eqtools Documentation

Release 1.1

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November 10, 2016

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Homepage: https://github.com/PSFCPlasmaTools/eqtools

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CHAPTER

ONE

OVERVIEW

eqtools is a Python package for working with magnetic equilibrium reconstructions from magnetic plasma confinement devices. At present, interfaces exist for data from the Alcator C-Mod and NSTX MDSplus trees as well as eqdsk a- and g-files. eqtools is designed to be flexible and extensible such that it can become a uniform interface to perform mapping operations and accessing equilibrium data for any magnetic confinement device, regardless of how the data are accessed.

The main class of eqtools is the Equilibrium, which contains all of the coordinate mapping functions as well as templates for methods to fetch data (primarily dictated to the quantities computed by EFIT). Subclasses such as EFITTree, CModEFITTree, NSTXEFITTree and EqdskReader implement specific methods to access the data and convert it to the form needed for the routines in Equilibrium. These classes are smart about caching intermediate results, so you will get a performance boost by using the same instance throughout your analysis of a given shot.

CHAPTER

TWO

INSTALLATION

The easiest way to install the latest release version is with *pip*:

pip install eqtools

To install from source, uncompress the source files and, from the directory containing *setup.py*, run the following command:

python setup.py install

Or, to build in place, run:

python setup.py build_ext --inplace

TUTORIAL: PERFORMING COORDINATE TRANSFORMS ON ALCATOR C-MOD DATA

The basic class for manipulating EFIT results stored in the Alcator C-Mod MDSplus tree is CModEFITTree. To load the data from a specific shot, simply create the CModEFITTree object with the shot number as the argument:

```
e = eqtools.CModEFITTree(1140729030)
```

The default EFIT to use is "ANALYSIS." If you want to use a different tree, such as "EFIT20," then you simply set this with the *tree* keyword:

```
e = eqtools.CModEFITTree(1140729030, tree='EFIT20')
```

eqtools understands units. The default is to convert all lengths to meters (whereas quantities in the tree are inconsistent – some are meters, some centimeters). If you want to specify a different default unit, use the *length_unit* keyword:

```
e = eqtools.CModEFITTree(1140729030, length_unit='cm')
```

Once this is loaded, you can access the data you would normally have to pull from specific nodes in the tree using convenient getter methods. For instance, to get the elongation as a function of time, you can run:

```
kappa = e.getElongation()
```

The timebase used for quantities like this is accessed with:

```
t = e.getTimeBase()
```

For length/area/volume quantities, eqtools understands units. The default is to return in whatever units you specified when creating the CModEFITTree, but you can override this with the *length_unit* keyword. For instance, to get the vertical position of the magnetic axis in mm, you can run:

```
Z_mag = e.getMagZ(length_unit='mm')
```

eqtools can map from almost any coordinate to any common flux surface label. For instance, say you want to know what the square root of normalized toroidal flux corresponding to a normalized flux surface volume of 0.5 is at t=1.0s. You can simply call:

```
rho = e.volnorm2phinorm(0.5, 1.0, sqrt=True)
```

If a list of times is provided, the default behavior is to evaluate all of the points to be converted at each of the times. So, to follow the mapping of normalized poloidal flux values [0.1, 0.5, 1.0] to outboard midplane major radius at time points [1.0, 1.25, 1.5, 1.75], you could call:

```
psinorm = e.psinorm2rmid([0.1, 0.5, 1.0], [1.0, 1.25, 1.5, 1.75])
```

This will return a 4-by-3 array: one row for each time, one column for each location. If you want to override this behavior and instead consider a sequence of (psi, t) points, set the *each_t* keyword to False:

```
psinorm = e.psinorm2rmid([0.3, 0.35], [1.0, 1.1], each_t=False)
```

This will return a two-element array with the Rmid values for (psinorm=0.3, t=1.0) and (psinorm=0.35, t=1.1).

For programmatically mapping between coordinates, the rho2rho() method is quite useful. To map from outboard midplane major radius to normalized flux surface volume, you can simply call:

```
e.rho2rho('Rmid', 'volnorm', 0.75, 1.0)
```

Finally, to get a look at the flux surfaces, simply run:

```
e.plotFlux()
```

CHAPTER

FOUR

PACKAGE REFERENCE

eqtools package

Submodules

eqtools.CModEFIT module

This module provides classes for working with C-Mod EFIT data.

```
 \textbf{class} \texttt{ eqtools.CModEFIT.CModEFITTree} (\textit{shot}, \textit{tree='ANALYSIS'}, \textit{length\_unit='m'}, \textit{gfile='g\_eqdsk'}, \\ \textit{afile='a\_eqdsk'}, \textit{tspline=False}, \textit{monotonic=False}) \\ \textbf{Bases: eqtools.EFIT.EFITTree}
```

Inherits EFITTree class. Machine-specific data handling class for Alcator C-Mod. Pulls EFIT data from selected MDS tree and shot, stores as object attributes. Each EFIT variable or set of variables is recovered with a corresponding getter method. Essential data for EFIT mapping are pulled on initialization (e.g. psirz grid). Additional data are pulled at the first request and stored for subsequent usage.

Intializes C-Mod version of EFITTree object. Pulls data from MDS tree for storage in instance attributes. Core attributes are populated from the MDS tree on initialization. Additional attributes are initialized as None, filled on the first request to the object.

- **shot** (long) int C-Mod shot index (long)
- **tree** str Optional input for EFIT tree, defaults to 'ANALYSIS' (i.e., EFIT data are under analysis::top.efit.results). For any string TREE (such as 'EFIT20') other than 'ANALYSIS', data are taken from TREE::top.results.
- **length_unit** str Sets the base unit used for any quantity whose dimensions are length to any power. Valid options are:

62	F 24
'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'de-	whatever the default in the tree is (no conversion is performed, units may
fault'	be inconsistent)

Default is 'm' (all units taken and returned in meters).

- **gfile** str Optional input for EFIT geqdsk location name, defaults to 'g_eqdsk' (i.e., EFIT data are under tree::top.results.G_EQDSK)
- afile str Optional input for EFIT aeqdsk location name, defaults to 'a_eqdsk' (i.e., EFIT data are under tree::top.results.A_EQDSK)
- **tspline** Boolean Sets whether or not interpolation in time is performed using a tricubic spline or nearest-neighbor interpolation. Tricubic spline interpolation requires at least four complete equilibria at different times. It is also assumed that they are functionally correlated, and that parameters do not vary out of their boundaries (derivative = 0 boundary condition). Default is False (use nearest neighbor interpolation).
- monotonic Boolean Sets whether or not the "monotonic" form of time window finding is used. If True, the timebase must be monotonically increasing. Default is False (use slower, safer method).

```
getFluxVol (length_unit=3)
          returns volume within flux surface [t,psi]
     getRmidPsi(length_unit=1)
          returns maximum major radius of each flux surface [t,psi]
     getF()
          returns F=RB_{Phi}(Psi), often calculated for grad-shafranov solutions [t,psi]
     getFluxPres()
          returns pressure at flux surface [t,psi]
     getFFPrime()
          returns FF' function used for grad-shafranov solutions [t,psi]
     getPPrime()
          returns plasma pressure gradient as a function of psi [t,psi]
     getQProfile()
          returns safety factor q [t,psi]
     getRLCFS (length_unit=1)
          returns R-values of LCFS position [t,n]
     getZLCFS (length unit=1)
          returns Z-values of LCFS position [t,n]
     getMachineCrossSectionFull()
          Pulls C-Mod cross-section data from tree, converts to plottable vector format for use in other plotting
          routines
               Parameters shot – (long) int C-Mod shot index (used for tree access)
               Returns The coordinates of the machine cross-section.
               Return type (x, y)
class eqtools.CModEFIT.CModEFITTreeProp (shot,
                                                               tree='ANALYSIS',
                                                                                      length unit='m'.
                                                   gfile='g_eqdsk',
                                                                     afile='a_eqdsk',
                                                                                        tspline=False,
                                                   monotonic=False)
     Bases: eqtools.CModEFIT.CModEFITTree, eqtools.core.PropertyAccessMixin
```

CModEFITTree with the PropertyAccessMixin added to enable property-style access. This is good for interactive use, but may drag the performance down.

eqtools.D3DEFIT module

eqtools.EFIT module

Bases: eqtools.core.Equilibrium

Inherits Equilibrium class. EFIT-specific data handling class for machines using standard EFIT tag names/tree structure with MDSplus. Constructor and/or data loading may need overriding in a machine-specific implementation. Pulls EFIT data from selected MDS tree and shot, stores as object attributes. Each EFIT variable or set of variables is recovered with a corresponding getter method. Essential data for EFIT mapping are pulled on initialization (e.g. psirz grid). Additional data are pulled at the first request and stored for subsequent usage.

Intializes EFITTree object. Pulls data from MDS tree for storage in instance attributes. Core attributes are populated from the MDS tree on initialization. Additional attributes are initialized as None, filled on the first request to the object.

Parameters

- **shot** int shot number
- tree tree MDSplus tree to open to fetch EFIT data.
- root str Root path for EFIT data in MDSplus tree.
- **length_unit** String. Sets the base unit used for any quantity whose dimensions are length to any power. Valid options are:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'de-	whatever the default in the tree is (no conversion is performed, units may
fault'	be inconsistent)

Default is 'm' (all units taken and returned in meters).

- **tspline** Boolean. Sets whether or not interpolation in time is performed using a tricubic spline or nearest-neighbor interpolation. Tricubic spline interpolation requires at least four complete equilibria at different times. It is also assumed that they are functionally correlated, and that parameters do not vary out of their boundaries (derivative = 0 boundary condition). Default is False (use nearest neighbor interpolation).
- monotonic Boolean. Sets whether or not the "monotonic" form of time window finding is used. If True, the timebase must be monotonically increasing. Default is False (use slower, safer method).

getInfo()

returns namedtuple of shot information

Returns

namedtuple containing

	shot	C-Mod shot index (long) EFIT tree (string)
	nr	size of R-axis for spatial grid
	nz nt	size of Z-axis for spatial grid size of timebase for flux grid
getTimeBase returns EFI		
getFluxGrid returns EFI		grid, [t,z,r]
getRGrid (len returns EFI	_	
getZGrid (len returns EFI	-	
getFluxAxis returns psi		netic axis [t]
getFluxLCFS returns psi		atrix [t]
getFluxVol (returns volu	-	unit=3) hin flux surface [t,psi]
getVolLCFS (returns volu	_	unit=3) hin LCFS [t]
getRmidPsi (returns max	-	unit=1) major radius of each flux surface [t,psi]
getRLCFS (len returns R-v	-	t=1) ELCFS position [t,n]
getZLCFS (len returns Z-v	-	t=1) FLCFS position [t,n]
surface. Th	RLCFS is is the	lse) S, ZLCFS values pulled from EFIT with explicitly-calculated contour of psinorm=1 on masked down by the limiter array using core.inPolygon, restricting the contour to the ace and the divertor legs.
		mask – Boolean. Default False. Set True to mask LCFS path to limiter outline Polygon). Set False to draw full contour of psi = psiLCFS.
getF() returns F=F	RB_{Ph	i}(Psi), often calculated for grad-shafranov solutions [t,psi]
getFluxPres returns pres		flux surface [t,psi]
<pre>getFFPrime() returns FF' function used for grad-shafranov solutions [t,psi]</pre>		
getPPrime() returns plas	sma pres	ssure gradient as a function of psi [t,psi]
getElongati returns LC		gation [t]

```
getUpperTriangularity()
     returns LCFS upper triangularity [t]
getLowerTriangularity()
     returns LCFS lower triangularity [t]
getSlhaping()
     pulls LCFS elongation and upper/lower triangularity
         Returns namedtuple containing {kappa, delta_u, delta_l}
getMagR (length_unit=1)
     returns magnetic-axis major radius [t]
getMagZ (length_unit=1)
     returns magnetic-axis Z [t]
getAreaLCFS (length_unit=2)
     returns LCFS cross-sectional area [t]
getAOut (length_unit=1)
     returns outboard-midplane minor radius at LCFS [t]
getRmidOut (length_unit=1)
     returns outboard-midplane major radius [t]
getGeometry (length_unit=None)
     pulls dimensional geometry parameters
         Returns namedtuple containing {magnetic-axis R,Z, LCFS area, outboard-midplane LCFS a,R}
getQProfile()
     returns safety factor q [t,psi]
getQ0()
     returns q on magnetic axis [t]
getQ95()
     returns q at 95% flux surface [t]
getQLCFS()
     returns q on LCFS [t]
getQ1Surf (length_unit=1)
     returns outboard-midplane minor radius of q=1 surface [t]
getQ2Surf (length_unit=1)
     returns outboard-midplane minor radius of q=2 surface [t]
getQ3Surf (length_unit=1)
     returns outboard-midplane minor radius of q=3 surface [t]
getQs (length_unit=1)
     pulls q values
         Returns namedtuple containing {q0,q95,qLCFS,rq1,rq2,rq3}
getBtVac()
     returns on-axis vacuum toroidal field [t]
getBtPla()
     returns on-axis plasma toroidal field [t]
getBpAvg()
     returns average poloidal field [t]
```

```
getFields()
     pulls vacuum and plasma toroidal field, avg poloidal field
         Returns namedtuple containing {btaxv,btaxp,bpolav}
getIpCalc()
     returns EFIT-calculated plasma current [t]
getIpMeas()
     returns magnetics-measured plasma current [t]
getJp()
     returns EFIT-calculated plasma current density Jp on flux grid [t,r,z]
getBetaT()
     returns EFIT-calculated toroidal beta [t]
getBetaP()
     returns EFIT-calculated poloidal beta [t]
getLi()
     returns EFIT-calculated internal inductance [t]
getBetas()
     pulls calculated betap, betat, internal inductance
         Returns namedtuple containing {betat,betap,Li}
getDiamagFlux()
     returns measured diamagnetic-loop flux [t]
getDiamagBetaT()
     returns diamagnetic-loop toroidal beta [t]
getDiamagBetaP()
     returns diamagnetic-loop avg poloidal beta [t]
getDiamagTauE()
     returns diamagnetic-loop energy confinement time [t]
getDiamagWp()
     returns diamagnetic-loop plasma stored energy [t]
getDiamag()
     pulls diamagnetic flux measurements, toroidal and poloidal beta, energy confinement time and stored
         Returns namedtuple containing {diamag. flux, betatd, betapd, tauDiamag, WDiamag}
getWMHD()
     returns EFIT-calculated MHD stored energy [t]
getTauMHD()
     returns EFIT-calculated MHD energy confinement time [t]
getPinj()
     returns EFIT-calculated injected power [t]
getWbdot()
     returns EFIT-calculated d/dt of magnetic stored energy [t]
getWpdot()
     returns EFIT-calculated d/dt of plasma stored energy [t]
```

getEnergy()

pulls EFIT-calculated energy parameters - stored energy, tau_E, injected power, d/dt of magnetic and plasma stored energy

Returns namedtuple containing {WMHD,tauMHD,Pinj,Wbdot,Wpdot}

getMachineCrossSection()

Returns R,Z coordinates of vacuum-vessel wall for masking, plotting routines.

Returns The requested data.

getMachineCrossSectionFull()

Returns R,Z coordinates of vacuum-vessel wall for plotting routines.

Absent additional vector-graphic data on machine cross-section, returns self.getMachineCrossSection().

Returns The requested data.

getCurrentSign()

Returns the sign of the current, based on the check in Steve Wolfe's IDL implementation efit_rz2psi.pro.

getParam(path)

backup function - path to parameter as input, returns desired variable acts as wrapper for MDS call

Parameters path – str The path to the MDSplus node you wish to pull in.

Returns The requested data.

eqtools.FromArrays module

Bases: eqtools.core.Equilibrium

Class to represent an equilibrium specified as arrays of data.

Create ArrayEquilibrium instance from arrays of data.

Has very little checking on the shape/type of the arrays at this point.

- psiRZ Array-like, (M, N, P). Flux values at M times, N Z locations and P R locations.
- rGrid Array-like, (P,). R coordinates that psiRZ is given at.
- **zGrid** Array-like, (N₁). Z coordinates that psiRZ is given at.
- **time** Array-like, (M,). Times that psiRZ is given at.
- q Array-like, (S, M). q profile evaluated at S values of psinorm from 0 to 1, given at M times.
- **fluxVol** Array-like, (S, M). Flux surface volumes evaluated at S values of psinorm from 0 to 1, given at M times.
- psiLCFS Array-like, (M,). Flux at the last closed flux surface, given at M times.
- **psiAxis** Array-like, (M,). Flux at the magnetic axis, given at M times.
- **rmag** Array-like, (M,). Radial coordinate of the magnetic axis, given at M times.
- zmag Array-like, (M,). Vertical coordinate of the magnetic axis, given at M times.
- Rout Outboard midplane radius of the last closed flux surface.

• **length_unit** – String. Sets the base unit used for any quantity whose dimensions are length to any power. Valid options are:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'de-	whatever the default in the tree is (no conversion is performed, units may
fault'	be inconsistent)

Default is 'm' (all units taken and returned in meters).

- **tspline** Boolean. Sets whether or not interpolation in time is performed using a tricubic spline or nearest-neighbor interpolation. Tricubic spline interpolation requires at least four complete equilibria at different times. It is also assumed that they are functionally correlated, and that parameters do not vary out of their boundaries (derivative = 0 boundary condition). Default is False (use nearest neighbor interpolation).
- **monotonic** Boolean. Sets whether or not the "monotonic" form of time window finding is used. If True, the timebase must be monotonically increasing. Default is False (use slower, safer method).
- verbose Boolean. Allows or blocks console readout during operation. Defaults to True, displaying useful information for the user. Set to False for quiet usage or to avoid console clutter for multiple instances.

```
getTimeBase()
     Returns a copy of the time base vector, array dimensions are (M<sub>1</sub>).
getFluxGrid()
     Returns a copy of the flux array, dimensions are (M, N, P), corresponding to (time, Z, R).
getRGrid (length_unit=1)
     Returns a copy of the radial grid, dimensions are (P,).
getZGrid(length unit=1)
     Returns a copy of the vertical grid, dimensions are (N<sub>1</sub>).
getQProfile()
     Returns safety factor q profile (over Q values of psinorm from 0 to 1), dimensions are (Q, M)
getFluxVol (length_unit=3)
     returns volume within flux surface [psi,t]
getFluxLCFS()
     returns psi at separatrix [t]
getFluxAxis()
     returns psi on magnetic axis [t]
getMagR (length_unit=1)
     returns magnetic-axis major radius [t]
getMagZ (length_unit=1)
```

returns magnetic-axis Z [t]

```
getRmidOut (length_unit=1)
    returns outboard-midplane major radius [t]
getRLCFS (length_unit=1)
getZLCFS (length_unit=1)
getCurrentSign ()
```

eqtools.NSTXEFIT module

This module provides classes for working with NSTX EFIT data.

Inherits EFITTree class. Machine-specific data handling class for the National Spherical Torus Experiment (NSTX). Pulls EFIT data from selected MDS tree and shot, stores as object attributes. Each EFIT variable or set of variables is recovered with a corresponding getter method. Essential data for EFIT mapping are pulled on initialization (e.g. psirz grid). Additional data are pulled at the first request and stored for subsequent usage.

Intializes NSTX version of EFITTree object. Pulls data from MDS tree for storage in instance attributes. Core attributes are populated from the MDS tree on initialization. Additional attributes are initialized as None, filled on the first request to the object.

Parameters

- shot (long) int NSTX shot index (long)
- **tree** str Optional input for EFIT tree, defaults to 'EFIT01' (i.e., EFIT data are under EFIT01::top.results).
- **length_unit** str Sets the base unit used for any quantity whose dimensions are length to any power. Valid options are:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'de-	whatever the default in the tree is (no conversion is performed, units may
fault'	be inconsistent)

Default is 'm' (all units taken and returned in meters).

- gfile str Optional input for EFIT geqdsk location name, defaults to 'g_eqdsk' (i.e., EFIT data are under tree::top.results.G_EQDSK)
- afile str Optional input for EFIT aeqdsk location name, defaults to 'a_eqdsk' (i.e., EFIT data are under tree::top.results.A_EQDSK)
- **tspline** Boolean Sets whether or not interpolation in time is performed using a tricubic spline or nearest-neighbor interpolation. Tricubic spline interpolation requires at least four complete equilibria at different times. It is also assumed that they are functionally correlated, and that parameters do not vary out of their boundaries (derivative = 0 boundary condition). Default is False (use nearest neighbor interpolation).

• monotonic – Boolean Sets whether or not the "monotonic" form of time window finding is used. If True, the timebase must be monotonically increasing. Default is False (use slower, safer method).

```
getFluxGrid()
     returns EFIT flux grid, [t,z,r]
getMachineCrossSection()
     Returns R,Z coordinates of vacuum-vessel wall for masking, plotting routines.
         Returns The requested data.
getFluxVol()
     Not implemented in NSTXEFIT tree.
     Returns volume within flux surface [psi,t]
getRmidPsi(length_unit=1)
     returns maximum major radius of each flux surface [t,psi]
getIpCalc()
     returns EFIT-calculated plasma current [t]
getVolLCFS (length unit=3)
     returns volume within LCFS [t]
getJp()
     Not implemented in NSTXEFIT tree.
     returns EFIT-calculated plasma current density Jp on flux grid [t,r,z]
rz2volnorm(*args, **kwargs)
     Calculated normalized volume of flux surfaces not stored in NSTX EFIT. All maping with Volnorm not
     implemented
```

psinorm2volnorm (*args, **kwargs)

Calculated normalized volume of flux surfaces not stored in NSTX EFIT. All maping with Volnorm not

```
class eqtools.NSTXEFIT.NSTXEFITTreeProp (shot, tree='EFIT01', length_unit='m', gfile='geqdsk', afile='aeqdsk', tspline=False, monotonic=False)
```

Bases: eqtools.NSTXEFIT.NSTXEFITTree, eqtools.core.PropertyAccessMixin

NSTXEFITTree with the PropertyAccessMixin added to enable property-style access. This is good for interactive use, but may drag the performance down.

eqtools.TCVLIUQE module

implemented

eqtools.AUGDATA module

eqtools.afilereader module

This module contains the AFileReader class, a lightweight data handler for a-file (time-history) datasets.

Classes:

AFileReader: Data-storage class for a-file data. Reads data from ASCII a-file, storing as copy-safe object attributes.

Class to read ASCII a-file (time-history data storage) into lightweight, user-friendly data structure.

A-files store data blocks of scalar time-history data for EFIT plasma equilibrium. Each parameter is read into a pseudo-private object attribute (marked by a leading underscore), followed by the standard EFIT variable names.

initialize object, reading from file.

Parameters afile – str path to a-file

eqtools.core module

This module provides the core classes for eqtools, including the base Equilibrium class.

```
exception eqtools.core.ModuleWarning
```

Bases: exceptions.Warning

Warning class to notify the user of unavailable modules.

x.__init__(...) initializes x; see help(type(x)) for signature

$class \verb| eqtools.core.PropertyAccessMixin|$

Bases: object

Mixin to implement access of getter methods through a property-type interface without the need to apply a decorator to every property.

For any getter *obj.getSomething()*, the call *obj.Something* will do the same thing.

This is accomplished by overriding __getattribute__() such that if an attribute *ATTR* does not exist it then attempts to call *self.getATTR()*. If *self.getATTR()* does not exist, an AttributeError will be raised as usual.

Also overrides __setattr__() such that it will raise an AttributeError when attempting to write an attribute *ATTR* for which there is already a method *getATTR*.

x.__init__(...) initializes x; see help(type(x)) for signature

eqtools.core.inPolygon(polyx, polyy, pointx, pointy)

Function calculating whether a given point is within a 2D polygon.

Given an array of X,Y coordinates describing a 2D polygon, checks whether a point given by x,y coordinates lies within the polygon. Operates via a ray-casting approach - the function projects a semi-infinite ray parallel to the positive horizontal axis, and counts how many edges of the polygon this ray intersects. For a simply-connected polygon, this determines whether the point is inside (even number of crossings) or outside (odd number of crossings) the polygon, by the Jordan Curve Theorem.

Parameters

- polyx (Array-like) Array of x-coordinates of the vertices of the polygon.
- polyy (Array-like) Array of y-coordinates of the vertices of the polygon.
- **pointx** (*Int or float*) x-coordinate of test point.
- **pointy** (*Int or float*) y-coordinate of test point.

Returns True/False result for whether the point is contained within the polygon.

Return type result (Boolean)

```
class eqtools.core.Equilibrium(length_unit='m', tspline=False, monotonic=False, verbose=True)
Bases: object
```

Abstract class of data handling object for magnetic reconstruction outputs.

Defines the mapping routines and method fingerprints necessary. Each variable or set of variables is recovered with a corresponding getter method. Essential data for mapping are pulled on initialization (psirz grid, for example) to frontload overhead. Additional data are pulled at the first request and stored for subsequent usage.

Note: This abstract class should not be used directly. Device- and code- specific subclasses are set up to account for inter-device/-code differences in data storage.

Parameters

• **length_unit** – Sets the base unit used for any quantity whose dimensions are length to any power. Valid options are:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'de-	whatever the default in the tree is (no conversion is performed, units may
fault'	be inconsistent)

Default is 'm' (all units taken and returned in meters).

- **tspline** Sets whether or not interpolation in time is performed using a tricubic spline or nearest-neighbor interpolation. Tricubic spline interpolation requires at least four complete equilibria at different times. It is also assumed that they are functionally correlated, and that parameters do not vary out of their boundaries (derivative = 0 boundary condition). Default is False (use nearest-neighbor interpolation).
- **monotonic** Sets whether or not the "monotonic" form of time window finding is used. If True, the timebase must be monotonically increasing. Default is False (use slower, safer method).
- verbose Allows or blocks console readout during operation. Defaults to True, displaying
 useful information for the user. Set to False for quiet usage or to avoid console clutter for
 multiple instances.

Kwtype length_unit String

Kwtype tspline Boolean

Kwtype monotonic Boolean

Kwtype verbose Boolean

Raises

- ValueError If *length_unit* is not a valid unit specifier.
- ValueError If tspline is True but module trispline did not load successfully.

rho2rho(origin, destination, *args, **kwargs)

Convert from one coordinate to another.

Parameters

• **origin** (String) – Indicates which coordinates the data are given in. Valid options are:

RZ	R,Z coordinates
psinorm	Normalized poloidal flux
phinorm	Normalized toroidal flux
volnorm	Normalized volume
Rmid	Midplane major radius
r/a	Normalized minor radius

Additionally, each valid option may be prepended with 'sqrt' to specify the square root of the desired unit.

• **destination** (String) – Indicates which coordinates to convert to. Valid options are:

psinorm	Normalized poloidal flux
phinorm	Normalized toroidal flux
volnorm	Normalized volume
Rmid	Midplane major radius
r/a	Normalized minor radius

Additionally, each valid option may be prepended with 'sqrt' to specify the square root of the desired unit.

- **rho** (*Array-like or scalar float*) Values of the starting coordinate to map to the new coordinate. Will be two arguments *R*, *Z* if *origin* is 'RZ'.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *rho*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *rho* (or the meshgrid of *R* and *Z* if *make_grid* is True).
- **sqrt** Set to True to return the square root of *rho*. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *rho* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *rho* or be a scalar. Default is True (evaluate ALL *rho* at EACH element in *t*).
- make_grid Only applicable if *origin* is 'RZ'. Set to True to pass R and Z through scipy.meshgrid() before evaluating. If this is set to True, R and Z must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **rho** Set to True to return r/a (normalized minor radius) instead of Rmid when *destination* is Rmid. Default is False (return major radius, Rmid).
- **length_unit** Length unit that quantities are given/returned in, as applicable. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from psinorm to Rmid/phinorm/volnorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype make_grid Boolean

Kwtype rho Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

rho or (*rho*, *time_idxs*)

- **rho** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *rho*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Raises ValueError – If *origin* is not one of the supported values.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value at r/a=0.6, t=0.26s:

```
psi_val = Eq_instance.rho2rho('r/a', 'psinorm', 0.6, 0.26)
```

Find psinorm values at r/a points 0.6 and 0.8 at the single time t=0.26s:

```
psi_arr = Eq_instance.rho2rho('r/a', 'psinorm', [0.6, 0.8], 0.26)
```

Find psinorm values at r/a of 0.6 at times t=[0.2s, 0.3s]:

```
psi\_arr = Eq\_instance.rho2rho('r/a', 'psinorm', 0.6, [0.2, 0.3])
```

Find psinorm values at (r/a, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
psi_arr = Eq_instance.rho2rho('r/a', 'psinorm', [0.6, 0.5], [0.2, 0.3], each_t=False)
```

rz2psi (*R*, *Z*, *t*, *return_t=False*, *make_grid=False*, *each_t=True*, *length_unit=1*) Converts the passed R, Z, t arrays to psi (unnormalized poloidal flux) values.

- **R** (*Array-like or scalar float*) Values of the radial coordinate to map to poloidal flux. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *Z* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *R* must have exactly one dimension.
- **Z** (*Array-like or scalar float*) Values of the vertical coordinate to map to poloidal flux. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *R* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *Z* must have exactly one dimension.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R*, *Z*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R* and *Z* (or their meshgrid if *make_grid* is True).
- each_t When True, the elements in *R*, *Z* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *R* and *Z* or be a scalar. Default is True (evaluate ALL *R*, *Z* at EACH element in *t*).
- make_grid Set to True to pass *R* and *Z* through scipy.meshgrid() before evaluating. If this is set to True, *R* and *Z* must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **length_unit** Length unit that *R*, *Z* are given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (use meters).

• **return_t** – Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype each_t Boolean

Kwtype make_grid Boolean

Kwtype length_unit String or 1

Kwtype return_t Boolean

Returns

psi or (psi, time idxs)

- **psi** (*Array or scalar float*) The unnormalized poloidal flux. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned. If *R* and *Z* both have the same shape then *psi* has this shape as well, unless the *make_grid* keyword was True, in which case *psi* has shape (len(*Z*), len(*R*)).
- **time_idxs** (Array with same shape as *psi*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psi value at R=0.6m, Z=0.0m, t=0.26s:

```
psi_val = Eq_instance.rz2psi(0.6, 0, 0.26)
```

Find psi values at (R, Z) points (0.6m, 0m) and (0.8m, 0m) at the single time t=0.26s. Note that the Z vector must be fully specified, even if the values are all the same:

```
psi_arr = Eq_instance.rz2psi([0.6, 0.8], [0, 0], 0.26)
```

Find psi values at (R, Z) points (0.6m, 0m) at times t=[0.2s, 0.3s]:

```
psi_arr = Eq_instance.rz2psi(0.6, 0, [0.2, 0.3])
```

Find psi values at (R, Z, t) points (0.6m, 0m, 0.2s) and (0.5m, 0.2m, 0.3s):

```
psi_arr = Eq_instance.rz2psi([0.6, 0.5], [0, 0.2], [0.2, 0.3], each_t=False)
```

Find psi values on grid defined by 1D vector of radial positions R and 1D vector of vertical positions Z at time t=0.2s:

```
psi_mat = Eq_instance.rz2psi(R, Z, 0.2, make_grid=True)
```

rz2psinorm (*R*, *Z*, *t*, return_t=False, sqrt=False, make_grid=False, each_t=True, length_unit=1) Calculates the normalized poloidal flux at the given (R, Z, t).

Uses the definition:

$$\texttt{psi_norm} = \frac{\psi - \psi(0)}{\psi(a) - \psi(0)}$$

- **R** (*Array-like or scalar float*) Values of the radial coordinate to map to psinorm. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *Z* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *R* must have exactly one dimension.
- **Z** (*Array-like or scalar float*) Values of the vertical coordinate to map to psinorm. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *R* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *Z* must have exactly one dimension.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R*, *Z*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R* and *Z* (or their meshgrid if *make_grid* is True).
- sqrt Set to True to return the square root of psinorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit rz2rho.pro. Default is False.
- each_t When True, the elements in R, Z are evaluated at each value in t. If True, t must have only one dimension (or be a scalar). If False, t must match the shape of R and Z or be a scalar. Default is True (evaluate ALL R, Z at EACH element in t).

- make_grid Set to True to pass *R* and *Z* through scipy.meshgrid() before evaluating. If this is set to True, *R* and *Z* must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **length_unit** Length unit that *R*, *Z* are given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (use meters).

• **return_t** – Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each t Boolean

Kwtype make_grid Boolean

Kwtype length_unit String or 1

Kwtype return_t Boolean

Returns

psinorm or (psinorm, time idxs)

- **psinorm** (*Array or scalar float*) The normalized poloidal flux. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned. If *R* and *Z* both have the same shape then *psinorm* has this shape as well, unless the *make_grid* keyword was True, in which case *psinorm* has shape (len(*Z*), len(*R*)).
- **time_idxs** (Array with same shape as *psinorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value at R=0.6m, Z=0.0m, t=0.26s:

```
psi_val = Eq_instance.rz2psinorm(0.6, 0, 0.26)
```

Find psinorm values at (R, Z) points (0.6m, 0m) and (0.8m, 0m) at the single time t=0.26s. Note that the Z vector must be fully specified, even if the values are all the same:

```
psi_arr = Eq_instance.rz2psinorm([0.6, 0.8], [0, 0], 0.26)
```

Find psinorm values at (R, Z) points (0.6m, 0m) at times t=[0.2s, 0.3s]:

```
psi\_arr = Eq\_instance.rz2psinorm(0.6, 0, [0.2, 0.3])
```

Find psinorm values at (R, Z, t) points (0.6m, 0m, 0.2s) and (0.5m, 0.2m, 0.3s):

```
psi_arr = Eq_instance.rz2psinorm([0.6, 0.5], [0, 0.2], [0.2, 0.3], each_t=False)
```

Find psinorm values on grid defined by 1D vector of radial positions R and 1D vector of vertical positions Z at time t=0.2s:

```
psi_mat = Eq_instance.rz2psinorm(R, Z, 0.2, make_grid=True)
```

rz2phinorm(*args, **kwargs)

Calculates the normalized toroidal flux.

Uses the definitions:

$$\mathrm{phi} = \int q(\psi) \, d\psi$$

$$\mathrm{phi} \mathrm{_norm} = \frac{\phi}{\phi(a)}$$

This is based on the IDL version efit rz2rho.pro by Steve Wolfe.

- **R** (*Array-like or scalar float*) Values of the radial coordinate to map to phinorm. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *Z* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *R* must have exactly one dimension.
- **Z** (*Array-like or scalar float*) Values of the vertical coordinate to map to phinorm. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *R* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *Z* must have exactly one dimension.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R*, *Z*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R* and *Z* (or their meshgrid if *make_grid* is True).
- sqrt Set to True to return the square root of phinorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit rz2rho.pro. Default is False.
- each_t When True, the elements in *R*, *Z* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *R* and *Z* or be a scalar. Default is True (evaluate ALL *R*, *Z* at EACH element in *t*).
- make_grid Set to True to pass *R* and *Z* through scipy.meshgrid() before evaluating. If this is set to True, *R* and *Z* must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **length_unit** Length unit that *R*, *Z* are given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- **kind** Specifies the type of interpolation to be performed in getting from psinorm to phinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each t Boolean

Kwtype make_grid Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

phinorm or (phinorm, time_idxs)

- **phinorm** (*Array or scalar float*) The normalized toroidal flux. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned. If *R* and *Z* both have the same shape then *phinorm* has this shape as well, unless the *make_grid* keyword was True, in which case *phinorm* has shape (len(*Z*), len(*R*)).
- **time_idxs** (Array with same shape as *phinorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single phinorm value at R=0.6m, Z=0.0m, t=0.26s:

```
phi_val = Eq_instance.rz2phinorm(0.6, 0, 0.26)
```

Find phinorm values at (R, Z) points (0.6m, 0m) and (0.8m, 0m) at the single time t=0.26s. Note that the Z vector must be fully specified, even if the values are all the same:

```
phi_arr = Eq_instance.rz2phinorm([0.6, 0.8], [0, 0], 0.26)
```

Find phinorm values at (R, Z) points (0.6m, 0m) at times t=[0.2s, 0.3s]:

```
phi_arr = Eq_instance.rz2phinorm(0.6, 0, [0.2, 0.3])
```

Find phinorm values at (R, Z, t) points (0.6m, 0m, 0.2s) and (0.5m, 0.2m, 0.3s):

```
phi_arr = Eq_instance.rz2phinorm([0.6, 0.5], [0, 0.2], [0.2, 0.3], each_t=False)
```

Find phinorm values on grid defined by 1D vector of radial positions R and 1D vector of vertical positions Z at time t=0.2s:

```
phi_mat = Eq_instance.rz2phinorm(R, Z, 0.2, make_grid=True)
```

```
rz2volnorm(*args, **kwargs)
```

Calculates the normalized flux surface volume.

Based on the IDL version efit_rz2rho.pro by Steve Wolfe.

- **R** (*Array-like or scalar float*) Values of the radial coordinate to map to volnorm. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *Z* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *R* must have exactly one dimension.
- **Z** (*Array-like or scalar float*) Values of the vertical coordinate to map to volnorm. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *R* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *Z* must have exactly one dimension.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R*, *Z*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R* and *Z* (or their meshgrid if *make grid* is True).
- **sqrt** Set to True to return the square root of volnorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *R*, *Z* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *R* and *Z* or be a scalar. Default is True (evaluate ALL *R*, *Z* at EACH element in *t*).
- make_grid Set to True to pass *R* and *Z* through scipy.meshgrid() before evaluating. If this is set to True, *R* and *Z* must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **length_unit** Length unit that *R*, *Z* are given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- **kind** Specifies the type of interpolation to be performed in getting from psinorm to volnorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each t Boolean

Kwtype make_grid Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

volnorm or (volnorm, time_idxs)

- **volnorm** (*Array or scalar float*) The normalized volume. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned. If *R* and *Z* both have the same shape then *volnorm* has this shape as well, unless the *make_grid* keyword was True, in which case *volnorm* has shape (len(*Z*), len(*R*)).
- **time_idxs** (Array with same shape as *volnorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single volnorm value at R=0.6m, Z=0.0m, t=0.26s:

```
psi_val = Eq_instance.rz2volnorm(0.6, 0, 0.26)
```

Find volnorm values at (R, Z) points (0.6m, 0m) and (0.8m, 0m) at the single time t=0.26s. Note that the Z vector must be fully specified, even if the values are all the same:

```
vol_arr = Eq_instance.rz2volnorm([0.6, 0.8], [0, 0], 0.26)
```

Find volnorm values at (R, Z) points (0.6m, 0m) at times t=[0.2s, 0.3s]:

```
vol_arr = Eq_instance.rz2volnorm(0.6, 0, [0.2, 0.3])
```

Find volnorm values at (R, Z, t) points (0.6m, 0m, 0.2s) and (0.5m, 0.2m, 0.3s):

```
vol_arr = Eq_instance.rz2volnorm([0.6, 0.5], [0, 0.2], [0.2, 0.3], each_t=False)
```

Find volnorm values on grid defined by 1D vector of radial positions R and 1D vector of vertical positions Z at time t=0.2s:

```
vol_mat = Eq_instance.rz2volnorm(R, Z, 0.2, make_grid=True)
```

```
rz2rmid(*args, **kwargs)
```

Maps the given points to the outboard midplane major radius, Rmid.

Based on the IDL version efit_rz2rmid.pro by Steve Wolfe.

- **R** (*Array-like or scalar float*) Values of the radial coordinate to map to Rmid. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *Z* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *R* must have exactly one dimension.
- **Z** (*Array-like or scalar float*) Values of the vertical coordinate to map to Rmid. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *R* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *Z* must have exactly one dimension.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R*, *Z*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R* and *Z* (or their meshgrid if *make grid* is True).
- **sqrt** Set to True to return the square root of Rmid. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *R*, *Z* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *R* and *Z* or be a scalar. Default is True (evaluate ALL *R*, *Z* at EACH element in *t*).
- make_grid Set to True to pass *R* and *Z* through scipy.meshgrid() before evaluating. If this is set to True, *R* and *Z* must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **rho** Set to True to return r/a (normalized minor radius) instead of Rmid. Default is False (return major radius, Rmid).
- **length_unit** Length unit that *R*, *Z* are given in, AND that *Rmid* is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from psinorm to Rmid. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype make_grid Boolean

Kwtype rho Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

Rmid or (*Rmid*, time_idxs)

- **Rmid** (*Array or scalar float*) The outboard midplan major radius. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned. If *R* and *Z* both have the same shape then *Rmid* has this shape as well, unless the *make_grid* keyword was True, in which case *Rmid* has shape (len(*Z*), len(*R*)).
- **time_idxs** (Array with same shape as *Rmid*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single Rmid value at R=0.6m, Z=0.0m, t=0.26s:

```
R_{mid_val} = Eq_{instance.rz2rmid(0.6, 0, 0.26)}
```

Find R_mid values at (R, Z) points (0.6m, 0m) and (0.8m, 0m) at the single time t=0.26s. Note that the Z vector must be fully specified, even if the values are all the same:

```
R_{mid_arr} = Eq_{instance.rz2rmid([0.6, 0.8], [0, 0], 0.26)}
```

Find Rmid values at (R, Z) points (0.6m, 0m) at times t=[0.2s, 0.3s]:

```
R_{mid\_arr} = Eq_{instance.rz2rmid(0.6, 0, [0.2, 0.3])
```

Find Rmid values at (R, Z, t) points (0.6m, 0m, 0.2s) and (0.5m, 0.2m, 0.3s):

```
R_mid_arr = Eq_instance.rz2rmid([0.6, 0.5], [0, 0.2], [0.2, 0.3], each_t=False)
```

Find Rmid values on grid defined by 1D vector of radial positions R and 1D vector of vertical positions Z at time t=0.2s:

```
R_mid_mat = Eq_instance.rz2rmid(R, Z, 0.2, make_grid=True)
```

```
rz2roa (*args, **kwargs)
```

Maps the given points to the normalized minor radius, r/a.

Based on the IDL version efit_rz2rmid.pro by Steve Wolfe.

- **R** (*Array-like or scalar float*) Values of the radial coordinate to map to r/a. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *Z* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *R* must have exactly one dimension.
- **Z** (*Array-like or scalar float*) Values of the vertical coordinate to map to r/a. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *R* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *Z* must have exactly one dimension.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R*, *Z*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R* and *Z* (or their meshgrid if *make_grid* is True).
- sqrt Set to True to return the square root of r/a. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *R*, *Z* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *R* and *Z* or be a scalar. Default is True (evaluate ALL *R*, *Z* at EACH element in *t*).
- make_grid Set to True to pass *R* and *Z* through scipy.meshgrid() before evaluating. If this is set to True, *R* and *Z* must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- length_unit Length unit that R, Z are given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from psinorm to Rmid. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype make grid Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

roa or (roa, time_idxs)

- **roa** (*Array or scalar float*) The normalized minor radius. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned. If *R* and *Z* both have the same shape then *roa* has this shape as well, unless the *make_grid* keyword was True, in which case *roa* has shape (len(*Z*), len(*R*)).
- **time_idxs** (Array with same shape as *roa*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class

Find single r/a value at R=0.6m, Z=0.0m, t=0.26s:

```
roa_val = Eq_instance.rz2roa(0.6, 0, 0.26)
```

Find r/a values at (R, Z) points (0.6m, 0m) and (0.8m, 0m) at the single time t=0.26s. Note that the Z vector must be fully specified, even if the values are all the same:

```
roa_arr = Eq_instance.rz2roa([0.6, 0.8], [0, 0], 0.26)
```

Find r/a values at (R, Z) points (0.6m, 0m) at times t=[0.2s, 0.3s]:

```
roa_arr = Eq_instance.rz2roa(0.6, 0, [0.2, 0.3])
```

Find r/a values at (R, Z, t) points (0.6m, 0m, 0.2s) and (0.5m, 0.2m, 0.3s):

```
roa_arr = Eq_instance.rz2roa([0.6, 0.5], [0, 0.2], [0.2, 0.3], each_t=False)
```

Find r/a values on grid defined by 1D vector of radial positions R and 1D vector of vertical positions Z at time t=0.2s:

```
roa_mat = Eq_instance.rz2roa(R, Z, 0.2, make_grid=True)
```

```
rz2rho (method, *args, **kwargs)
```

Convert the passed (R, Z, t) coordinates into one of several coordinates.

Parameters

• method (String) – Indicates which coordinates to convert to. Valid options are:

psinorm	Normalized poloidal flux
phinorm	Normalized toroidal flux
volnorm	Normalized volume
Rmid	Midplane major radius
r/a	Normalized minor radius

Additionally, each valid option may be prepended with 'sqrt' to specify the square root of the desired unit.

- **R** (*Array-like or scalar float*) Values of the radial coordinate to map to *rho*. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *Z* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *R* must have exactly one dimension.
- **Z** (*Array-like or scalar float*) Values of the vertical coordinate to map to *rho*. If *R* and *Z* are both scalar values, they are used as the coordinate pair for all of the values in *t*. Must have the same shape as *R* unless the *make_grid* keyword is set. If the *make_grid* keyword is True, *Z* must have exactly one dimension.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R*, *Z*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R* and *Z* (or their meshgrid if *make_grid* is True).
- **sqrt** Set to True to return the square root of *rho*. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit rz2rho.pro. Default is False.
- each_t When True, the elements in *R*, *Z* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *R* and *Z* or be a scalar. Default is True (evaluate ALL *R*, *Z* at EACH element in *t*).
- make_grid Set to True to pass R and Z through scipy.meshgrid() before evaluating. If this is set to True, R and Z must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **rho** Set to True to return r/a (normalized minor radius) instead of Rmid when *destination* is Rmid. Default is False (return major radius, Rmid).

• **length_unit** – Length unit that *R*, *Z* are given in, AND that *Rmid* is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from psinorm to Rmid/phinorm/volnorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype make_grid Boolean

Kwtype rho Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

rho or (rho, time_idxs)

- **rho** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *rho*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Raises ValueError – If *method* is not one of the supported values.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value at R=0.6m, Z=0.0m, t=0.26s:

```
psi_val = Eq_instance.rz2rho('psinorm', 0.6, 0, 0.26)
```

Find psinorm values at (R, Z) points (0.6m, 0m) and (0.8m, 0m) at the single time t=0.26s. Note that the Z vector must be fully specified, even if the values are all the same:

```
psi_arr = Eq_instance.rz2rho('psinorm', [0.6, 0.8], [0, 0], 0.26)
```

Find psinorm values at (R, Z) points (0.6m, 0m) at times t=[0.2s, 0.3s]:

```
psi_arr = Eq_instance.rz2rho('psinorm', 0.6, 0, [0.2, 0.3])
```

Find psinorm values at (R, Z, t) points (0.6m, 0m, 0.2s) and (0.5m, 0.2m, 0.3s):

```
psi_arr = Eq_instance.rz2rho('psinorm', [0.6, 0.5], [0, 0.2], [0.2, 0.3], each_t=False)
```

Find psinorm values on grid defined by 1D vector of radial positions R and 1D vector of vertical positions Z at time t=0.2s:

```
psi_mat = Eq_instance.rz2rho('psinorm', R, Z, 0.2, make_grid=True)
```

rmid2roa (*R_mid*, *t*, *each_t=True*, *return_t=False*, *sqrt=False*, *time_idxs=None*, *length_unit=1*) Convert the passed (R mid, t) coordinates into r/a.

Parameters

- **R_mid** (*Array-like or scalar float*) Values of the outboard midplane major radius to map to r/a.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R_mid*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R_mid*.
- sqrt Set to True to return the square root of r/a. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *R_mid* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *R_mid* or be a scalar. Default is True (evaluate ALL *R_mid* at EACH element in *t*).
- **length_unit** Length unit that *R_mid* is given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length unit is 1 or None, meters are assumed. The default value is 1 (use meters).

• **return_t** – Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype length_unit String or 1

Kwtype return_t Boolean

Returns

roa or (roa, time idxs)

- roa (*Array or scalar float*) Normalized midplane minor radius. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *roa*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single r/a value at R mid=0.6m, t=0.26s:

```
roa_val = Eq_instance.rmid2roa(0.6, 0.26)
```

Find roa values at R_mid points 0.6m and 0.8m at the single time t=0.26s.:

```
roa_arr = Eq_instance.rmid2roa([0.6, 0.8], 0.26)
```

Find roa values at R_{mid} of 0.6m at times t=[0.2s, 0.3s]:

```
roa_arr = Eq_instance.rmid2roa(0.6, [0.2, 0.3])
```

Find r/a values at (R_mid, t) points (0.6m, 0.2s) and (0.5m, 0.3s):

```
roa_arr = Eq_instance.rmid2roa([0.6, 0.5], [0.2, 0.3], each_t=False)
```

rmid2psinorm(R_mid, t, **kwargs)

Calculates the normalized poloidal flux corresponding to the passed R_mid (mapped outboard midplane major radius) values.

Parameters

- **R_mid** (*Array-like or scalar float*) Values of the outboard midplane major radius to map to psinorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R_mid*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R_mid*.
- sqrt Set to True to return the square root of psinorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *R_mid* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *R_mid* or be a scalar. Default is True (evaluate ALL *R_mid* at EACH element in *t*).
- **length_unit** Length unit that *R_mid* is given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from Rmid to psinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

psinorm or (*psinorm*, *time_idxs*)

- **psinorm** (*Array or scalar float*) Normalized poloidal flux. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *psinorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value for Rmid=0.7m, t=0.26s:

```
psinorm_val = Eq_instance.rmid2psinorm(0.7, 0.26)
```

Find psinorm values at R_mid values of 0.5m and 0.7m at the single time t=0.26s:

```
psinorm_arr = Eq_instance.rmid2psinorm([0.5, 0.7], 0.26)
```

Find psinorm values at R_mid=0.5m at times t=[0.2s, 0.3s]:

```
psinorm_arr = Eq_instance.rmid2psinorm(0.5, [0.2, 0.3])
```

Find psinorm values at (R_mid, t) points (0.6m, 0.2s) and (0.5m, 0.3s):

```
psinorm_arr = Eq_instance.rmid2psinorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

rmid2phinorm(*args, **kwargs)

Calculates the normalized toroidal flux.

Uses the definitions:

$$\label{eq:phi_norm} \begin{split} \mathrm{phi} &= \int q(\psi)\,d\psi \\ \mathrm{phi_norm} &= \frac{\phi}{\phi(a)} \end{split}$$

This is based on the IDL version efit_rz2rho.pro by Steve Wolfe.

Parameters

- **R_mid** (*Array-like or scalar float*) Values of the outboard midplane major radius to map to phinorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R_mid*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R_mid*.
- **sqrt** Set to True to return the square root of phinorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *R_mid* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *R_mid* or be a scalar. Default is True (evaluate ALL *R_mid* at EACH element in *t*).
- **length_unit** Length unit that *R_mid* is given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (use meters).

• kind – Specifies the type of interpolation to be performed in getting from Rmid to psinorm and psinorm to phinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.

• **return_t** – Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

phinorm or (phinorm, time_idxs)

- **phinorm** (*Array or scalar float*) Normalized toroidal flux. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *phinorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single phinorm value at R_mid=0.6m, t=0.26s:

```
phi_val = Eq_instance.rmid2phinorm(0.6, 0.26)
```

Find phinorm values at R_mid points 0.6m and 0.8m at the single time t=0.26s:

```
phi_arr = Eq_instance.rmid2phinorm([0.6, 0.8], 0.26)
```

Find phinorm values at R_mid point 0.6m at times t=[0.2s, 0.3s]:

```
phi_arr = Eq_instance.rmid2phinorm(0.6, [0.2, 0.3])
```

Find phinorm values at (R, t) points (0.6m, 0.2s) and (0.5m, 0.3s):

```
phi_arr = Eq_instance.rmid2phinorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

```
rmid2volnorm(*args, **kwargs)
```

Calculates the normalized flux surface volume.

Based on the IDL version efit_rz2rho.pro by Steve Wolfe.

Parameters

- **R_mid** (*Array-like or scalar float*) Values of the outboard midplane major radius to map to volnorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R_mid*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R_mid*.

- **sqrt** Set to True to return the square root of volnorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *R_mid* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *R_mid* or be a scalar. Default is True (evaluate ALL *R_mid* at EACH element in *t*).
- **length_unit** Length unit that *R_mid* is given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from Rmid to psinorm and psinorm to volnorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

volnorm or (*volnorm*, *time_idxs*)

- **volnorm** (*Array or scalar float*) Normalized volume. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *volnorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single volnorm value at R_mid=0.6m, t=0.26s:

```
vol_val = Eq_instance.rmid2volnorm(0.6, 0.26)
```

Find volnorm values at R_mid points 0.6m and 0.8m at the single time t=0.26s:

```
vol_arr = Eq_instance.rmid2volnorm([0.6, 0.8], 0.26)
```

Find volnorm values at R_mid points 0.6m at times t=[0.2s, 0.3s]:

```
vol_arr = Eq_instance.rmid2volnorm(0.6, [0.2, 0.3])
```

Find volnorm values at (R_mid, t) points (0.6m, 0.2s) and (0.5m, 0.3s):

```
vol_arr = Eq_instance.rmid2volnorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

```
rmid2rho (method, R_mid, t, **kwargs)
```

Convert the passed (R mid, t) coordinates into one of several coordinates.

Parameters

• method (String) – Indicates which coordinates to convert to. Valid options are:

psinorm	Normalized poloidal flux
phinorm	Normalized toroidal flux
volnorm	Normalized volume
r/a	Normalized minor radius

Additionally, each valid option may be prepended with 'sqrt' to specify the square root of the desired unit.

- **R_mid** (*Array-like or scalar float*) Values of the outboard midplane major radius to map to rho.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *R_mid*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *R_mid*.
- sqrt Set to True to return the square root of rho. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *R_mid* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *R_mid* or be a scalar. Default is True (evaluate ALL *R_mid* at EACH element in *t*).
- **length_unit** Length unit that *R_mid* is given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from Rmid to psinorm and psinorm to volnorm or phinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

rho or (rho, time_idxs)

- **rho** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *rho*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value at R_mid=0.6m, t=0.26s:

```
psi_val = Eq_instance.rmid2rho('psinorm', 0.6, 0.26)
```

Find psinorm values at R_mid points 0.6m and 0.8m at the single time t=0.26s.:

```
psi_arr = Eq_instance.rmid2rho('psinorm', [0.6, 0.8], 0.26)
```

Find psinorm values at R_{mid} of 0.6m at times t=[0.2s, 0.3s]:

```
psi_arr = Eq_instance.rmid2rho('psinorm', 0.6, [0.2, 0.3])
```

Find psinorm values at (R_mid, t) points (0.6m, 0.2s) and (0.5m, 0.3s):

```
psi_arr = Eq_instance.rmid2rho('psinorm', [0.6, 0.5], [0.2, 0.3], each_t=False)
```

roa2rmid (roa, t, each_t=True, return_t=False, time_idxs=None, length_unit=1)
Convert the passed (r/a, t) coordinates into Rmid.

Parameters

- roa (Array-like or scalar float) Values of the normalized minor radius to map to Rmid.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *roa*. If the *each_t* keyword is True, then *t* must be scalar

or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *roa*.

- each_t When True, the elements in *roa* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *roa* or be a scalar. Default is True (evaluate ALL *roa* at EACH element in *t*).
- **length_unit** Length unit that *Rmid* is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (use meters).

• **return_t** – Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype each_t Boolean

Kwtype length_unit String or 1

Kwtype return_t Boolean

Returns

Rmid or (Rmid, time idxs)

- **Rmid** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *Rmid*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single R mid value at r/a=0.6, t=0.26s:

```
R_mid_val = Eq_instance.roa2rmid(0.6, 0.26)
```

Find R_mid values at r/a points 0.6 and 0.8 at the single time t=0.26s.:

```
R_{mid\_arr} = Eq_{instance.roa2rmid([0.6, 0.8], 0.26)}
```

Find R_mid values at r/a of 0.6 at times t=[0.2s, 0.3s]:

```
R_{mid_arr} = Eq_{instance.roa2rmid(0.6, [0.2, 0.3])}
```

Find R_mid values at (roa, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
R_{mid}_{arr} = Eq_{instance.roa2rmid([0.6, 0.5], [0.2, 0.3], each_t=False)}
```

roa2psinorm(*args, **kwargs)

Convert the passed (r/a, t) coordinates into psinorm.

Parameters

- roa (Array-like or scalar float) Values of the normalized minor radius to map to psinorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *roa*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *roa*.
- sqrt Set to True to return the square root of psinorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *roa* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *roa* or be a scalar. Default is True (evaluate ALL *roa* at EACH element in *t*).
- kind Specifies the type of interpolation to be performed in getting from Rmid to psinorm and psinorm to volnorm or phinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

psinorm or (*psinorm*, *time_idxs*)

- **psinorm** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *psinorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value at r/a=0.6, t=0.26s:

```
psinorm_val = Eq_instance.roa2psinorm(0.6, 0.26)

Find psinorm values at r/a points 0.6 and 0.8 at the single time t=0.26s.:

psinorm_arr = Eq_instance.roa2psinorm([0.6, 0.8], 0.26)

Find psinorm values at r/a of 0.6 at times t=[0.2s, 0.3s]:

psinorm_arr = Eq_instance.roa2psinorm(0.6, [0.2, 0.3])

Find psinorm values at (roa, t) points (0.6, 0.2s) and (0.5, 0.3s):

psinorm_arr = Eq_instance.roa2psinorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

roa2phinorm(*args, **kwargs)

Convert the passed (r/a, t) coordinates into phinorm.

Parameters

- roa (Array-like or scalar float) Values of the normalized minor radius to map to phinorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *roa*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *roa*.
- **sqrt** Set to True to return the square root of phinorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- **each_t** When True, the elements in *roa* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *roa* or be a scalar. Default is True (evaluate ALL *roa* at EACH element in *t*).
- kind Specifies the type of interpolation to be performed in getting from Rmid to psinorm and psinorm to volnorm or phinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each t Boolean

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

phinorm or (phinorm, time idxs)

• **phinorm** (*Array or scalar float*) - The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.

• **time_idxs** (Array with same shape as *phinorm*) - The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single phinorm value at r/a=0.6, t=0.26s:

```
phinorm_val = Eq_instance.roa2phinorm(0.6, 0.26)
```

Find phinorm values at r/a points 0.6 and 0.8 at the single time t=0.26s.:

```
phinorm_arr = Eq_instance.roa2phinorm([0.6, 0.8], 0.26)
```

Find phinorm values at r/a of 0.6 at times t=[0.2s, 0.3s]:

```
phinorm_arr = Eq_instance.roa2phinorm(0.6, [0.2, 0.3])
```

Find phinorm values at (roa, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
phinorm_arr = Eq_instance.roa2phinorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

```
roa2volnorm(*args, **kwargs)
```

Convert the passed (r/a, t) coordinates into volnorm.

Parameters

- roa (Array-like or scalar float) Values of the normalized minor radius to map to volnorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *roa*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *roa*.
- **sqrt** Set to True to return the square root of volnorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *roa* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *roa* or be a scalar. Default is True (evaluate ALL *roa* at EACH element in *t*).
- **kind** Specifies the type of interpolation to be performed in getting from Rmid to psinorm and psinorm to volnorm or phinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

volnorm or (volnorm, time_idxs)

- **volnorm** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *volnorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single volnorm value at r/a=0.6, t=0.26s:

```
volnorm_val = Eq_instance.roa2volnorm(0.6, 0.26)
```

Find volnorm values at r/a points 0.6 and 0.8 at the single time t=0.26s.:

```
volnorm_arr = Eq_instance.roa2volnorm([0.6, 0.8], 0.26)
```

Find volnorm values at r/a of 0.6 at times t=[0.2s, 0.3s]:

```
volnorm_arr = Eq_instance.roa2volnorm(0.6, [0.2, 0.3])
```

Find volnorm values at (roa, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
volnorm_arr = Eq_instance.roa2volnorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

```
roa2rho (method, *args, **kwargs)
```

Convert the passed (r/a, t) coordinates into one of several coordinates.

Parameters

• method (String) – Indicates which coordinates to convert to. Valid options are:

psinorm	Normalized poloidal flux
phinorm	Normalized toroidal flux
volnorm	Normalized volume
Rmid	Midplane major radius

Additionally, each valid option may be prepended with 'sqrt' to specify the square root of the desired unit.

- roa (Array-like or scalar float) Values of the normalized minor radius to map to rho.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *roa*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *roa*.
- sqrt Set to True to return the square root of rho. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.

- each_t When True, the elements in *roa* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *roa* or be a scalar. Default is True (evaluate ALL *roa* at EACH element in *t*).
- **length_unit** Length unit that *Rmid* is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from Rmid to psinorm and psinorm to volnorm or phinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each t Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

rho or (rho, time_idxs)

- **rho** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *rho*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value at r/a=0.6, t=0.26s:

```
psi_val = Eq_instance.roa2rho('psinorm', 0.6, 0.26)
```

Find psinorm values at r/a points 0.6 and 0.8 at the single time t=0.26s:

```
psi_arr = Eq_instance.roa2rho('psinorm', [0.6, 0.8], 0.26)
```

Find psinorm values at r/a of 0.6 at times t=[0.2s, 0.3s]:

```
psi_arr = Eq_instance.roa2rho('psinorm', 0.6, [0.2, 0.3])
```

Find psinorm values at (r/a, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
psi_arr = Eq_instance.roa2rho('psinorm', [0.6, 0.5], [0.2, 0.3], each_t=False)
```

psinorm2rmid (psi_norm, t, **kwargs)

Calculates the outboard R_mid location corresponding to the passed psinorm (normalized poloidal flux) values.

Parameters

- **psi_norm** (*Array-like or scalar float*) Values of the normalized poloidal flux to map to Rmid.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *psi_norm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *psi_norm*.
- sqrt Set to True to return the square root of Rmid. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in psi_norm are evaluated at each value in t. If True, t must have only one dimension (or be a scalar). If False, t must match the shape of psi_norm or be a scalar. Default is True (evaluate ALL psi_norm at EACH element in t).
- **rho** Set to True to return r/a (normalized minor radius) instead of Rmid. Default is False (return major radius, Rmid).
- **length_unit** Length unit that *Rmid* is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (use meters).

• kind – Specifies the type of interpolation to be performed in getting from psinorm to Rmid. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.

• **return_t** – Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype rho Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

Rmid or (*Rmid*, time_idxs)

- **Rmid** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *Rmid*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single R_mid value for psinorm=0.7, t=0.26s:

```
R_mid_val = Eq_instance.psinorm2rmid(0.7, 0.26)
```

Find R_mid values at psi_norm values of 0.5 and 0.7 at the single time t=0.26s:

```
R_{mid_arr} = Eq_{instance.psinorm2rmid([0.5, 0.7], 0.26)
```

Find R_mid values at psi_norm=0.5 at times t=[0.2s, 0.3s]:

```
R_mid_arr = Eq_instance.psinorm2rmid(0.5, [0.2, 0.3])
```

Find R_mid values at (psinorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
R_{mid} arr = Eq_instance.psinorm2rmid([0.6, 0.5], [0.2, 0.3], each_t=False)
```

```
psinorm2roa (psi_norm, t, **kwargs)
```

Calculates the normalized minor radius location corresponding to the passed psi_norm (normalized poloidal flux) values.

Parameters

- **psi_norm** (*Array-like or scalar float*) Values of the normalized poloidal flux to map to r/a.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *psi_norm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *psi_norm*.

- sqrt Set to True to return the square root of r/a. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in psi_norm are evaluated at each value in t. If True, t must have only one dimension (or be a scalar). If False, t must match the shape of psi_norm or be a scalar. Default is True (evaluate ALL psi_norm at EACH element in t).
- kind Specifies the type of interpolation to be performed in getting from psinorm to Rmid. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

roa or (roa, time idxs)

- roa (*Array or scalar float*) Normalized midplane minor radius. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *roa*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single r/a value for psinorm=0.7, t=0.26s:

```
roa_val = Eq_instance.psinorm2roa(0.7, 0.26)
```

Find r/a values at psi_norm values of 0.5 and 0.7 at the single time t=0.26s:

```
roa_arr = Eq_instance.psinorm2roa([0.5, 0.7], 0.26)
```

Find r/a values at psi norm=0.5 at times t=[0.2s, 0.3s]:

```
roa_arr = Eq_instance.psinorm2roa(0.5, [0.2, 0.3])
```

Find r/a values at (psinorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
roa_arr = Eq_instance.psinorm2roa([0.6, 0.5], [0.2, 0.3], each_t=False)
```

psinorm2volnorm (psi_norm, t, **kwargs)

Calculates the normalized volume corresponding to the passed psi_norm (normalized poloidal flux) values.

Parameters

- **psi_norm** (*Array-like or scalar float*) Values of the normalized poloidal flux to map to volnorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *psi_norm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *psi_norm*.
- sqrt Set to True to return the square root of volnorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *psi_norm* are evaluated at each value in t. If True, t must have only one dimension (or be a scalar). If False, t must match the shape of *psi_norm* or be a scalar. Default is True (evaluate ALL *psi_norm* at EACH element in t).
- kind Specifies the type of interpolation to be performed in getting from psinorm to volnorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

volnorm or (volnorm, time_idxs)

- **volnorm** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *volnorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single volnorm value for psinorm=0.7, t=0.26s:

```
volnorm_val = Eq_instance.psinorm2volnorm(0.7, 0.26)
```

Find volnorm values at psi_norm values of 0.5 and 0.7 at the single time t=0.26s:

```
volnorm_arr = Eq_instance.psinorm2volnorm([0.5, 0.7], 0.26)
```

Find volnorm values at psi_norm=0.5 at times t=[0.2s, 0.3s]:

```
volnorm_arr = Eq_instance.psinorm2volnorm(0.5, [0.2, 0.3])
```

Find volnorm values at (psinorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
volnorm_arr = Eq_instance.psinorm2volnorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

psinorm2phinorm(psi_norm, t, **kwargs)

Calculates the normalized toroidal flux corresponding to the passed psi_norm (normalized poloidal flux) values.

Parameters

- **psi_norm** (*Array-like or scalar float*) Values of the normalized poloidal flux to map to phinorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *psi_norm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *psi_norm*.
- sqrt Set to True to return the square root of phinorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in psi_norm are evaluated at each value in t. If True, t must have only one dimension (or be a scalar). If False, t must match the shape of psi_norm or be a scalar. Default is True (evaluate ALL psi_norm at EACH element in t).
- **kind** Specifies the type of interpolation to be performed in getting from psinorm to phinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype kind String or non-negative int

Kwtype return t Boolean

Returns

phinorm or (phinorm, time_idxs)

- **phinorm** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *phinorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single phinorm value for psinorm=0.7, t=0.26s:

```
phinorm_val = Eq_instance.psinorm2phinorm(0.7, 0.26)
```

Find phinorm values at psi_norm values of 0.5 and 0.7 at the single time t=0.26s:

```
phinorm_arr = Eq_instance.psinorm2phinorm([0.5, 0.7], 0.26)
```

Find phinorm values at psi_norm=0.5 at times t=[0.2s, 0.3s]:

```
phinorm_arr = Eq_instance.psinorm2phinorm(0.5, [0.2, 0.3])
```

Find phinorm values at (psinorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
phinorm_arr = Eq_instance.psinorm2phinorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

psinorm2rho (method, *args, **kwargs)

Convert the passed (psinorm, t) coordinates into one of several coordinates.

Parameters

• method (String) – Indicates which coordinates to convert to. Valid options are:

phinorm	Normalized toroidal flux
volnorm	Normalized volume
Rmid	Midplane major radius
r/a	Normalized minor radius

Additionally, each valid option may be prepended with 'sqrt' to specify the square root of the desired unit.

- **psi_norm** (*Array-like or scalar float*) Values of the normalized poloidal flux to map to rho.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *psi_norm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *psi_norm*.
- sqrt Set to True to return the square root of rho. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *psi_norm* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *psi_norm* or be a scalar. Default is True (evaluate ALL *psi_norm* at EACH element in *t*).
- **rho** Set to True to return r/a (normalized minor radius) instead of Rmid. Default is False (return major radius, Rmid).
- **length_unit** Length unit that *Rmid* is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from psinorm to Rmid/phinorm/volnorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype rho Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

rho or (rho, time_idxs)

- **rho** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *rho*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Raises ValueError – If *method* is not one of the supported values.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single phinorm value at psinorm=0.6, t=0.26s:

```
phi_val = Eq_instance.psinorm2rho('phinorm', 0.6, 0.26)
```

Find phinorm values at phinorm of 0.6 and 0.8 at the single time t=0.26s:

```
phi_arr = Eq_instance.psinorm2rho('phinorm', [0.6, 0.8], 0.26)
```

Find phinorm values at psinorm of 0.6 at times t=[0.2s, 0.3s]:

```
phi_arr = Eq_instance.psinorm2rho('phinorm', 0.6, [0.2, 0.3])
```

Find phinorm values at (psinorm, t) points (0.6, 0.2s) and (0.5m, 0.3s):

```
phi_arr = Eq_instance.psinorm2rho('phinorm', [0.6, 0.5], [0.2, 0.3], each_t=False)
```

phinorm2psinorm(phinorm, t, **kwargs)

Calculates the normalized poloidal flux corresponding to the passed phinorm (normalized toroidal flux) values.

Parameters

- **phinorm** (*Array-like or scalar float*) Values of the normalized toroidal flux to map to psinorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *phinorm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *phinorm*.
- sqrt Set to True to return the square root of psinorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *phinorm* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *phinorm* or be a scalar. Default is True (evaluate ALL *phinorm* at EACH element in *t*).
- **kind** Specifies the type of interpolation to be performed in getting from phinorm to psinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

psinorm or (*psinorm*, *time_idxs*)

- **psinorm** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *psinorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value for phinorm=0.7, t=0.26s:

```
psinorm_val = Eq_instance.phinorm2psinorm(0.7, 0.26)
```

Find psinorm values at phinorm values of 0.5 and 0.7 at the single time t=0.26s:

```
psinorm_arr = Eq_instance.phinorm2psinorm([0.5, 0.7], 0.26)
```

Find psinorm values at phinorm=0.5 at times t=[0.2s, 0.3s]:

```
psinorm_arr = Eq_instance.phinorm2psinorm(0.5, [0.2, 0.3])
```

Find psinorm values at (phinorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
psinorm_arr = Eq_instance.phinorm2psinorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

phinorm2volnorm(*args, **kwargs)

Calculates the normalized flux surface volume corresponding to the passed phinorm (normalized toroidal flux) values.

Parameters

- **phinorm** (*Array-like or scalar float*) Values of the normalized toroidal flux to map to volnorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *phinorm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *phinorm*.
- sqrt Set to True to return the square root of volnorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *phinorm* are evaluated at each value in t. If True, t must have only one dimension (or be a scalar). If False, t must match the shape of *phinorm* or be a scalar. Default is True (evaluate ALL *phinorm* at EACH element in t).
- kind Specifies the type of interpolation to be performed in getting from phinorm to psinorm and psinorm to volnorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

volnorm or (*volnorm*, *time_idxs*)

- **volnorm** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *volnorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single volnorm value for phinorm=0.7, t=0.26s:

```
volnorm_val = Eq_instance.phinorm2volnorm(0.7, 0.26)
```

Find volnorm values at phinorm values of 0.5 and 0.7 at the single time t=0.26s:

```
volnorm_arr = Eq_instance.phinorm2volnorm([0.5, 0.7], 0.26)
```

Find volnorm values at phinorm=0.5 at times t=[0.2s, 0.3s]:

```
volnorm_arr = Eq_instance.phinorm2volnorm(0.5, [0.2, 0.3])
```

Find volnorm values at (phinorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
volnorm_arr = Eq_instance.phinorm2volnorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

phinorm2rmid(*args, **kwargs)

Calculates the mapped outboard midplane major radius corresponding to the passed phinorm (normalized toroidal flux) values.

Parameters

- **phinorm** (*Array-like or scalar float*) Values of the normalized toroidal flux to map to Rmid.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *phinorm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *phinorm*.
- sqrt Set to True to return the square root of Rmid. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *phinorm* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *phinorm* or be a scalar. Default is True (evaluate ALL *phinorm* at EACH element in *t*).
- **rho** Set to True to return r/a (normalized minor radius) instead of Rmid. Default is False (return major radius, Rmid).
- **length_unit** Length unit that *Rmid* is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- **kind** Specifies the type of interpolation to be performed in getting from phinorm to psinorm and psinorm to Rmid. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype rho Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

Rmid or (Rmid, time_idxs)

- **Rmid** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *Rmid*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single Rmid value for phinorm=0.7, t=0.26s:

```
Rmid_val = Eq_instance.phinorm2rmid(0.7, 0.26)
```

Find Rmid values at phinorm values of 0.5 and 0.7 at the single time t=0.26s:

```
Rmid_arr = Eq_instance.phinorm2rmid([0.5, 0.7], 0.26)
```

Find Rmid values at phinorm=0.5 at times t=[0.2s, 0.3s]:

```
Rmid_arr = Eq_instance.phinorm2rmid(0.5, [0.2, 0.3])
```

Find Rmid values at (phinorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
Rmid_arr = Eq_instance.phinorm2rmid([0.6, 0.5], [0.2, 0.3], each_t=False)
```

phinorm2roa (phi_norm, t, **kwargs)

Calculates the normalized minor radius corresponding to the passed phinorm (normalized toroidal flux) values.

Parameters

- **phinorm** (*Array-like or scalar float*) Values of the normalized toroidal flux to map to r/a.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *phinorm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *phinorm*.
- **sqrt** Set to True to return the square root of r/a. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *phinorm* are evaluated at each value in t. If True, t must have only one dimension (or be a scalar). If False, t must match the shape of *phinorm* or be a scalar. Default is True (evaluate ALL *phinorm* at EACH element in t).
- **kind** Specifies the type of interpolation to be performed in getting from phinorm to psinorm and psinorm to Rmid. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype kind String or non-negative int

Kwtype return t Boolean

Returns

roa or (roa, time_idxs)

- roa (*Array or scalar float*) Normalized midplane minor radius. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *roa*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single r/a value for phinorm=0.7, t=0.26s:

```
roa_val = Eq_instance.phinorm2roa(0.7, 0.26)
```

Find r/a values at phinorm values of 0.5 and 0.7 at the single time t=0.26s:

```
roa_arr = Eq_instance.phinorm2roa([0.5, 0.7], 0.26)
```

Find r/a values at phinorm=0.5 at times t=[0.2s, 0.3s]:

```
roa_arr = Eq_instance.phinorm2roa(0.5, [0.2, 0.3])
```

Find r/a values at (phinorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
roa_arr = Eq_instance.phinorm2roa([0.6, 0.5], [0.2, 0.3], each_t=False)
```

phinorm2rho (method, *args, **kwargs)

Convert the passed (phinorm, t) coordinates into one of several coordinates.

Parameters

• method (String) – Indicates which coordinates to convert to. Valid options are:

psinorm	Normalized poloidal flux
volnorm	Normalized volume
Rmid	Midplane major radius
r/a	Normalized minor radius

Additionally, each valid option may be prepended with 'sqrt' to specify the square root of the desired unit.

- **phinorm** (*Array-like or scalar float*) Values of the normalized toroidal flux to map to rho.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *phinorm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *phinorm*.
- sqrt Set to True to return the square root of rho. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- **each_t** When True, the elements in *phinorm* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *phinorm* or be a scalar. Default is True (evaluate ALL *phinorm* at EACH element in *t*).
- **rho** Set to True to return r/a (normalized minor radius) instead of Rmid. Default is False (return major radius, Rmid).
- **length_unit** Length unit that *Rmid* is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from psinorm to Rmid/phinorm/volnorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype rho Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

rho or (rho, time_idxs)

- **rho** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *rho*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Raises ValueError – If *method* is not one of the supported values.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value at phinorm=0.6, t=0.26s:

```
psi_val = Eq_instance.phinorm2rho('psinorm', 0.6, 0.26)
```

Find psinorm values at phinorm of 0.6 and 0.8 at the single time t=0.26s:

```
psi_arr = Eq_instance.phinorm2rho('psinorm', [0.6, 0.8], 0.26)
```

Find psinorm values at phinorm of 0.6 at times t=[0.2s, 0.3s]:

```
psi_arr = Eq_instance.phinorm2rho('psinorm', 0.6, [0.2, 0.3])
```

Find psinorm values at (phinorm, t) points (0.6, 0.2s) and (0.5m, 0.3s):

```
psi_arr = Eq_instance.phinorm2rho('psinorm', [0.6, 0.5], [0.2, 0.3], each_t=False)
```

volnorm2psinorm(*args, **kwargs)

Calculates the normalized poloidal flux corresponding to the passed volnorm (normalized flux surface volume) values.

Parameters

- **volnorm** (*Array-like or scalar float*) Values of the normalized flux surface volume to map to psinorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *volnorm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *volnorm*.
- sqrt Set to True to return the square root of psinorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *volnorm* are evaluated at each value in t. If True, t must have only one dimension (or be a scalar). If False, t must match the shape of *volnorm* or be a scalar. Default is True (evaluate ALL *volnorm* at EACH element in t).
- **kind** Specifies the type of interpolation to be performed in getting from volnorm to psinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

psinorm or (*psinorm*, *time_idxs*)

- **psinorm** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *psinorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value for volnorm=0.7, t=0.26s:

```
psinorm_val = Eq_instance.volnorm2psinorm(0.7, 0.26)
```

Find psinorm values at volnorm values of 0.5 and 0.7 at the single time t=0.26s:

```
psinorm_arr = Eq_instance.volnorm2psinorm([0.5, 0.7], 0.26)
```

Find psinorm values at volnorm=0.5 at times t=[0.2s, 0.3s]:

```
psinorm_arr = Eq_instance.volnorm2psinorm(0.5, [0.2, 0.3])
```

Find psinorm values at (volnorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
psinorm_arr = Eq_instance.volnorm2psinorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

volnorm2phinorm(*args, **kwargs)

Calculates the normalized toroidal flux corresponding to the passed volnorm (normalized flux surface volume) values.

Parameters

- **volnorm** (*Array-like or scalar float*) Values of the normalized flux surface volume to map to phinorm.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *volnorm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *volnorm*.
- sqrt Set to True to return the square root of phinorm. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- each_t When True, the elements in *volnorm* are evaluated at each value in t. If True, t must have only one dimension (or be a scalar). If False, t must match the shape of *volnorm* or be a scalar. Default is True (evaluate ALL *volnorm* at EACH element in t).
- kind Specifies the type of interpolation to be performed in getting from volnorm to psinorm and psinorm to phinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype kind String or non-negative int

Kwtype return t Boolean

Returns

phinorm or (*phinorm*, *time_idxs*)

- **phinorm** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *phinorm*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single phinorm value for volnorm=0.7, t=0.26s:

```
phinorm_val = Eq_instance.volnorm2phinorm(0.7, 0.26)
```

Find phinorm values at volnorm values of 0.5 and 0.7 at the single time t=0.26s:

```
phinorm_arr = Eq_instance.volnorm2phinorm([0.5, 0.7], 0.26)
```

Find phinorm values at volnorm=0.5 at times t=[0.2s, 0.3s]:

```
phinorm_arr = Eq_instance.volnorm2phinorm(0.5, [0.2, 0.3])
```

Find phinorm values at (volnorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
phinorm_arr = Eq_instance.volnorm2phinorm([0.6, 0.5], [0.2, 0.3], each_t=False)
```

```
volnorm2rmid(*args, **kwargs)
```

Calculates the mapped outboard midplane major radius corresponding to the passed volnorm (normalized flux surface volume) values.

Parameters

- **volnorm** (*Array-like or scalar float*) Values of the normalized flux surface volume to map to Rmid.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *volnorm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *volnorm*.
- sqrt Set to True to return the square root of Rmid. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- **each_t** When True, the elements in *volnorm* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *volnorm* or be a scalar. Default is True (evaluate ALL *volnorm* at EACH element in *t*).
- **rho** Set to True to return r/a (normalized minor radius) instead of Rmid. Default is False (return major radius, Rmid).
- **length_unit** Length unit that *Rmid* is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from volnorm to psinorm and psinorm to Rmid. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each t Boolean

Kwtype rho Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

Rmid or (Rmid, time_idxs)

- **Rmid** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *Rmid*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single Rmid value for volnorm=0.7, t=0.26s:

```
Rmid_val = Eq_instance.volnorm2rmid(0.7, 0.26)
```

Find Rmid values at volnorm values of 0.5 and 0.7 at the single time t=0.26s:

```
Rmid_arr = Eq_instance.volnorm2rmid([0.5, 0.7], 0.26)
```

Find Rmid values at volnorm=0.5 at times t=[0.2s, 0.3s]:

```
Rmid_arr = Eq_instance.volnorm2rmid(0.5, [0.2, 0.3])
```

Find Rmid values at (volnorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
Rmid_arr = Eq_instance.volnorm2rmid([0.6, 0.5], [0.2, 0.3], each_t=False)
```

volnorm2roa(*args, **kwargs)

Calculates the normalized minor radius corresponding to the passed volnorm (normalized flux surface volume) values.

Parameters

- **volnorm** (*Array-like or scalar float*) Values of the normalized flux surface volume to map to r/a.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *volnorm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *volnorm*.
- sqrt Set to True to return the square root of r/a. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- **each_t** When True, the elements in *volnorm* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *volnorm* or be a scalar. Default is True (evaluate ALL *volnorm* at EACH element in *t*).
- **kind** Specifies the type of interpolation to be performed in getting from volnorm to psinorm and psinorm to Rmid. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype kind String or non-negative int

Kwtype return t Boolean

Returns

roa or (roa, time_idxs)

- roa (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *roa*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single r/a value for volnorm=0.7, t=0.26s:

```
roa_val = Eq_instance.volnorm2roa(0.7, 0.26)
```

Find r/a values at volnorm values of 0.5 and 0.7 at the single time t=0.26s:

```
roa_arr = Eq_instance.volnorm2roa([0.5, 0.7], 0.26)
```

Find r/a values at volnorm=0.5 at times t=[0.2s, 0.3s]:

```
roa_arr = Eq_instance.volnorm2roa(0.5, [0.2, 0.3])
```

Find r/a values at (volnorm, t) points (0.6, 0.2s) and (0.5, 0.3s):

```
roa_arr = Eq_instance.volnorm2roa([0.6, 0.5], [0.2, 0.3], each_t=False)
```

volnorm2rho (method, *args, **kwargs)

Convert the passed (volnorm, t) coordinates into one of several coordinates.

Parameters

• method (String) – Indicates which coordinates to convert to. Valid options are:

psinorm	Normalized poloidal flux
phinorm	Normalized toroidal flux
Rmid	Midplane major radius
r/a	Normalized minor radius

Additionally, each valid option may be prepended with 'sqrt' to specify the square root of the desired unit.

- **volnorm** (*Array-like or scalar float*) Values of the normalized flux surface volume to map to rho.
- **t** (*Array-like or scalar float*) Times to perform the conversion at. If *t* is a single value, it is used for all of the elements of *volnorm*. If the *each_t* keyword is True, then *t* must be scalar or have exactly one dimension. If the *each_t* keyword is False, *t* must have the same shape as *volnorm*.
- sqrt Set to True to return the square root of rho. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False.
- **each_t** When True, the elements in *volnorm* are evaluated at each value in *t*. If True, *t* must have only one dimension (or be a scalar). If False, *t* must match the shape of *volnorm* or be a scalar. Default is True (evaluate ALL *volnorm* at EACH element in *t*).
- **rho** Set to True to return r/a (normalized minor radius) instead of Rmid. Default is False (return major radius, Rmid).
- **length_unit** Length unit that *Rmid* is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (use meters).

- kind Specifies the type of interpolation to be performed in getting from volnorm to Rmid/phinorm/psinorm. This is passed to scipy.interpolate.interpld. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interpld for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **return_t** Set to True to return a tuple of (*rho*, *time_idxs*), where *time_idxs* is the array of time indices actually used in evaluating *rho* with nearest-neighbor interpolation. (This is mostly present as an internal helper.) Default is False (only return *rho*).

Kwtype sqrt Boolean

Kwtype each_t Boolean

Kwtype rho Boolean

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Kwtype return_t Boolean

Returns

rho or (*rho*, *time_idxs*)

- **rho** (*Array or scalar float*) The converted coordinates. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array is returned.
- **time_idxs** (Array with same shape as *rho*) The indices (in self.getTimeBase()) that were used for nearest-neighbor interpolation. Only returned if *return_t* is True.

Raises ValueError – If *method* is not one of the supported values.

Examples

All assume that $Eq_instance$ is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value at volnorm=0.6, t=0.26s:

```
psi_val = Eq_instance.volnorm2rho('psinorm', 0.6, 0.26)
```

Find psinorm values at volnorm of 0.6 and 0.8 at the single time t=0.26s:

```
psi_arr = Eq_instance.volnorm2rho('psinorm', [0.6, 0.8], 0.26)
Find psinorm values at volnorm of 0.6 at times t=[0.2s, 0.3s]:
```

```
psi_arr = Eq_instance.volnorm2rho('psinorm', 0.6, [0.2, 0.3])
```

Find psinorm values at (volnorm, t) points (0.6, 0.2s) and (0.5m, 0.3s):

```
psi_arr = Eq_instance.volnorm2rho('psinorm', [0.6, 0.5], [0.2, 0.3], each_t=False)
```

getMagRSpline (length_unit=1, kind='nearest')

Gets the univariate spline to interpolate R_mag as a function of time.

Only used if the instance was created with keyword tspline=True.

Parameters

• length_unit - Length unit that R_mag is returned in. If a string is given, it must be a valid unit specifier:

'm' meters 'cm' centimeters 'mm' millimeters 'in' inches 'ft' feet 'yd' yards 'smoot' smoots 'cubit' cubits 'hand' hands 'default' meters

If length unit is 1 or None, meters are assumed. The default value is 1 (R out returned in meters).

• kind – Specifies the type of interpolation to be performed in getting from t to R_mag. This is passed to scipy interpolate interp1d. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interp1d for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.

Kwtype length unit String or 1

Kwtype kind String or non-negative int

Returns scipy.interpolate.interp1d to convert from t to R_mid.

```
getMagZSpline (length_unit=1, kind='nearest')
```

Gets the univariate spline to interpolate Z mag as a function of time.

Generated for completeness of the core position calculation when using tspline = True

Parameters

• length_unit - Length unit that R_mag is returned in. If a string is given, it must be a valid unit specifier:

'm' meters 'cm' centimeters 'mm' millimeters 'in' inches 'ft' feet 'yd' yards 'smoot' smoots 'cubit' cubits 'hand' hands 'default' meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (R_out returned in meters).

• kind – Specifies the type of interpolation to be performed in getting from t to R mag. This is passed to scipy interpolate interp1d. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interp1d for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.

Kwtype length unit String or 1

Kwtype kind String or non-negative int

Returns scipy.interpolate.interp1d to convert from t to R mid.

```
getRmidOutSpline (length_unit=1, kind='nearest')
```

Gets the univariate spline to interpolate R_mid_out as a function of time.

Generated for completeness of the core position calculation when using tspline = True

Parameters

• length_unit – Length unit that R_mag is returned in. If a string is given, it must be a valid unit specifier:

'm' meters 'cm' centimeters 'mm' millimeters 'in' inches 'ft' feet 'yd' yards 'smoot' smoots 'cubit' cubits 'hand' hands 'default' meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (R_out returned in meters).

• **kind** – Specifies the type of interpolation to be performed in getting from t to R_mag. This is passed to scipy.interpolate.interp1d. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interp1d for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.

Kwtype length_unit String or 1

Kwtype kind String or non-negative int

Returns scipy.interpolate.interp1d to convert from t to R_mid.

```
getAOutSpline (length_unit=1, kind='nearest')
```

Gets the univariate spline to interpolate a_out as a function of time.

Parameters

• **length_unit** – Length unit that a_out is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If *length_unit* is 1 or None, meters are assumed. The default value is 1 (a_out returned in meters).

• **kind** – Specifies the type of interpolation to be performed in getting from t to a_out. This is passed to scipy.interpolate.interp1d. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interp1d for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.

Kwtype length unit String or 1

Kwtype kind String or non-negative int

Returns scipy.interpolate.interp1d to convert from t to a_out.

getInfo()

Abstract method. See child classes for implementation.

Returns namedtuple of instance parameters (shot, equilibrium type, size, timebase, etc.)

getTimeBase()

Abstract method. See child classes for implementation.

Returns timebase array [t]

getFluxGrid()

Abstract method. See child classes for implementation.

returns 3D grid of psi(r,z,t)

The array returned should have the following dimensions: First dimension: time Second dimension: Z Third dimension: R

getRGrid()

Abstract method. See child classes for implementation.

Returns vector of R-values for psiRZ grid [r]

getZGrid()

Abstract method. See child classes for implementation.

Returns vector of Z-values for psiRZ grid [z]

getFluxAxis()

Abstract method. See child classes for implementation.

Returns psi at magnetic axis [t]

getFluxLCFS()

Abstract method. See child classes for implementation.

Returns psi a separatrix [t]

getRLCFS()

Abstract method. See child classes for implementation.

Returns R-positions (n points) mapping LCFS [t,n]

getZLCFS()

Abstract method. See child classes for implementation.

Returns Z-positions (n points) mapping LCFS [t,n]

remapLCFS()

Abstract method. See child classes for implementation.

Overwrites stored R,Z positions of LCFS with explicitly calculated psinorm=1 surface. This surface is then masked using core.inPolygon() to only draw within vacuum vessel, the end result replacing RLCFS, ZLCFS with an R,Z array showing the divertor legs of the flux surface in addition to the core-enclosing closed flux surface.

getFluxVol()

Abstract method. See child classes for implementation.

Returns volume contained within flux surface as function of psi [psi,t]. Psi assumed to be evenly-spaced grid on [0,1]

getVolLCFS()

Abstract method. See child classes for implementation.

Returns plasma volume within LCFS [t]

getRmidPsi()

Abstract method. See child classes for implementation.

Returns outboard-midplane major radius of flux surface [t,psi]

getF()

Abstract method. See child classes for implementation.

Returns F=RB_{Phi}(Psi), often calculated for grad-shafranov solutions [psi,t]

getFluxPres()

Abstract method. See child classes for implementation.

Returns calculated pressure profile [psi,t]. Psi assumed to be evenly-spaced grid on [0,1]

getFFPrime()

Abstract method. See child classes for implementation.

Returns FF' function used for grad-shafranov solutions [psi,t]

getPPrime()

Abstract method. See child classes for implementation.

Returns plasma pressure gradient as a function of psi [psi,t]

getElongation()

Abstract method. See child classes for implementation.

Returns LCFS elongation [t]

getUpperTriangularity()

Abstract method. See child classes for implementation.

Returns LCFS upper triangularity [t]

getLowerTriangularity()

Abstract method. See child classes for implementation.

Returns LCFS lower triangularity [t]

getShaping()

Abstract method. See child classes for implementation.

Returns dimensionless shaping parameters for plasma. Namedtuple containing {LCFS elongation, LCFS upper/lower triangularity}

getMagR()

Abstract method. See child classes for implementation.

Returns magnetic-axis major radius [t]

getMagZ()

Abstract method. See child classes for implementation.

Returns magnetic-axis Z [t]

getAreaLCFS()

Abstract method. See child classes for implementation.

Returns LCFS surface area [t]

getAOut()

Abstract method. See child classes for implementation.

Returns outboard-midplane minor radius [t]

getRmidOut()

Abstract method. See child classes for implementation.

Returns outboard-midplane major radius [t]

getGeometry()

Abstract method. See child classes for implementation.

Returns dimensional geometry parameters Namedtuple containing {mag axis R,Z, LCFS area, volume, outboard-midplane major radius}

getQProfile()

Abstract method. See child classes for implementation.

Returns safety factor q profile [psi,t] Psi assumed to be evenly-spaced grid on [0,1]

getQ0()

Abstract method. See child classes for implementation.

Returns q on magnetic axis [t]

getQ95()

Abstract method. See child classes for implementation.

Returns q on 95% flux surface [t]

getQLCFS()

Abstract method. See child classes for implementation.

Returns q on LCFS [t]

getQ1Surf()

Abstract method. See child classes for implementation.

Returns outboard-midplane minor radius of q=1 surface [t]

getQ2Surf()

Abstract method. See child classes for implementation.

Returns outboard-midplane minor radius of q=2 surface [t]

getQ3Surf()

Abstract method. See child classes for implementation.

Returns outboard-midplane minor radius of q=3 surface [t]

getQs()

Abstract method. See child classes for implementation.

Returns specific q-profile values. Namedtuple containing {q0, q95, qLCFS, minor radius of q=1,2,3 surfaces}

getBtVac()

Abstract method. See child classes for implementation.

Returns vacuum on-axis toroidal field [t]

getBtPla()

Abstract method. See child classes for implementation.

Returns plasma on-axis toroidal field [t]

getBpAvg()

Abstract method. See child classes for implementation.

Returns average poloidal field [t]

getFields()

Abstract method. See child classes for implementation.

Returns magnetic-field values. Namedtuple containing {Btor on magnetic axis (plasma and vacuum), avg Bpol}

getIpCalc()

Abstract method. See child classes for implementation.

Returns calculated plasma current [t]

getIpMeas()

Abstract method. See child classes for implementation.

Returns measured plasma current [t]

getJp()

Abstract method. See child classes for implementation.

Returns grid of calculated toroidal current density [t,z,r]

getBetaT()

Abstract method. See child classes for implementation.

Returns calculated global toroidal beta [t]

getBetaP()

Abstract method. See child classes for implementation.

Returns calculated global poloidal beta [t]

getLi()

Abstract method. See child classes for implementation.

Returns calculated internal inductance of plasma [t]

getBetas()

Abstract method. See child classes for implementation.

Returns calculated betas and inductance. Namedtuple of {betat,betap,Li}

getDiamagFlux()

Abstract method. See child classes for implementation.

Returns diamagnetic flux [t]

getDiamagBetaT()

Abstract method. See child classes for implementation.

Returns diamagnetic-loop toroidal beta [t]

getDiamagBetaP()

Abstract method. See child classes for implementation.

Returns diamagnetic-loop poloidal beta [t]

getDiamagTauE()

Abstract method. See child classes for implementation.

Returns diamagnetic-loop energy confinement time [t]

getDiamagWp()

Abstract method. See child classes for implementation.

Returns diamagnetic-loop plasma stored energy [t]

getDiamag()

Abstract method. See child classes for implementation.

Returns diamagnetic measurements of plasma parameters. Namedtuple of {diamag. flux, betat, betap from coils, tau_E from diamag., diamag. stored energy}

getWMHD()

Abstract method. See child classes for implementation.

Returns calculated MHD stored energy [t]

getTauMHD()

Abstract method. See child classes for implementation.

Returns calculated MHD energy confinement time [t]

getPinj()

Abstract method. See child classes for implementation.

Returns calculated injected power [t]

getCurrentSign()

Abstract method. See child classes for implementation.

Returns calculated current direction, where CCW = +

getWbdot()

Abstract method. See child classes for implementation.

Returns calculated d/dt of magnetic stored energy [t]

getWpdot()

Abstract method. See child classes for implementation.

Returns calculated d/dt of plasma stored energy [t]

getEnergy()

Abstract method. See child classes for implementation.

Returns stored-energy parameters. Namedtuple of {stored energy, confinement time, injected power, d/dt of magnetic, plasma stored energy}

getParam(path)

Abstract method. See child classes for implementation.

Backup function: takes parameter name for variable, returns variable directly. Acts as wrapper to direct data-access routines from within object.

getMachineCrossSection()

Abstract method. See child classes for implementation.

Returns (R,Z) coordinates of vacuum wall cross-section for plotting/masking routines.

getMachineCrossSectionFull()

Abstract method. See child classes for implementation.

Returns (R,Z) coordinates of machine wall cross-section for plotting routines. Returns a more detailed cross-section than getLimiter(), generally a vector map displaying non-critical cross-section information. If this is unavailable, this should point to self.getMachineCrossSection(), which pulls the limiter outline stored by default in data files e.g. g-eqdsk files.

```
gfile (time=None, nw=None, nh=None, shot=None, name=None, tunit='ms', title='EQTOOLS', nbbbs=100)
```

```
plotFlux (fill=True, mask=True)
```

Plots flux contours directly from psi grid.

Parameters fill – Set True to plot filled contours. Set False (default) to plot white-background color contours.

Kwtype fill Boolean

eqtools.eqdskreader module

This module contains the EqdskReader class, which creates Equilibrium class functionality for equilibria stored in eqdsk files from EFIT(a- and g-files).

Classes:

EqdskReader: class inheriting Equilibrium reading g- and a-files for equilibrium data.

Equilibrium subclass working from eqdsk ASCII-file equilibria.

Inherits mapping and structural data from Equilibrium, populates equilibrium and profile data from g- and a-files for a selected shot and time window.

Create instance of EqdskReader.

Generates object and reads data from selected g-file (either manually set or autodetected based on user shot and time selection), storing as object attributes for usage in Equilibrium mapping methods.

Calling structure - user may call class with shot and time (ms) values, set by keywords (or positional placement allows calling without explicit keyword syntax). EqdskReader then attempts to construct filenames from the shot/time, of the form 'g[shot].[time]' and 'a[shot].[time]'. Alternately, the user may skip this input and explicitly set paths to the g- and/or a-files, using the gfile and afile keyword arguments. If both types of calls are set, the explicit g-file and a-file paths override the auto-generated filenames from the shot and time.

Parameters

- **shot** Int. Shot index.
- time Int. Time index (typically ms). Shot and Time used to autogenerate filenames.
- gfile String. Manually selects ASCII file for equilibrium read.
- afile String. Manually selects ASCII file for time-history read.
- **length_unit** String. Flag setting length unit for equilibrium scales. Defaults to 'm' for lengths in meters.
- **verbose** Boolean. When set to False, suppresses terminal outputs during CSV read. Defaults to True (prints terminal output).

Raises

- IOError if both name/shot and explicit filenames are not set.
- ValueError if the g-file cannot be found, or if multiple valid g/a-files are found.

getInfo()

returns namedtuple of equilibrium information

Returns

namedtuple containing

shot	shot index
time	time point of g-file
nr	size of R-axis of spatial grid
nz	size of Z-axis of spatial grid
efittype	EFIT calculation type (magnetic, kinetic, MSE)

readAFile (afile)

Reads a-file (scalar time-history data) to pull additional equilibrium data not found in g-file, populates remaining data (initialized as None) in object.

Parameters afile – String. Path to ASCII a-file.

Raises IOError – If afile is not found.

rz2psi(R, Z, *args, **kwargs)

Converts passed, R,Z arrays to psi values.

Wrapper for Equilibrium.rz2psi masking out timebase dependence.

Parameters

- **R** Array-like or scalar float. Values of the radial coordinate to map to poloidal flux. If the make_grid keyword is True, R must have shape (len_R,).
- **Z** Array-like or scalar float. Values of the vertical coordinate to map to poloidal flux. Must have the same shape as R unless the make_grid keyword is set. If the make_grid keyword is True, Z must have shape (len_Z,).
- *args Slot for time input for consistent syntax with Equilibrium.rz2psi. will return dummy value for time if input in EqdskReader.
- make_grid Boolean. Set to True to pass R and Z through meshgrid before evaluating. If this is set to True, R and Z must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **length_unit** String or 1. Length unit that R and Z are being given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (R and Z given in meters).

• **kwargs – Other keywords (i.e., return_t) to rz2psi are valid (necessary for proper inheritance and usage in other mapping routines) but will return dummy values.

Returns

Array or scalar float. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array instance is returned. If R and Z both have the same shape then

psi has this shape as well. If the make_grid keyword was True then psi has shape (len(Z), len(R)).

Return type psi

rz2psinorm(R, Z, *args, **kwargs)

Calculates the normalized poloidal flux at the given (R,Z). Wrapper for Equilibrium.rz2psinorm masking out timebase dependence.

Uses the definition: $psi_norm = (psi - psi(0)) / (psi(a) - psi(0))$

Parameters

- **R** Array-like or scalar float. Values of the radial coordinate to map to normalized poloidal flux. If R and Z are both scalar values, they are used as the coordinate pair for all of the values in t. Must have the same shape as Z unless the make_grid keyword is set. If the make_grid keyword is True, R must have shape (len_R,).
- **Z** Array-like or scalar float. Values of the vertical coordinate to map to normalized poloidal flux. If R and Z are both scalar values, they are used as the coordinate pair for all of the values in t. Must have the same shape as R unless the make_grid keyword is set. If the make_grid keyword is True, Z must have shape (len_Z,).
- *args Slot for time input for consistent syntax with Equilibrium.rz2psinorm. will return dummy value for time if input in EqdskReader.
- sqrt Boolean. Set to True to return the square root of normalized flux. Only the square root of positive psi_norm values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False (return psinorm).
- make_grid Boolean. Set to True to pass R and Z through meshgrid before evaluating. If this is set to True, R and Z must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **length_unit** String or 1. Length unit that R and Z are being given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (R and Z given in meters).

 **kwargs – Other keywords passed to Equilibrium.rz2psinorm are valid, but will return dummy values (i.e. for timebase keywords)

Returns

Array or scalar float. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array instance is returned. If R and Z both have the same shape then psinorm has this shape as well. If the make_grid keyword was True then psinorm has shape (len(Z), len(R)).

Return type psinorm

Examples

All assume that Eq_instance is a valid instance EqdskReader:

Find single psinorm value at R=0.6m, Z=0.0m:

```
psi_val = Eq_instance.rz2psinorm(0.6, 0)
```

Find psinorm values at (R, Z) points (0.6m, 0m) and (0.8m, 0m). Note that the Z vector must be fully specified, even if the values are all the same:

```
psi_arr = Eq_instance.rz2psinorm([0.6, 0.8], [0, 0])
```

Find psinorm values on grid defined by 1D vector of radial positions R and 1D vector of vertical positions Z:

```
psi_mat = Eq_instance.rz2psinorm(R, Z, make_grid=True)
```

```
rz2phinorm(R, Z, *args, **kwargs)
```

Calculates normalized toroidal flux at a given (R,Z).

Wrapper for Equilibrium.rz2phinorm masking out timebase dependence.

Parameters

- **R** Array-like or scalar float. Values of the radial coordinate to map to normalized toroidal flux. Must have the same shape as Z unless the make_grid keyword is set. If the make_grid keyword is True, R must have shape (len_R₁).
- **Z** Array-like or scalar float. Values of the vertical coordinate to map to normalized toroidal flux. Must have the same shape as R unless the make_grid keyword is set. If the make_grid keyword is True, Z must have shape (len_Z,).
- *args Slot for time input for consistent syntax with Equilibrium.rz2phinorm. will return dummy value for time if input in EqdskReader.
- **sqrt** Boolean. Set to True to return the square root of normalized flux. Only the square root of positive phi_norm values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False (return phinorm).
- make_grid Boolean. Set to True to pass R and Z through meshgrid before evaluating. If this is set to True, R and Z must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **kind** String or non-negative int. Specifies the type of interpolation to be performed in getting from psinorm to phinorm. This is passed to scipy.interpolate.interp1d. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interp1d for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **length_unit** String or 1. Length unit that R and Z are being given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (R and Z given in meters).

• **kwargs – Other keywords passed to Equilibrium.rz2phinorm are valid, but will return dummy values (i.e. for timebase keywords)

Returns

Array or scalar float. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array instance is returned. If R and Z both have the same shape then phinorm has this shape as well. If the make_grid keyword was True then phinorm has shape (len(Z), len(R)).

Return type phinorm

Examples

All assume that Eq_instance is a valid instance of EqdskReader.

Find single phinorm value at R=0.6m, Z=0.0m:

```
phi_val = Eq_instance.rz2phinorm(0.6, 0)
```

Find phinorm values at (R, Z) points (0.6m, 0m) and (0.8m, 0m). Note that the Z vector must be fully specified, even if the values are all the same:

```
phi_arr = Eq_instance.rz2phinorm([0.6, 0.8], [0, 0])
```

Find phinorm values on grid defined by 1D vector of radial positions R and 1D vector of vertical positions Z:

```
phi_mat = Eq_instance.rz2phinorm(R, Z, make_grid=True)
```

```
rz2volnorm(*args, **kwargs)
```

Calculates the normalized flux surface volume.

Not implemented for EqdskReader, as necessary parameter is not read from a/g-files.

```
Raises NotImplementedError - in all cases.
```

```
rz2rho (method, R, Z, t=False, sqrt=False, make_grid=False, kind='cubic', length_unit=1)
```

Convert the passed (R, Z) coordinates into one of several normalized coordinates. Wrapper for Equilibrium.rz2rho masking timebase dependence.

Parameters

• method – String. Indicates which normalized coordinates to use. Valid options are:

psinorm	Normalized poloidal flux
phinorm	Normalized toroidal flux
volnorm	Normalized volume

- **R** Array-like or scalar float. Values of the radial coordinate to map to normalized coordinate. Must have the same shape as Z unless the make_grid keyword is set. If the make_grid keyword is True, R must have shape (len_R,).
- **Z** Array-like or scalar float. Values of the vertical coordinate to map to normalized coordinate. Must have the same shape as R unless the make_grid keyword is set. If the make_grid keyword is True, Z must have shape (len_Z,).
- t indeterminant. Provides duck typing for inclusion of t values. Passed t values either as an Arg or Kwarg are neglected.
- sqrt Boolean. Set to True to return the square root of normalized coordinate. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False (return normalized coordinate itself).
- make_grid Boolean. Set to True to pass R and Z through meshgrid before evaluating. If this is set to True, R and Z must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **kind** String or non-negative int. Specifies the type of interpolation to be performed in getting from psinorm to phinorm or volnorm. This is passed to scipy.interpolate.interp1d. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interp1d for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- **length_unit** String or 1. Length unit that R and Z are being given in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (R and Z given in meters).

Kwtype kind phinorm and volnorm only

Returns

Array or scalar float. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array instance is returned. If R and Z both have the same shape then rho has this shape as well. If the make_grid keyword was True then rho has shape (len(Z), len(R)).

Return type rho

Raises ValueError – If method is not one of the supported values.

Examples

All assume that Eq_instance is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single psinorm value at R=0.6m, Z=0.0m:

```
psi_val = Eq_instance.rz2rho('psinorm', 0.6, 0)
```

Find psinorm values at (R, Z) points (0.6m, 0m) and (0.8m, 0m). Note that the Z vector must be fully specified, even if the values are all the same:

```
psi_arr = Eq_instance.rz2rho('psinorm', [0.6, 0.8], [0, 0])
```

Find psinorm values on grid defined by 1D vector of radial positions R and 1D vector of vertical positions Z:

```
psi_mat = Eq_instance.rz2rho('psinorm', R, Z, make_grid=True)
```

rz2rmid (R, Z, t=False, sqrt=False, make_grid=False, rho=False, kind='cubic', length_unit=1)

Maps the given points to the outboard midplane major radius, R_mid. Wrapper for Equilibrium.rz2rmid masking timebase dependence.

Based on the IDL version efit_rz2rmid.pro by Steve Wolfe.

Parameters

- **R** Array-like or scalar float. Values of the radial coordinate to map to midplane radius. Must have the same shape as Z unless the make_grid keyword is set. If the make_grid keyword is True, R must have shape (len_R₁).
- **Z** Array-like or scalar float. Values of the vertical coordinate to map to midplane radius. Must have the same shape as R unless the make_grid keyword is set. If the make_grid keyword is True, Z must have shape (len_Z,).
- t indeterminant. Provides duck typing for inclusion of t values. Passed t values either as an Arg or Kwarg are neglected.
- sqrt Boolean. Set to True to return the square root of midplane radius. Only the square root of positive values is taken. Negative values are replaced with zeros, consistent with Steve Wolfe's IDL implementation efit_rz2rho.pro. Default is False (return R_mid itself).
- make_grid Boolean. Set to True to pass R and Z through meshgrid before evaluating. If this is set to True, R and Z must each only have a single dimension, but can have different lengths. Default is False (do not form meshgrid).
- **rho** Boolean. Set to True to return r/a (normalized minor radius) instead of R_mid. Default is False (return major radius, R_mid).
- **kind** String or non-negative int. Specifies the type of interpolation to be performed in getting from psinorm to R_mid. This is passed to scipy.interpolate.interp1d. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer, it specifies the order of spline to use. See the documentation for interp1d for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.
- length_unit -

String or 1. Length unit that R and Z are being given in AND that R_mid is returned in. If a string is given, it must be a valid unit specifier:

'm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
'yd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (R and Z given in meters, R_mid returned in meters).

Returns

Array or scalar float. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array instance is returned. If R and Z both have the same shape then R_mid has this shape as well. If the make_grid keyword was True then R_mid has shape (len(Z), len(R)).

Return type R_mid

Examples

All assume that Eq_instance is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single R mid value at R=0.6m, Z=0.0m:

```
R_mid_val = Eq_instance.rz2rmid(0.6, 0)
```

Find R_mid values at (R, Z) points (0.6m, 0m) and (0.8m, 0m). Note that the Z vector must be fully specified, even if the values are all the same:

```
R_{mid\_arr} = Eg_{instance.rz2rmid([0.6, 0.8], [0, 0])}
```

Find R_mid values on grid defined by 1D vector of radial positions R and 1D vector of vertical positions Z:

```
R_mid_mat = Eq_instance.rz2rmid(R, Z, make_grid=True)
```

psinorm2rmid (psi_norm, t=False, rho=False, kind='cubic', length_unit=1)

Calculates the outboard R_mid location corresponding to the passed psi_norm (normalized poloidal flux) values.

Parameters

- **psi_norm** Array-like or scalar float. Values of the normalized poloidal flux to map to midplane radius.
- t indeterminant. Provides duck typing for inclusion of t values. Passed t values either as an Arg or Kwarg are neglected.
- **rho** Boolean. Set to True to return r/a (normalized minor radius) instead of R_mid. Default is False (return major radius, R_mid).
- **kind** String or non-negative int. Specifies the type of interpolation to be performed in getting from psinorm to R_mid. This is passed to scipy.interpolate.interp1d. Valid options are: 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' If this keyword is an integer,

it specifies the order of spline to use. See the documentation for interp1d for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.

• **length_unit** – String or 1. Length unit that R_mid is returned in. If a string is given, it must be a valid unit specifier:

ʻm'	meters
'cm'	centimeters
'mm'	millimeters
'in'	inches
'ft'	feet
ʻyd'	yards
'smoot'	smoots
'cubit'	cubits
'hand'	hands
'default'	meters

If length_unit is 1 or None, meters are assumed. The default value is 1 (R_mid returned in meters).

Returns

Array or scalar float. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array instance is returned.

Return type R_mid

Examples

All assume that Eq_instance is a valid instance of the appropriate extension of the Equilibrium abstract class.

Find single R_mid value for psinorm=0.7:

```
R_mid_val = Eq_instance.psinorm2rmid(0.7)
```

Find R_mid values at psi_norm values of 0.5 and 0.7. Note that the Z vector must be fully specified, even if the values are all the same:

```
R_mid_arr = Eq_instance.psinorm2rmid([0.5, 0.7])
```

psinorm2volnorm(*args, **kwargs)

Calculates the outboard R_mid location corresponding to psi_norm (normalized poloidal flux) values. Not implemented for EqdskReader, as necessary parameter is not read from a/g-files.

psinorm2phinorm (psi_norm, t=False, kind='cubic')

Calculates the normalized toroidal flux corresponding to the passed psi_norm (normalized poloidal flux) values.

Parameters

- **psi_norm** Array-like or scalar float. Values of the normalized poloidal flux to map to normalized toroidal flux.
- t indeterminant. Provides duck typing for inclusion of t values. Passed t values either as an Arg or Kwarg are neglected.

• **kind** – String or non-negative int. Specifies the type of interpolation to be performed in getting from psinorm to phinorm. This is 'linear', 'nearest', 'zero', 'slinear', 'quadratic', 'cubic' passed to scipy.interpolate.interp1d. Valid options are: If this keyword is an integer, it specifies the order of spline to use. See the documentation for interp1d for more details. Default value is 'cubic' (3rd order spline interpolation). On some builds of scipy, this can cause problems, in which case you should try 'linear' until you can rebuild your scipy install.

Returns

Array or scalar float. If all of the input arguments are scalar, then a scalar is returned. Otherwise, a scipy Array instance is returned.

Return type phinorm

Examples

All assume that Eq_instance is a valid instance of the appropriate extension of the Equilibrium abstract class.

```
Find single phinorm value for psinorm=0.7:
```

```
phinorm_val = Eq_instance.psinorm2phinorm(0.7)
```

Find phinorm values at psi_norm values of 0.5 and 0.7. Note that the Z vector must be fully specified, even if the values are all the same:

```
phinorm_arr = Eq_instance.psinorm2phinorm([0.5, 0.7])
```

getTimeBase()

Returns EFIT time point

getCurrentSign()

Returns the sign of the current, based on the check in Steve Wolfe's IDL implementation efit_rz2psi.pro.

getFluxGrid()

Returns EFIT flux grid, [r,z]

getRGrid (length_unit=1)

Returns EFIT R-axis [r]

getZGrid (length_unit=1)

Returns EFIT Z-axis [z]

getFluxAxis()

Returns psi on magnetic axis

getFluxLCFS()

Returns psi at separatrix

getRLCFS (length_unit=1)

Returns array of R-values of LCFS

getZLCFS (length_unit=1)

Returns array of Z-values of LCFS

remapLCFS (mask=False)

Overwrites RLCFS, ZLCFS values pulled from EFIT with explicitly-calculated contour of psinorm=1 surface.

```
(using inPolygon). Set False to draw full contour of psi = psiLCFS.
getFluxVol()
getVolLCFS (length_unit=3)
     Returns volume with LCFS.
         Raises ValueError – if a-file data is not read.
getRmidPsi()
     Returns outboard-midplane major radius of flux surfaces.
     Data not read from a/g-files, not implemented for EqdskReader.
         Raises NotImplementedError – RmidPsi not read from a/g-files.
getF()
     returns F=RB_{Phi}(Psi), often calculated for grad-shafranov solutions [psi,t]
getFluxPres()
     Returns pressure on flux surface p(psi)
getFFPrime()
     returns FF' function used for grad-shafranov solutions [psi,t]
getPPrime()
     returns plasma pressure gradient as a function of psi [psi,t]
getElongation()
     Returns elongation of LCFS.
         Raises ValueError – if a-file data is not read.
getUpperTriangularity()
     Returns upper triangularity of LCFS.
         Raises ValueError – if a-file data is not read.
getLowerTriangularity()
     Returns lower triangularity of LCFS.
         Raises ValueError – if a-file data is not read.
getShaping()
     Pulls LCFS elongation, upper/lower triangularity.
         Returns namedtuple containing [kappa,delta_u,delta_l].
         Raises ValueError – if a-file data is not read.
getMagR (length_unit=1)
     Returns major radius of magnetic axis.
         Raises ValueError – if a-file data is not read.
getMagZ (length_unit=1)
     Returns Z of magnetic axis.
         Raises ValueError – if a-file data is not read.
getAreaLCFS (length_unit=2)
     Returns surface area of LCFS.
         Raises ValueError – if a-file data is not read.
```

Parameters mask – Boolean. Default False. Set True to mask LCFS path to limiter outline

```
getAOut (length_unit=1)
     Returns outboard-midplane minor radius of LCFS.
         Raises ValueError – if a-file data is not read.
getRmidOut (length_unit=1)
     Returns outboard-midplane major radius of LCFS.
         Raises ValueError – if a-file data is not read.
getGeometry (length_unit=None)
     Pulls dimensional geometry parameters.
         Returns namedtuple containing [Rmag,Zmag,AreaLCFS,aOut,RmidOut]
         Parameters length unit - TODO
         Raises ValueError – if a-file data is not read.
getQProfile()
     Returns safety factor q(psi).
     Returns safety factor q on-axis, q0.
         Raises ValueError – if a-file data is not read.
getQ95()
     Returns safety factor q at 95% flux surface.
         Raises ValueError – if a-file data is not read.
getQLCFS()
     Returns safety factor q at LCFS (interpolated).
         Raises ValueError – if a-file data is not loaded.
getQ1Surf (length_unit=1)
     Returns outboard-midplane minor radius of q=1 surface.
         Raises ValueError – if a-file data is not read.
getQ2Surf (length_unit=1)
     Returns outboard-midplane minor radius of q=2 surface.
         Raises ValueError – if a-file data is not read.
getQ3Surf (length_unit=1)
     Returns outboard-midplane minor radius of q=3 surface.
         Raises ValueError – if a-file data is not read.
getQs (length unit=1)
     Pulls q-profile data.
         Returns namedtuple containing [q0,q95,qLCFS,rq1,rq2,rq3]
         Raises ValueError – if a-file data is not read.
getBtVac()
     Returns vacuum toroidal field on-axis.
         Raises ValueError – if a-file data is not read.
getBtPla()
     Returns plasma toroidal field on-axis.
```

```
Raises ValueError – if a-file data is not read.
getBpAvg()
     Returns average poloidal field.
         Raises ValueError – if a-file data is not read.
getFields()
     Pulls vacuum and plasma toroidal field, poloidal field data.
         Returns namedtuple containing [BtVac,BtPla,BpAvg]
         Raises ValueError – if a-file data is not read.
getIpCalc()
     Returns EFIT-calculated plasma current.
getIpMeas()
     Returns measured plasma current.
         Raises ValueError – if a-file data is not read.
qetJp()
     Returns (r,z) grid of toroidal plasma current density.
     Data not read from g-file, not implemented for EqdskReader.
         Raises NotImplementedError - Jp not read from g-file.
getBetaT()
     Returns EFIT-calculated toroidal beta.
         Raises ValueError – if a-file data is not read.
getBetaP()
     Returns EFIT-calculated poloidal beta.
         Raises ValueError – if a-file data is not read
getLi()
     Returns internal inductance of plasma.
         Raises ValueError – if a-file data is not read.
getBetas()
     Pulls EFIT-calculated betas and internal inductance.
         Returns namedtuple containing [betat,betap,Li]
         Raises ValueError – if a-file data is not read.
getDiamagFlux()
     Returns diamagnetic flux.
         Raises ValueError – if a-file data is not read.
getDiamagBetaT()
     Returns diamagnetic-loop measured toroidal beta.
         Raises ValueError – if a-file data is not read.
getDiamagBetaP()
     Returns diamagnetic-loop measured poloidal beta.
         Raises ValueError – if a-file data is not read.
```

getDiamagTauE()

Returns diamagnetic-loop energy confinement time.

Raises ValueError – if a-file data is not read.

getDiamagWp()

Returns diamagnetic-loop measured stored energy.

Raises ValueError – if a-file data is not read.

getDiamag()

Pulls diamagnetic flux, diamag. measured toroidal and poloidal beta, stored energy, and energy confinement time.

Returns namedtuple containing [diaFlux,diaBetat,diaBetap,diaTauE,diaWp]

Raises ValueError – if a-file data is not read

getWMHD()

Returns EFIT-calculated stored energy.

Raises ValueError – if a-file data is not read.

getTauMHD()

Returns EFIT-calculated energy confinement time.

Raises ValueError – if a-file data is not read.

getPinj()

Returns EFIT injected power.

Raises ValueError – if a-file data is not read.

getWbdot()

Returns EFIT d/dt of magnetic stored energy

Raises ValueError – if a-file data is not read.

getWpdot()

Returns EFIT d/dt of plasma stored energy.

Raises ValueError – if a-file data is not read.

getEnergy()

Pulls EFIT stored energy, energy confinement time, injected power, and d/dt of magnetic and plasma stored energy.

Returns namedtuple containing [WMHD,tauMHD,Pinj,Wbdot,Wpdot]

Raises ValueError – if a-file data is not read.

getParam (name)

Backup function, applying a direct path input for tree-like data storage access for parameters not typically found in Equilbrium object. Directly calls attributes read from g/a-files in copy-safe manner.

Parameters name - String. Parameter name for value stored in EqdskReader instance.

Raises AttributeError – raised if no attribute is found.

getMachineCrossSection()

Method to pull machine cross-section from data storage, convert to standard format for plotting routine.

getMachineCrossSectionFull()

Returns vectorization of machine cross-section.

Absent additional data (not found in eqdsks) simply returns self.getMachineCrossSection().

```
gfile (time=None, nw=None, nh=None, shot=None, name=None, tunit='ms', title='EQTOOLS', nbbbs=100)
```

```
plotFlux (fill=True, mask=True)
```

streamlined plotting of flux contours directly from psi grid

Parameters

- fill Boolean. Default True. Set True to plot filled contours of flux delineated by black outlines. Set False to instead plot color-coded line contours on a blank background.
- mask Boolean. Default True. Set True to draw a clipping mask based on the limiter outline for the flux contours. Set False to draw the full RZ grid.

eqtools.filewriter module

```
eqtools.filewriter.gfile(obj, tin, nw=None, nh=None, shot=None, name=None, tunit='ms', ti-
tle='EQTOOLS', nbbbs=100)
eqtools.filewriter.findLCFS(rgrid, zgrid, psiRZ, rcent, zcent, psiLCFS, nbbbs=100)
```

eqtools.pfilereader module

This module contains the PFileReader class, a lightweight data handler for p-file (radial profile) datasets.

Classes:

PFileReader: Data-storage class for p-file data. Reads data from ASCII p-file, storing as copy-safe object attributes.

```
class eqtools.pfilereader.PFileReader (pfile, verbose=True)
    Bases: object
```

Class to read ASCII p-file (profile data storage) into lightweight, user-friendly data structure.

P-files store data blocks containing the following: a header with parameter name, parameter units, x-axis units, and number of data points, followed by values of axis x, parameter y, and derivative dy/dx. Each parameter block is read into a namedtuple storing ['name','npts','units','xunits','x','y','dydx'], with each namedtuple stored as an attribute of the PFileReader instance. This gracefully handles variable formats of p-files (differing versions of p-files will have different parameters stored). Data blocks are accessed as attributes in a copy-safe manner.

Creates instance of PFileReader.

Parameters

- **pfile** String. Path to ASCII p-file to be loaded.
- **verbose** Boolean. Option to print message on object creation listing available data parameters. Defaults to True.

eqtools.trispline module

This module provides interface to the tricubic spline interpolator. It also contains an enhanced bivariate spline which generates bounds errors.

```
class eqtools.trispline.Spline (z, y, x, f, regular=True, fast=False)
```

Tricubic interpolating spline with forced edge derivative equal zero conditions. It assumes a cartesian grid.

Create a new Spline instance.

Parameters

- **z** (*1-dimensional float array*) Values of the positions of the 1st Dimension of f. Must be monotonic without duplicates.
- y (1-dimensional float array) Values of the positions of the 2nd dimension of f. Must be monotonic without duplicates.
- **x** (*1-dimensional float array*) Values of the positions of the 3rd dimension of f. Must be monotonic without duplicates.
- **f** (3-dimensional float array) f[z,y,x]. NaN and Inf will hamper performance and affect interpolation in 4x4x4 space about its value.
- regular If the grid is known to be regular, forces matrix-based fast evaluation of interpolation.
- fast Outdated input to test the indexing performance of the c code vs internal python handling.

Kwtype regular Boolean

Kwtype fast Boolean

Raises

- ValueError If any of the dimensions do not match specified f dim
- ValueError If x,y, or z are not monotonic

Examples

```
temp
```

```
ev (z1, y1, x1)
```

evaluates tricubic spline at point (x1,y1,z1) which is f[z1,y1,x1].

Parameters

- **z1** (*scalar float or 1-dimensional float*) Position in z dimension. (First dimension of 3d valued grid)
- y1 (scalar float or 1-dimensional float) Position in y dimension. (Second dimension of 3d valued grid)
- **x1** (*scalar float or 1-dimensional float*) Position in x dimension. (Third dimension of 3d valued grid)

Returns

val

• val (array or scalar float) - The interpolated value at (x1,y1,z1).

Raises ValueError - If any of the dimensions exceed the evaluation boundary of the grid

Examples

Data is grouped into the grid voxels so as to reuse calculated spline coefficients, thus speeding evaluation. It is recommended that it is evaluated outside of for loops to best utilize this feature.

```
class eqtools.trispline.RectBivariateSpline (x, y, z, bbox=[None, None, None, None], <math>kx=3, ky=3, s=0, bounds\_error=True, fill\_value=nan)

Bases: scipy.interpolate.fitpack2.RectBivariateSpline
```

the lack of a graceful bounds error causes the fortran to fail hard. This masks scipy.interpolate.RectBivariateSpline with a proper bound checker and value filler such that it will not fail in use for EqTools

Can be used for both smoothing and interpolating data.

Parameters

- x (1-dimensional float array) 1-D array of coordinates in monotonically increasing order.
- y (1-dimensional float array) 1-D array of coordinates in monotonically increasing order.
- **z** (2-dimensional float array) 2-D array of data with shape (x.size,y.size).
- **bbox** Sequence of length 4 specifying the boundary of the rectangular approximation domain. By default, bbox=[min(x,tx),max(x,tx), min(y,ty),max(y,ty)].
- **kx** Degrees of the bivariate spline. Default is 3.
- ky Degrees of the bivariate spline. Default is 3.
- s Positive smoothing factor defined for estimation condition, sum((w[i]*(z[i]-s(x[i], y[i])))**2, axis=0) <= s Default is s=0, which is for interpolation.

Kwtype bbox 1-dimensional float

Kwtype kx integer

Kwtype ky integer

Kwtype s float

ev(xi, yi)

Evaluate the rectBiVariateSpline at (xi,yi). (x,y)values are checked for being in the bounds of the interpolated data.

Parameters

- xi (float array) input x dimensional values
- yi (float array) input x dimensional values

Returns

```
evaluated spline at points (x[i], y[i]), i=0,...,len(x)-1
```

Return type val (float array)

Module contents

Provides classes for interacting with magnetic equilibrium data in a variety of formats.

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