tor flux deriv over 2pi half'
- character(len= *), parameter **read_wout_mod::ln_jdotb** = 'J dot B'
- character(len= *), parameter **read_wout_mod::ln_bgrv** = 'B dot grad v'
- character(len= *), parameter **read_wout_mod::ln_merc** = 'Mercier criterion'
- character(len= *), parameter **read_wout_mod::ln_mshear** = 'Shear Mercier'
- character(len= *), parameter **read_wout_mod::ln_mwell** = 'Well Mercier'
- character(len= *), parameter **read_wout_mod::ln_mcurr** = 'Current Mercier'

- character(len= *), parameter **read_wout_mod::ln_mgeo** = 'Geodesic Mercier'
- character(len= *), parameter **read_wout_mod::ln_equip** = 'Average force balance'
- character(len= *), parameter **read_wout_mod::ln_fsq** = 'Residual decay'
- character(len= *), parameter **read_wout_mod::ln_wdot** = 'Wdot decay'
- character(len= *), parameter **read_wout_mod::ln_extcur** = 'External coil currents'
- character(len= *), parameter **read_wout_mod::ln_fsqr** = 'Residual decay - radial'
- character(len= *), parameter **read_wout_mod::ln_fsqz** = 'Residual decay - vertical'
- character(len= *), parameter **read_wout_mod::ln_fsqli** = 'Residual decay - hoop'
- character(len= *), parameter **read_wout_mod::ln_ftolv** = 'Residual decay - requested'
- character(len= *), parameter **read_wout_mod::ln_curlab** = 'External current names'
- character(len= *), parameter **read_wout_mod::ln_rmnc** = 'cosmn component of cylindrical R, full mesh'
- character(len= *), parameter **read_wout_mod::ln_zmns** = 'sinmn component of cylindrical Z, full mesh'
- character(len= *), parameter **read_wout_mod::ln_lmns** = 'sinmn component of lambda, half mesh'
- character(len= *), parameter **read_wout_mod::ln_gmnc** = 'cosmn component of [jacobian](#), half mesh'
- character(len= *), parameter **read_wout_mod::ln_bmnc** = 'cosmn component of mod-B, half mesh'
- character(len= *), parameter **read_wout_mod::ln_bsubumnc** = 'cosmn covariant u-component of B, half mesh'
- character(len= *), parameter **read_wout_mod::ln_bsubvmnc** = 'cosmn covariant v-component of B, half mesh'
- character(len= *), parameter **read_wout_mod::ln_bsubsmns** = 'sinmn covariant s-component of B, full mesh'
- character(len= *), parameter **read_wout_mod::ln_bsubumnc_sur** = 'cosmn bsubu of B, surface'
- character(len= *), parameter **read_wout_mod::ln_bsubvmnc_sur** = 'cosmn bsubv of B, surface'
- character(len= *), parameter **read_wout_mod::ln_bsupumnc_sur** = 'cosmn bsupu of B, surface'
- character(len= *), parameter **read_wout_mod::ln_bsupvmnc_sur** = 'cosmn bsupv of B, surface'
- character(len= *), parameter **read_wout_mod::ln_bsupumnc** = 'BSUPUmnc half'
- character(len= *), parameter **read_wout_mod::ln_bsupvmnc** = 'BSUPVmnc half'
- character(len= *), parameter **read_wout_mod::ln_rmns** = 'sinmn component of cylindrical R, full mesh'
- character(len= *), parameter **read_wout_mod::ln_zmnc** = 'cosmn component of cylindrical Z, full mesh'
- character(len= *), parameter **read_wout_mod::ln_lmnc** = 'cosmn component of lambda, half mesh'
- character(len= *), parameter **read_wout_mod::ln_gmns** = 'sinmn component of [jacobian](#), half mesh'
- character(len= *), parameter **read_wout_mod::ln_bmns** = 'sinmn component of mod-B, half mesh'
- character(len= *), parameter **read_wout_mod::ln_bsubumns** = 'sinmn covariant u-component of B, half mesh'
- character(len= *), parameter **read_wout_mod::ln_bsubvmns** = 'sinmn covariant v-component of B, half mesh'
- character(len= *), parameter **read_wout_mod::ln_bsubsmnc** = 'cosmn covariant s-component of B, full mesh'
- character(len= *), parameter **read_wout_mod::ln_bsubumns_sur** = 'sinmn bsubu of B, surface'
- character(len= *), parameter **read_wout_mod::ln_bsubvmns_sur** = 'sinmn bsubv of B, surface'
- character(len= *), parameter **read_wout_mod::ln_bsupumns_sur** = 'sinmn bsupu of B, surface'
- character(len= *), parameter **read_wout_mod::ln_bsupvmns_sur** = 'sinmn bsupv of B, surface'
- character(len= *), parameter **read_wout_mod::ln_bsupumns** = 'BSUPUmns half'
- character(len= *), parameter **read_wout_mod::ln_bsupvmns** = 'BSUPVmns half'
- character(len= *), parameter **read_wout_mod::ln_rbc** = 'Initial boundary R cos(mu-nv) coefficients'
- character(len= *), parameter **read_wout_mod::ln_zbs** = 'Initial boundary Z sin(mu-nv) coefficients'
- character(len= *), parameter **read_wout_mod::ln_rbs** = 'Initial boundary R sin(mu-nv) coefficients'
- character(len= *), parameter **read_wout_mod::ln_zbc** = 'Initial boundary Z cos(mu-nv) coefficients'
- character(len= *), parameter **read_wout_mod::ln_potvac** = 'Vacuum Potential on Boundary'
- integer **read_wout_mod::nfp**
- integer **read_wout_mod::ns**
- integer **read_wout_mod::mpol**
- integer **read_wout_mod::ntor**
- integer **read_wout_mod::mnmax**
- integer **read_wout_mod::mnmax_nyq**

- integer `read_wout_mod::itfsq`
- integer `read_wout_mod::niter`
- integer `read_wout_mod::iasym`
- integer `read_wout_mod::ierr_vmec`
- integer `read_wout_mod::imse`
- integer `read_wout_mod::itse`
- integer `read_wout_mod::nstore_seq`
- integer `read_wout_mod::isnodes`
- integer `read_wout_mod::ipnodes`
- integer `read_wout_mod::imatch_phiedge`
- integer `read_wout_mod::isigng`
- integer `read_wout_mod::mnyq`
- integer `read_wout_mod::nnyq`
- integer `read_wout_mod::ntmax`
- real(rprec) `read_wout_mod::wb`
- real(rprec) `read_wout_mod::wp`
- real(rprec) `read_wout_mod::gamma`
- real(rprec) `read_wout_mod::pfac`
- real(rprec) `read_wout_mod::rmax_surf`
- real(rprec) `read_wout_mod::rmin_surf`
- real(rprec) `read_wout_mod::zmax_surf`
- real(rprec) `read_wout_mod::aspect`
- real(rprec) `read_wout_mod::betatot`
- real(rprec) `read_wout_mod::betapol`
- real(rprec) `read_wout_mod::betator`
- real(rprec) `read_wout_mod::betaxis`
- real(rprec) `read_wout_mod::b0`
- real(rprec) `read_wout_mod::tswgt`
- real(rprec) `read_wout_mod::msewgt`
- real(rprec) `read_wout_mod::flmwgt`
- real(rprec) `read_wout_mod::bcwgt`
- real(rprec) `read_wout_mod::phidiam`
- real(rprec) `read_wout_mod::version_`
- real(rprec) `read_wout_mod::delphid`
- real(rprec) `read_wout_mod::ionlarmor`
- real(rprec) `read_wout_mod::volavgb`
- real(rprec) `read_wout_mod::fsql`
- real(rprec) `read_wout_mod::fsqr`
- real(rprec) `read_wout_mod::fsqz`
- real(rprec) `read_wout_mod::ftolv`
- real(rprec) `read_wout_mod::aminor`
- real(rprec) `read_wout_mod::rmajor`
- real(rprec) `read_wout_mod::volume`
- real(rprec) `read_wout_mod::rbtor`
- real(rprec) `read_wout_mod::rbtor0`
- real(rprec) `read_wout_mod::itor`
- real(rprec) `read_wout_mod::machsq`
- real(rprec), dimension(:,:), allocatable `read_wout_mod::rzi_local`
- real(rprec), dimension(:,:), allocatable `read_wout_mod::rmnc`
- real(rprec), dimension(:,:), allocatable `read_wout_mod::zmns`
- real(rprec), dimension(:,:), allocatable `read_wout_mod::lmns`
- real(rprec), dimension(:,:), allocatable `read_wout_mod::rmns`
- real(rprec), dimension(:,:), allocatable `read_wout_mod::zmnc`
- real(rprec), dimension(:,:), allocatable `read_wout_mod::lmnc`
- real(rprec), dimension(:,:), allocatable `read_wout_mod::bmnc`

- `real(rprec), dimension(:, :), allocatable read_wout_mod::gmnc`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bsubumnc`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bsubvmnc`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bsubsmns`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bsupumnc`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bsupvmnc`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::curvmnc`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::currumnc`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bbc`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::raxis`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::zaxis`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bmns`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::gmns`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bsubumns`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bsubvmns`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bsubsmnc`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bsupumns`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::bsupvmns`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::currumns`
- `real(rprec), dimension(:, :), allocatable read_wout_mod::curvmns`
- `real(rprec), dimension(:), allocatable read_wout_mod::iotas`
- `real(rprec), dimension(:), allocatable read_wout_mod::iotaf`
- `real(rprec), dimension(:), allocatable read_wout_mod::presf`
- `real(rprec), dimension(:), allocatable read_wout_mod::phipf`
- `real(rprec), dimension(:), allocatable read_wout_mod::mass`
- `real(rprec), dimension(:), allocatable read_wout_mod::pres`
- `real(rprec), dimension(:), allocatable read_wout_mod::beta_vol`
- `real(rprec), dimension(:), allocatable read_wout_mod::xm`
- `real(rprec), dimension(:), allocatable read_wout_mod::xn`
- `real(rprec), dimension(:), allocatable read_wout_mod::qfact`
- `real(rprec), dimension(:), allocatable read_wout_mod::chipf`
- `real(rprec), dimension(:), allocatable read_wout_mod::phi`
- `real(rprec), dimension(:), allocatable read_wout_mod::chi`
- `real(rprec), dimension(:), allocatable read_wout_mod::xm_nyq`
- `real(rprec), dimension(:), allocatable read_wout_mod::xn_nyq`
- `real(rprec), dimension(:), allocatable read_wout_mod::phip`
- `real(rprec), dimension(:), allocatable read_wout_mod::buco`
- `real(rprec), dimension(:), allocatable read_wout_mod::bvco`
- `real(rprec), dimension(:), allocatable read_wout_mod::vp`
- `real(rprec), dimension(:), allocatable read_wout_mod::overr`
- `real(rprec), dimension(:), allocatable read_wout_mod::jcuru`
- `real(rprec), dimension(:), allocatable read_wout_mod::jcurv`
- `real(rprec), dimension(:), allocatable read_wout_mod::specw`
- `real(rprec), dimension(:), allocatable read_wout_mod::jdotb`
- `real(rprec), dimension(:), allocatable read_wout_mod::bdotgradv`
- `real(rprec), dimension(:), allocatable read_wout_mod::fsqt`
- `real(rprec), dimension(:), allocatable read_wout_mod::wdot`
- `real(rprec), dimension(:), allocatable read_wout_mod::am`
- `real(rprec), dimension(:), allocatable read_wout_mod::ac`
- `real(rprec), dimension(:), allocatable read_wout_mod::ai`
- `real(rprec), dimension(:), allocatable read_wout_mod::am_aux_s`
- `real(rprec), dimension(:), allocatable read_wout_mod::am_aux_f`
- `real(rprec), dimension(:), allocatable read_wout_mod::ac_aux_s`
- `real(rprec), dimension(:), allocatable read_wout_mod::ac_aux_f`
- `real(rprec), dimension(:), allocatable read_wout_mod::ai_aux_s`

- real(rprec), dimension(:), allocatable **read_wout_mod::ai_aux_f**
- real(rprec), dimension(:), allocatable **read_wout_mod::dmerc**
- real(rprec), dimension(:), allocatable **read_wout_mod::dshear**
- real(rprec), dimension(:), allocatable **read_wout_mod::dwell**
- real(rprec), dimension(:), allocatable **read_wout_mod::dcurr**
- real(rprec), dimension(:), allocatable **read_wout_mod::dgeod**
- real(rprec), dimension(:), allocatable **read_wout_mod::equif**
- real(rprec), dimension(:), allocatable **read_wout_mod::extcur**
- real(rprec), dimension(:), allocatable **read_wout_mod::sknots**
- real(rprec), dimension(:), allocatable **read_wout_mod::ystark**
- real(rprec), dimension(:), allocatable **read_wout_mod::y2stark**
- real(rprec), dimension(:), allocatable **read_wout_mod::pknots**
- real(rprec), dimension(:), allocatable **read_wout_mod::ythom**
- real(rprec), dimension(:), allocatable **read_wout_mod::y2thom**
- real(rprec), dimension(:), allocatable **read_wout_mod::anglemse**
- real(rprec), dimension(:), allocatable **read_wout_mod::rmid**
- real(rprec), dimension(:), allocatable **read_wout_mod::qmid**
- real(rprec), dimension(:), allocatable **read_wout_mod::shear**
- real(rprec), dimension(:), allocatable **read_wout_mod::presmid**
- real(rprec), dimension(:), allocatable **read_wout_mod::alfa**
- real(rprec), dimension(:), allocatable **read_wout_mod::curmid**
- real(rprec), dimension(:), allocatable **read_wout_mod::rstark**
- real(rprec), dimension(:), allocatable **read_wout_mod::qmeas**
- real(rprec), dimension(:), allocatable **read_wout_mod::datastark**
- real(rprec), dimension(:), allocatable **read_wout_mod::rthom**
- real(rprec), dimension(:), allocatable **read_wout_mod::datathom**
- real(rprec), dimension(:), allocatable **read_wout_mod::dsiobt**
- real(rprec), dimension(:), allocatable **read_wout_mod::potvac**
- logical **read_wout_mod::lasym**
- logical **read_wout_mod::lthreed**
- logical **read_wout_mod::lwout_opened** = .false.
- character **read_wout_mod::mgrid_file**
- character **read_wout_mod::input_extension**
- character **read_wout_mod::pmass_type**
- character **read_wout_mod::pcurr_type**
- character **read_wout_mod::piota_type**

7.84 src/readin.f90 File Reference

Functions/Subroutines

- subroutine **readin** (input_file, ier_flag)

7.85 src/reset_params.f90 File Reference

Functions/Subroutines

- subroutine [reset_params](#)

7.85.1 Function/Subroutine Documentation

7.85.1.1 `reset_params()`

```
subroutine reset_params
```

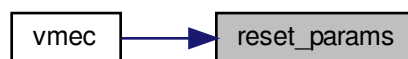
m=1 constraint (=t: apply correct, polar constraint; =f, apply approx. constraint)

Assume scaled mode; read in from mgrid in free-bdy mode

Definition at line 2 of file `reset_params.f90`.

Referenced by `vmec()`.

Here is the caller graph for this function:



7.86 `src/residue.f90` File Reference

Functions/Subroutines

- subroutine **residue** (gcr, gcx, gcl)
- subroutine **constrain_m1** (gcr, gcx)
- subroutine **scale_m1** (gcr, gcx)

7.87 `src/restart_iter.f90` File Reference

Functions/Subroutines

- subroutine **restart_iter** (time_step)

7.88 `src/safe_open_mod.f` File Reference

Functions/Subroutines

- subroutine **safe_open_mod::safe_open** (iunit, istat, filename, filestat, fileform, record_in, access_in, delim_in)

7.89 src/scalfor.f90 File Reference

Functions/Subroutines

- subroutine **scalfor** (gcx, axm, bxm, axd, bxd, cx, iflag)

7.90 src/solver.f90 File Reference

Functions/Subroutines

- subroutine **solver** (amat, b, m, nrhs, info)

7.91 src/spectrum.f90 File Reference

Functions/Subroutines

- subroutine **spectrum** (rmn, zmn)

7.92 src/spline_akima.f File Reference

Functions/Subroutines

- subroutine **spline_akima** (x, y, xx, yy, npts, iflag)

7.93 src/spline_akima_int.f File Reference

Functions/Subroutines

- subroutine **spline_akima_int** (x, y, xx, yy, npts, iflag)

7.94 src/spline_cubic.f File Reference

Functions/Subroutines

- subroutine **spline_cubic** (x, y, xx, yy, n, iflag)
- subroutine **spline_nr** (x, y, n, yp1, ypn, y2)
- subroutine **splint_nr** (xa, ya, y2a, n, x, y)

7.95 src/spline_cubic_int.f File Reference

Functions/Subroutines

- subroutine **spline_cubic_int** (x, y, xx, yy, n, iflag)
- subroutine **spline_int** (x, y, n, yp1, ypn, y2)
- subroutine **splint_int** (xa, ya, y2a, n, x, y)

7.96 src/symforce.f90 File Reference

Functions/Subroutines

- subroutine **symforce** (ars, brs, crs, azs, bzs, czs, bls, cls, rcs, zcs, ara, bra, cra, aza, bza, cza, bla, cla, rca, zca)
- subroutine **symoutput** (bsq, gsqr, bsubu, bsubv, bsupu, bsupv, bsubs, bsqa, gsqrta, bsubua, bsubva, bsupua, bsupva, bsubsa)

7.97 src/symrzi.f90 File Reference

Functions/Subroutines

- subroutine **symrzi** (r1s, rus, rvs, z1s, zus, zvs, lus, lvs, rcons, zcons, r1a, rua, rva, z1a, zua, zva, lua, lva, rcona, zcona)

7.98 src/tolower.f90 File Reference

Functions/Subroutines

- subroutine **tolower** (string)

7.99 src/tomnsp.f90 File Reference

Functions/Subroutines

- subroutine **tomnsp** (frzl_array, armn, brmn, crmn, azmn, bzm, czmn, blmn, clmn, arcn, azcn)
- subroutine **tomnsa** (frzl_array, armn, brmn, crmn, azmn, bzm, czmn, blmn, clmn, arcn, azcn)

7.100 src/totzsp.f90 File Reference

Functions/Subroutines

- subroutine **totzsps** (rzi_array, r11, ru1, rv1, z11, zu1, zv1, lu1, lv1, rcn1, zcn1)
- subroutine **convert_sym** (rmnss, zmncs)
- subroutine **totzspa** (rzi_array, r11, ru1, rv1, z11, zu1, zv1, lu1, lv1, rcn1, zcn1)
- subroutine **convert_asym** (rmnsc, zmnc)

7.100.1 Function/Subroutine Documentation

7.100.1.1 totzsps()

```

subroutine totzsps (
    real(rprec), dimension(ns,0:ntor,0:mpoll,3*ntmax), intent(inout), target rzl_←
array,
    real(rprec), dimension(ns*nzeta*ntheta3,0:1), intent(out) r11,
    real(rprec), dimension(ns*nzeta*ntheta3,0:1), intent(out) ru1,
    real(rprec), dimension(ns*nzeta*ntheta3,0:1), intent(out) rv1,
    real(rprec), dimension(ns*nzeta*ntheta3,0:1), intent(out) z11,
    real(rprec), dimension(ns*nzeta*ntheta3,0:1), intent(out) zu1,
    real(rprec), dimension(ns*nzeta*ntheta3,0:1), intent(out) zv1,
    real(rprec), dimension(ns*nzeta*ntheta3,0:1), intent(out) lu1,
    real(rprec), dimension(ns*nzeta*ntheta3,0:1), intent(out) lv1,
    real(rprec), dimension(ns*nzeta*ntheta3,0:1), intent(out) rcn1,
    real(rprec), dimension(ns*nzeta*ntheta3,0:1), intent(out) zcn1 )

```

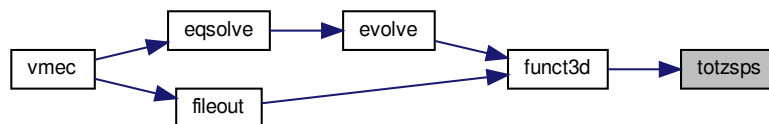
Parameters

out	<i>r11</i>	R
out	<i>ru1</i>	dR/dTheta
out	<i>rv1</i>	dR/dZeta
out	<i>z11</i>	Z
out	<i>zu1</i>	dZ/dTheta
out	<i>zv1</i>	dZ/dZeta
out	<i>lu1</i>	dLambda/dTheta
out	<i>lv1</i>	-dLambda/dZeta
out	<i>rcn1</i>	TODO: what is this?
out	<i>zcn1</i>	TODO: what is this?

Definition at line 2 of file totzsp.f90.

Referenced by funct3d().

Here is the caller graph for this function:



7.101 src/tridslv.f90 File Reference

Functions/Subroutines

- subroutine **tridslv** (a, d, b, c, jmin, jmax, mnd1, ns, nrhs)

7.102 src/vmec.f90 File Reference

Main program of VMEC.

Functions/Subroutines

- program `vmec`
Main program of VMEC.

7.102.1 Detailed Description

Main program of VMEC.

7.103 src/wrout.f90 File Reference

Functions/Subroutines

- subroutine `wrout` (bsq, gsqrt, bsubu, bsubv, bsubs, bsupv, bsupu, rzl_array, gc_array, ier_flag)

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