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# **NIRFASTerFF**

***Release 1.0.1***

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NIRFAST was originally developed in 2001 as a MATLAB-based package used to model near-infrared light propagation through tissue and reconstruct images of optical biomarkers. At its core, NIRFAST is tool that does finite element modeling of the medium in which the light propagates, and calculates the fluence field by solving the diffusion equation.

A fully overhauled version, titled NIRFASTer, was published in 2018, thanks to the work of Dr. Stanisław Wojtkiewicz. In the new version, GPU support was added to the key algorithms, giving the software a dramatic boost in performance. The CPU versions of the algorithms were re-implemented in C++ with multithreading enabled, and the performance was improved considerably.

In this version, now titled NIRFASTerFF (Fast and Furious), the entire toolbox is re-written with Python as its interfacing language, while fully inter-operatable with the original Matlab version. The algorithms, running on both GPU and CPU, are yet again improved for even better performance.

This manual is a detailed documentation of all APIs in the package. Please also refer to the demos to see how the package is used in difference applications.



## SUMMARY OF THE FUNCTIONALITIES

Mesh types supported: standard, fluorescence, and DCS

FEM solver calculates: CW fluence, FD fluence, TPSF, direct TR moments for standard and fluorescence mesh, and G1/g1 curve for DCS mesh

Analytical solution in semi-infinite medium for: CW/FD fluence, TPSF, and DCS G1/g1 curves

Jacobian matrices: CW for standard, fluorescence, and DCS mesh, and FD for standard and fluorescence mesh





## **LINK TO THE MATLAB VERSION**

The original Matlab-based NIRFAST and NIRFASTer are still available for download, but we will gradually drop our support for them.

<https://github.com/nirfast-admin/NIRFAST>

<https://github.com/nirfaster/NIRFASTer>



## REFERENCES

If you use our package, please cite,

- H. Dehghani, M.E. Eames, P.K. Yalavarthy, S.C. Davis, S. Srinivasan, C.M. Carpenter, B.W. Pogue, and K.D. Paulsen, "Near infrared optical tomography using NIRFAST: Algorithm for numerical model and image reconstruction," Communications in Numerical Methods in Engineering, vol. 25, 711-732 (2009)  
[doi:10.1002/cnm.1162](https://doi.org/10.1002/cnm.1162)



## API DOCUMENTATION

### 4.1 nirfasterff

#### Modules

<i>nirfasterff.base</i>	Core classes used in the package
<i>nirfasterff.forward</i>	Functions for forward data calculation
<i>nirfasterff.inverse</i>	Calculation of the Jacobian matrices and a basic Tikhonov regularization function
<i>nirfasterff.io</i>	Some functions for reading/writing certain data types.
<i>nirfasterff.lib</i>	Low-level functions implemented in C/C++, on both GPU and CPU
<i>nirfasterff.math</i>	Some low-level functions used by the forward solvers.
<i>nirfasterff.meshing</i>	Functions used for mesh generation and quality check
<i>nirfasterff.utils</i>	Utility functions and auxiliary classes frequently used in the package.
<i>nirfasterff.visualize</i>	Functions for basic data visualization

#### 4.1.1 nirfasterff.base

Core classes used in the package

#### Modules

<i>nirfasterff.base.data</i>	Defining some data classes, which are the return types of the fem data calculation functions
<i>nirfasterff.base.dcs_mesh</i>	Define the DCS mesh class.
<i>nirfasterff.base.fluor_mesh</i>	Define the fluorescence mesh class
<i>nirfasterff.base.optodes</i>	Define the optode class, an instance of which can be either a source or a detector
<i>nirfasterff.base.stnd_mesh</i>	Define the standard mesh class

## nirfasterff.base.data

Defining some data classes, which are the return types of the fem data calculation functions

### Classes

<i>DCSdata()</i>	Class holding DCS data.
<i>FDdata()</i>	Class holding FD/CW data.
<i>FLdata()</i>	Class holding FD/CW fluorescence data.
<i>TPSFdata()</i>	Class holding time-resolved TPSF data.
<i>TRMomentsdata()</i>	Class holding time-resolved moments data calculated using Mellin transform.
<i>flTPSFdata()</i>	Class holding fluorescence time-resolved TPSF data.
<i>flTRMomentsdata()</i>	Class holding fluorescence TR moments data calculated using Mellin transform.
<i>meshvol()</i>	Small class holding the information needed for converting between mesh and volumetric space.

### nirfasterff.base.data.DCSdata

#### class nirfasterff.base.data.DCSdata

Bases: object

Class holding DCS data.

##### **phi**

steady-state fluence from each source. If mesh contains non-empty field vol, this will be represented on the grid.

Last dimension has the size of the number of sources

This is the same as nirfasterff.base.FDdata.phi, when modulation frequency is zero

##### **Type**

double NumPy array

##### **link**

Defining all the channels (i.e. source-detector pairs). Copied from mesh.link

##### **Type**

int32 NumPy array

##### **amplitude**

Steady-state amplitude of each channel. Size (NChannel,)

This is the same as nirfasterff.base.FDdata.amplitude, when modulation frequency is zero

##### **Type**

double NumPy vector

##### **tau\_DCS**

time vector in seconds

##### **Type**

double NumPy vector

**phi\_DCS**

G1 in medium from each source at each time step . If mesh contains non-empty field vol, this will be represented on the grid

shape[-1] equals length of tau\_DCS, and shape[-2] equals number of sources

**Type**

double NumPy array

**G1\_DCS**

G1 curve as is calculated from the correlation diffusion equation. Size: (NChannel, NTime)

**Type**

double NumPy array

**g1\_DCS**

g1 curve, i.e. G1 normalized by amplitudes. Size: (NChannel, NTime)

**Type**

double NumPy array

**vol**

Information needed to convert between volumetric and mesh space. Copied from mesh.vol

**Type**

nirfasterff.base.meshvol

**\_\_init\_\_()****Methods**

<code>__init__()</code>	
<code>isvol()</code>	Checks if data is in volumetric space.
<code>togrid(mesh)</code>	Convert data to volumetric space as is defined in mesh.vol.
<code>tomesh(mesh)</code>	Convert data back to mesh space using information defined in mesh.vol.

**isvol()**

Checks if data is in volumetric space.

**Returns**

True if data is in volumetric space, False if not.

**Return type**

bool

**togrid(mesh)**

Convert data to volumetric space as is defined in mesh.vol. If it is empty, the function does nothing.

If data is already in volumetric space, function casts data to the new volumetric space

CAUTION: This OVERRIDES the fields phi and phi\_DCS

**Parameters**

**mesh** (*nirfasterff.base.dcsmesh*) – mesh whose .vol attribute is used to do the conversion.

**Return type**

None.

**tomesh**(*mesh*)

Convert data back to mesh space using information defined in mesh.vol. If data.vol is empty, the function does nothing.

CAUTION: This OVERRIDES fields phi and phi\_DCS

**Parameters**

**mesh** (*nirfasterff.base.dcsmesh*) – mesh whose .vol attribute is used to do the conversion.

**Return type**

None.

**nirfasterff.base.data.FDdata****class nirfasterff.base.data.FDdata**

Bases: object

Class holding FD/CW data.

**phi**

Fluence from each source. If mesh contains non-empty field vol, this will be represented on the grid. Last dimension has the size of the number of sources

**Type**

double Numpy array

**complex**

Complex amplitude of each channel. Same as amplitude in case of CW data

**Type**

double or complex double Numpy vector

**link**

Defining all the channels (i.e. source-detector pairs). Copied from mesh.link

**Type**

int32 NumPy array

**amplitude**

Absolute amplitude of each channel. I.e. amplitude=abs(complex)

**Type**

double Numpy vector

**phase**

phase data of each channel. All zero in case of CW data

**Type**

double Numpy vector

**vol**

Information needed to convert between volumetric and mesh space. Copied from mesh.vol

**Type**

nirfasterff.base.meshvol



`__init__()`

## Methods

<code>__init__()</code>	
<code>isvol()</code>	Checks if data is in volumetric space.
<code>togrid(mesh)</code>	Convert data to volumetric space as is defined in mesh.vol.
<code>tomesh(mesh)</code>	Convert data back to mesh space using information defined in mesh.vol.

`isvol()`

Checks if data is in volumetric space.

### Returns

True if data is in volumetric space, False if not.

### Return type

bool

`togrid(mesh)`

Convert data to volumetric space as is defined in mesh.vol. If it is empty, the function does nothing.

If data is already in volumetric space, function casts data to the new volumetric space

CAUTION: This OVERRIDES the field phi

### Parameters

**mesh** (*nirfasterff.base.stndmesh*) – mesh whose .vol attribute is used to do the conversion.

### Return type

None.

`tomesh(mesh)`

Convert data back to mesh space using information defined in mesh.vol. If data.vol is empty, the function does nothing.

CAUTION: This OVERRIDES the field phi

### Parameters

**mesh** (*nirfasterff.base.stndmesh*) – mesh whose .vol attribute is used to do the conversion.

### Return type

None.

**nirfasterff.base.data.FLdata****class** nirfasterff.base.data.FLdata

Bases: object

Class holding FD/CW fluorescence data.

**phix**

intrinsic fluence from each source at excitation wavelength. If mesh contains non-empty field vol, this will be represented on the grid. Last dimension has the size of the number of sources

**Type**

double Numpy array

**phimm**

intrinsic from each source at emission wavelength.

**Type**

double Numpy array

**phi fl**

fluorescence emission fluence

**Type**

double Numpy array

**complexxx**

Complex amplitude of each channel, intrinsic excitation. Same as amplitude in case of CW data

**Type**

double or complex double Numpy vector

**complexmm**

Complex amplitude of each channel, intrinsic emission. Same as amplitude in case of CW data

**Type**

double or complex double Numpy vector

**complexfl**

Complex amplitude of each channel, fluorescence emission. Same as amplitude in case of CW data

**Type**

double or complex double Numpy vector

**link**

Defining all the channels (i.e. source-detector pairs). Copied from mesh.link

**Type**

int32 NumPy array

**amplitudex**

Absolute amplitude of each channel, intrinsic excitation. I.e. `amplitudex=abs(complexxx)`

**Type**

double Numpy vector

**amplitudemmm**

Absolute amplitude of each channel, intrinsic emission. I.e. `amplitudemmm=abs(complexmm)`

**Type**

double Numpy vector

**amplitudefl**

Absolute amplitude of each channel, fluorescence emission. I.e. `amplitudefl=abs(complexfl)`

**Type**

double Numpy vector

**phasesx**

phase data of each channel, intrinsic excitation. All zero in case of CW data

**Type**

double Numpy vector

**phasemm**

phase data of each channel, intrinsic emission. All zero in case of CW data

**Type**

double Numpy vector

**phasefl**

phase data of each channel, fluorescence emission. All zero in case of CW data

**Type**

double Numpy vector

**vol**

Information needed to convert between volumetric and mesh space. Copied from `mesh.vol`

**Type**

`nirfasterff.base.meshvol`

**\_\_init\_\_()****Methods**

<code>__init__()</code>	
<code>isvol()</code>	Checks if data is in volumetric space.
<code>togrid(mesh)</code>	Convert data to volumetric space as is defined in <code>mesh.vol</code> .
<code>tomesh(mesh)</code>	Convert data back to mesh space using information defined in <code>mesh.vol</code> .

**isvol()**

Checks if data is in volumetric space.

**Returns**

True if data is in volumetric space, False if not.

**Return type**

bool

**togrid(mesh)**

Convert data to volumetric space as is defined in `mesh.vol`. If it is empty, the function does nothing.

If data is already in volumetric space, function casts data to the new volumetric space

CAUTION: This OVERRIDES the fields `phix`, `phimm`, and `phifl`, if they are defined

**Parameters**

**mesh** (*nirfasterff.base.fluormesh*) – mesh whose .vol attribute is used to do the conversion.

**Return type**

None.

**tomesh**(*mesh*)

Convert data back to mesh space using information defined in mesh.vol. If data.vol is empty, the function does nothing.

CAUTION: This OVERRIDES fields phix, phimm, and phifl, if they are defined

**Parameters**

**mesh** (*nirfasterff.base.fluormesh*) – mesh whose .vol attribute is used to do the conversion.

**Return type**

None.

**nirfasterff.base.data.TPSFdata****class nirfasterff.base.data.TPSFdata**

Bases: object

Class holding time-resolved TPSF data.

**phi**

TPSF from each source at each spatial location. If mesh contains non-empty field vol, this will be represented on the grid

Shape: NNodes x num\_sources x time\_steps

OR: len(xgrid) x len(ygrid) x len(zgrid) x num\_sources x time\_steps

None by default, and only contains data if 'field' option is set to True when calculating forward data.

**Type**

double NumPy array or None

**time**

time vector, in seconds

**Type**

double NumPy vector

**tpsf**

TPSF measured at each channel. Size: (NChannels, time\_steps)

**Type**

double NumPy array

**link**

Defining all the channels (i.e. source-detector pairs). Copied from mesh.link

**Type**

int32 NumPy array

**vol**

Information needed to convert between volumetric and mesh space. Copied from mesh.vol

**Type**

nirfasterff.base.meshvol

**\_\_init\_\_()****Methods**

<code>__init__()</code>	
<code>isvol()</code>	Checks if data is in volumetric space.
<code>togrid(mesh)</code>	Convert data to volumetric space as is defined in mesh.vol.
<code>tomesh(mesh)</code>	Convert data back to mesh space using information defined in mesh.vol.

**isvol()**

Checks if data is in volumetric space.

**Returns**

True if data is in volumetric space, False if not.

**Return type**

bool

**togrid(mesh)**

Convert data to volumetric space as is defined in mesh.vol. If it is empty or data.phi==None, the function does nothing.

If data is already in volumetric space, function casts data to the new volumetric space

CAUTION: This OVERRIDES the field phi, if it is defined

**Parameters**

**mesh** (*nirfasterff.base.stndmesh*) – mesh whose .vol attribute is used to do the conversion.

**Return type**

None.

**tomesh(mesh)**

Convert data back to mesh space using information defined in mesh.vol. If data.vol is empty or data.phi==None, the function does nothing.

CAUTION: This OVERRIDES field phi, if it is defined

**Parameters**

**mesh** (*nirfasterff.base.fluormesh*) – mesh whose .vol attribute is used to do the conversion.

**Return type**

None.

**nirfasterff.base.data.TRMomentsdata****class** nirfasterff.base.data.TRMomentsdata

Bases: object

Class holding time-resolved moments data calculated using Mellin transform.

**phi**

moments from each source at each spatial location. If mesh contains non-empty field vol, this will be represented on the grid

Shape: NNodes x num\_sources x (max\_moment\_order + 1)

OR: len(xgrid) x len(ygrid) x len(zgrid) x num\_sources x (max\_moment\_order + 1)

None by default, and only contains data if 'field' option is set to True when calculating forward data.

**Type**

double Numpy array or None

**moments**

moments for each channel. i-th column contains i-th moment as measured at each channel. Size: (NChannels, max\_moment\_order + 1)

**Type**

double Numpy vector

**link**

Defining all the channels (i.e. source-detector pairs). Copied from mesh.link

**Type**

int32 NumPy array

**vol**

Information needed to convert between volumetric and mesh space. Copied from mesh.vol

**Type**

nirfasterff.base.meshvol

**\_\_init\_\_()****Methods**

<code>__init__()</code>	
<code>isvol()</code>	Checks if data is in volumetric space.
<code>togrid(mesh)</code>	Convert data to volumetric space as is defined in mesh.vol.
<code>tomesh(mesh)</code>	Convert data back to mesh space using information defined in mesh.vol.

**isvol()**

Checks if data is in volumetric space.

**Returns**

True if data is in volumetric space, False if not.

**Return type**

bool

**togrid**(*mesh*)

Convert data to volumetric space as is defined in *mesh.vol*. If it is empty or *data.phi==None*, the function does nothing.

If data is already in volumetric space, function casts data to the new volumetric space

CAUTION: This OVERRIDES the field *phi*, if it is defined

**Parameters**

**mesh** (*nirfasterff.base.stndmesh*) – mesh whose *.vol* attribute is used to do the conversion.

**Return type**

None.

**tomesh**(*mesh*)

Convert data back to mesh space using information defined in *mesh.vol*. If *data.vol* is empty or *data.phi==None*, the function does nothing.

CAUTION: This OVERRIDES field *phi*, if it is defined

**Parameters**

**mesh** (*nirfasterff.base.fluormesh*) – mesh whose *.vol* attribute is used to do the conversion.

**Return type**

None.

**nirfasterff.base.data.fITPSFdata****class** nirfasterff.base.data.fITPSFdata

Bases: object

Class holding fluorescence time-resolved TPSF data.

**phix**

TPSF at excitation wavelength from each source at each spatial location. If mesh contains non-empty field *vol*, this will be represented on the grid

Shape: NNodes x num\_sources x time\_steps

OR: len(xgrid) x len(ygrid) x len(zgrid) x num\_sources x time\_steps

None by default, and only contains data if 'field' option is set to True when calculating forward data.

**Type**

double NumPy array

**phi**

similar to *phix*, but for fluorescence emission

**Type**

double NumPy array

**time**

time vector, in seconds

**Type**

double NumPy vector

**tpsfx**

TPSF measured at each channel, excitation. Size: (NChannels, time\_steps)

**Type**

double NumPy array

**link**

Defining all the channels (i.e. source-detector pairs). Copied from mesh.link

**Type**

int32 NumPy array

**vol**

Information needed to convert between volumetric and mesh space. Copied from mesh.vol

**Type**

nirfasterff.base.meshvol

**\_\_init\_\_()****Methods**

<code>__init__()</code>	
<code>isvol()</code>	Checks if data is in volumetric space.
<code>togrid(mesh)</code>	Convert data to volumetric space as is defined in mesh.vol.
<code>tomesh(mesh)</code>	Convert data back to mesh space using information defined in mesh.vol.

**isvol()**

Checks if data is in volumetric space.

**Returns**

True if data is in volumetric space, False if not.

**Return type**

bool

**togrid(*mesh*)**

Convert data to volumetric space as is defined in mesh.vol. If it is empty or data.phix==None, the function does nothing.

If data is already in volumetric space, function casts data to the new volumetric space

CAUTION: This OVERRIDES the fields phix and phifl, if they are defined

**Parameters**

**mesh** (*nirfasterff.base.fluormesh*) – mesh whose .vol attribute is used to do the conversion.

**Return type**

None.



**tomesh**(*mesh*)

Convert data back to mesh space using information defined in `mesh.vol`. If `data.vol` is empty or `data.phix==None`, the function does nothing.

CAUTION: This OVERRIDES fields `phix` and `phiFl`, if they are defined

**Parameters**

**mesh** (*nirfasterff.base.fluormesh*) – mesh whose `.vol` attribute is used to do the conversion.

**Return type**

None.

**nirfasterff.base.data.flTRMomentsdata****class** nirfasterff.base.data.flTRMomentsdata

Bases: object

Class holding fluorescence TR moments data calculated using Mellin transform.

**phix**

moments at excitation wavelength from each source at each spatial location. If mesh contains non-empty field `vol`, this will be represented on the grid

Shape: `NNodes x num_sources x (max_moment_order + 1)`

OR: `len(xgrid) x len(ygrid) x len(zgrid) x num_sources x (max_moment_order + 1)`

None by default, and only contains data if ‘field’ option is set to True when calculating forward data.

**Type**

double NumPy array

**phiFl**

similar to `phix`, but for fluorescence emission

**Type**

double NumPy array

**momentsx**

moments for each channel, excitation. `i`-th column contains `i`-th moment. Size: (`NChannels`, `max_moment_order + 1`)

**Type**

double NumPy array

**momentsfL**

moments for each channel, fluorescence emission. `i`-th column contains `i`-th moment. Size: (`NChannels`, `max_moment_order + 1`)

**Type**

double NumPy array

**link**

Defining all the channels (i.e. source-detector pairs). Copied from `mesh.link`

**Type**

int32 NumPy array

**vol**

Information needed to convert between volumetric and mesh space. Copied from mesh.vol

**Type**

nirfasterff.base.meshvol

**\_\_init\_\_()****Methods**

<code>__init__()</code>	
<code>isvol()</code>	Checks if data is in volumetric space.
<code>togrid(mesh)</code>	Convert data to volumetric space as is defined in mesh.vol.
<code>tomesh(mesh)</code>	Convert data back to mesh space using information defined in mesh.vol.

**isvol()**

Checks if data is in volumetric space.

**Returns**

True if data is in volumetric space, False if not.

**Return type**

bool

**togrid(*mesh*)**

Convert data to volumetric space as is defined in mesh.vol. If it is empty or data.phix==None, the function does nothing.

If data is already in volumetric space, function casts data to the new volumetric space

CAUTION: This OVERRIDES the fields phix and phifl, if they are defined

**Parameters**

**mesh** (*nirfasterff.base.fluormesh*) – mesh whose .vol attribute is used to do the conversion.

**Return type**

None.

**tomesh(*mesh*)**

Convert data back to mesh space using information defined in mesh.vol. If data.vol is empty or data.phix==None, the function does nothing.

CAUTION: This OVERRIDES fields phix and phifl, if they are defined

**Parameters**

**mesh** (*nirfasterff.base.fluormesh*) – mesh whose .vol attribute is used to do the conversion.

**Return type**

None.

**nirfasterff.base.data.meshvol****class** nirfasterff.base.data.**meshvol**

Bases: object

Small class holding the information needed for converting between mesh and volumetric space. Values calculated by nirfasterff.base.\*mesh.gen\_intmat

Note that the volumetric space, defined by xgrid, ygrid, and zgrid (empty for 2D mesh), must be uniform

**xgrid**

x grid of the volumetric space. In mm

**Type**

double Numpy array

**ygrid**

y grid of the volumetric space. In mm

**Type**

double Numpy array

**zgrid**

z grid of the volumetric space. In mm. Empty for 2D meshes

**Type**

double Numpy array

**mesh2grid**

matrix converting a vector in mesh space to volumetric space, done by mesh2grid.dot(data)

The result is vectorized in 'F' (Matlab) order

Size: (len(xgrid)\*len(ygrid)\*len(zgrid), NNodes)

**Type**

double CSR sparse matrix

**gridinmesh**

indices (one-based) of data points in the volumetric space that are within the mesh space, vectorized in 'F' order.

**Type**

int32 NumPy array

**res**

resolution in x, y, z (if 3D) direction, in mm. Size (2,) or (3,)

**Type**

double NumPy array

**grid2mesh**

matrix converting volumetric data, vectorized in 'F' order, to mesh space. Done by grid2mesh.dot(data)

Size (Nnodes, len(xgrid)\*len(ygrid)\*len(ygrid))

**Type**

double CSR sparse matrix

**meshingrid**

indices (one-based) of data points in the mesh space that are within the volumetric space

**Type**

int32 NumPy array

`__init__()`

**Methods**

`__init__()`

**nirfasterff.base.dcs\_mesh**

Define the DCS mesh class. Assuming all motions are Brownian.

**Classes**

`dcsmesh()`

Main class for standard mesh.

**nirfasterff.base.dcs\_mesh.dcsmesh**

**class** nirfasterff.base.dcs\_mesh.dcsmesh

Bases: object

Main class for standard mesh. The methods should cover most of the commonly-used functionalities

**name**

name of the mesh. Default: 'EmptyMesh'

**Type**

str

**nodes**

locations of nodes in the mesh. Unit: mm. Size (NNodes, dim)

**Type**

double NumPy array

**bndvtx**

indicator of whether a node is at boundary (1) or internal (0). Size (NNodes,)

**Type**

double NumPy array

**type**

type of the mesh. It is always 'dcs'.

**Type**

str

**mua**

absorption coefficient ( $\text{mm}^{-1}$ ) at each node. Size (NNodes,)

**Type**

double NumPy array

**kappa**

diffusion coefficient (mm) at each node. Size (NNodes,). Defined as  $1/(3*(\text{mua} + \text{mus}))$

**Type**

double NumPy array

**ri**

refractive index at each node. Size (NNodes,)

**Type**

double NumPy array

**mus**

reduced scattering coefficient ( $\text{mm}^{-1}$ ) at each node. Size (NNodes,)

**Type**

(double NumPy array

**wv\_DCS**

wavelength used (nm)

**Type**

double

**alpha**

fraction of photon scattering events that occur from moving particles in the medium (a.u.). Size (NNodes, NFlow)

**Type**

double NumPy array

**Db**

effective diffusion coefficient in Brownian motion ( $\text{mm}^2/\text{s}$ ). Size (NNodes, NFlow)

**Type**

double NumPy array

**aDb**

Defined as  $\text{np.sum}(a*Db, \text{axis}=1)$ . This lumped parameter ( $\text{mm}^2/\text{s}$ ) is what is actually used in data generation. Size (NNodes,)

**Type**

double NumPy array

**elements**

triangulation (tetrahedrons or triangles) of the mesh, Size (NElements, dim+1)

Row i contains the indices of the nodes that form tetrahedron/triangle i

One-based indexing for direct interoperability with the Matlab version

**Type**

double NumPy array

**region**

region labeling of each node. Starting from 1. Size (NNodes,)

**Type**

double NumPy array

**source**

information about the sources

**Type**

nirfasterff.base.optode

**meas**

information about the the detectors

**Type**

nirfasterff.base.optode

**link**

list of source-detector pairs, i.e. channels. Size (NChannels,3)

First column: source; Second column: detector; Third column: active (1) or not (0)

**Type**

int32 NumPy array

**c**

light speed (mm/sec) at each node. Size (NNodes,). Defined as  $c_0/r_i$ , where  $c_0$  is the light speed in vacuum

**Type**

double NumPy array

**ksi**

photon fluence rate scale factor on the mesh-outside\_mesh boundary as derived from Fresnel's law. Size (NNodes,)

**Type**

double NumPy array

**element\_area**

volume/area ( $\text{mm}^3$  or  $\text{mm}^2$ ) of each element. Size (NElements,)

**Type**

double NumPy array

**support**

total volume/area of all the elements each node belongs to. Size (NNodes,)

**Type**

double NumPy array

**vol**

object holding information for converting between mesh and volumetric space.

**Type**

nirfasterff.base.meshvol

**\_\_init\_\_()**

## Methods

<code>__init__()</code>	
<code>change_prop(idx, prop)</code>	Change optical properties (mua, musp, ri, alpha and Db) at nodes specified in idx, and automatically change fields kappa, c, and ksi as well
<code>femdata(tvec[, solver, opt])</code>	Calculates steady-state fluences and G1/g1 curves for each source using a FEM solver, and then the boudary measurables for each channel
<code>from_copy(mesh)</code>	Deep copy all fields from another mesh.
<code>from_file(file)</code>	Read from classic NIRFAST mesh (ASCII) format, not checking the correctness of the loaded integration functions.
<code>from_mat(matfile[, varname])</code>	Read from Matlab .mat file that contains a NIRFASTer mesh struct.
<code>from_solid(ele, nodes[, prop, src, det, link])</code>	Construct a NIRFASTer mesh from a 3D solid mesh generated by a mesher.
<code>from_volume(vol[, param, prop, src, det, link])</code>	Construct mesh from a segmented 3D volume using the built-in CGAL mesher.
<code>gen_intmat(xgrid, ygrid[, zgrid])</code>	Calculate the information needed to convert data between mesh and volumetric space, specified by x, y, z (if 3D) grids.
<code>isvol()</code>	Check if conversion matrices between mesh and volumetric spaces are calculated
<code>jacobian(tvec[, normalize, solver, opt])</code>	Calculates the Jacobian matrix
<code>save_nirfast(filename)</code>	Save mesh in the classic NIRFASTer ASCII format, which is directly compatible with the Matlab version
<code>set_prop(prop)</code>	Set optical properties of the whole mesh, using information provided in prop.
<code>touch_optodes()</code>	Moves all optodes (if non fixed) and recalculate the integration functions (i.e. barycentric coordinates).

### `change_prop(idx, prop)`

Change optical properties (mua, musp, ri, alpha and Db) at nodes specified in idx, and automatically change fields kappa, c, and ksi as well

#### Parameters

- **idx** (*list or NumPy array or -1*) – zero-based indices of nodes to change. If *idx* == -1, function changes all the nodes
- **prop** (*list or NumPy array of length 6*) – new optical properties to be assigned to the specified nodes. [region mua(mm-1) musp(mm-1) ri alpha Db].

#### Return type

None.

### `femdata(tvec, solver=utils.get_solver(), opt=utils.SolverOptions())`

Calculates steady-state fluences and G1/g1 curves for each source using a FEM solver, and then the boudary measurables for each channel

Assumes Brownian motion, that is,  $\langle \Delta r^2 \rangle = 6 * \alpha Db * \tau$

If *mesh.vol* is set, fluence and G1 data will be represented in volumetric space

See `femdata_DCS()` and `femdata_stnd_CW()` for details

#### Parameters

- **tvec** (*double NumPy array*) – time vector (i.e. :math:`\tau` au) for the G1 curve, in seconds. It is usually a good idea to use log scale
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (`nirfasterff.utils.SolverOptions`, *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See `SolverOptions()` for details

#### Returns

- **data** (`nirfasterff.base.DCSdata`) – contains fluence, G1 curve, and g1 curve calculated at each spatial location, and also the boundary data.

See `DCSdata()` for details.

- **info** (`nirfasterff.utils.ConvergenceInfo`) – convergence information of the solver when calculating the fluence field.

See `ConvergenceInfo()` for details

#### `from_copy(mesh)`

Deep copy all fields from another mesh.

##### Parameters

**mesh** (`nirfasterff.base.stndmesh`) – the mesh to copy from.

##### Return type

None.

#### `from_file(file)`

Read from classic NIRFAST mesh (ASCII) format, not checking the correctness of the loaded integration functions.

All fields after loading should be directly compatible with Matlab version.

##### Parameters

**file** (*str*) – name of the mesh. Any extension will be ignored.

##### Return type

None.

### Examples

```
>>> mesh = nirfasterff.base.dcsmesh()
>>> mesh.from_file('meshname')
```

#### `from_mat(matfile, varname=None)`

Read from Matlab .mat file that contains a NIRFASTer mesh struct. All fields copied as is without error checking.

##### Parameters

- **matfile** (*str*) – name of the .mat file to load. Use of extension is optional.



- **varname** (*str*, *optional*) – if your .mat file contains multiple variables, use this argument to specify which one to load. The default is None.

When *varname==None*, *matfile* should contain exactly one structure, which is a NIRFASTer mesh, or the function will do nothing

#### Return type

None.

**from\_solid**(*ele*, *nodes*, *prop=None*, *src=None*, *det=None*, *link=None*)

Construct a NIRFASTer mesh from a 3D solid mesh generated by a mesher. Similar to the `solidmesh2nirfast` function in Matlab version.

Can also set the optical properties and optodes if supplied

#### Parameters

- **ele** (*int/double NumPy array*) – element list in one-based indexing. If four columns, all nodes will be labeled as region 1

If five columns, the last column will be used for region labeling.

- **nodes** (*double NumPy array*) – node locations in the mesh. Unit: mm. Size (NNodes,3).

- **prop** (*double NumPy array, optional*) – If not *None*, calls `dcsmesh.set_prop()` and sets the optical properties in the mesh. The default is None.

See [set\\_prop\(\)](#) for details.

- **src** (*nirfasterff.base.optode, optional*) – If not *None*, sets the sources and moves them to the appropriate locations. The default is None.

See [touch\\_sources\(\)](#) for details.

- **det** (*nirfasterff.base.optode, optional*) – If not *None*, sets the detectors and moves them to the appropriate locations. The default is None.

See [touch\\_detectors\(\)](#) for details.

- **link** (*int32 NumPy array, optional*) – If not *None*, sets the channel information. Uses one-based indexing. The default is None.

Each row represents a channel, in the form of [*src*, *det*, *active*], where *active* is 0 or 1

If *link* contains only two columns, all channels are considered active.

#### Return type

None.

**from\_volume**(*vol*, *param=utils.MeshingParams()*, *prop=None*, *src=None*, *det=None*, *link=None*)

Construct mesh from a segmented 3D volume using the built-in CGAL mesher. Calls `stndmesh.from_solid` after meshing step.

#### Parameters

- **vol** (*uint8 NumPy array*) – 3D segmented volume to be meshed. 0 is considered as outside. Regions labeled using unique integers.

- **param** (*nirfasterff.utils.MeshingParams, optional*) – parameters used in the CGAL mesher. If not specified, uses the default parameters defined in `nirfasterff.utils.MeshingParams()`.

Please modify fields `xPixelSpacing`, `yPixelSpacing`, and `SliceThickness` if your volume doesn't have [1,1,1] resolution

See [MeshingParams\(\)](#) for details.

- **prop** (*double NumPy array, optional*) – If not *None*, calls *dcsmesh.set\_prop()* and sets the optical properties in the mesh. The default is *None*.

See [set\\_prop\(\)](#) for details.

- **src** (*nirfasterff.base.optode, optional*) – If not *None*, sets the sources and moves them to the appropriate locations. The default is *None*.

See [touch\\_sources\(\)](#) for details.

- **det** (*nirfasterff.base.optode, optional*) – If not *None*, sets the detectors and moves them to the appropriate locations. The default is *None*.

See [touch\\_detectors\(\)](#) for details.

- **link** (*int32 NumPy array, optional*) – If not *None*, sets the channel information. Uses one-based indexing. The default is *None*.

Each row represents a channel, in the form of *[src, det, active]*, where *active* is 0 or 1

If *link* contains only two columns, all channels are considered active.

#### Return type

*None*.

#### **gen\_intmat** (*xgrid, ygrid, zgrid=[]*)

Calculate the information needed to convert data between mesh and volumetric space, specified by x, y, z (if 3D) grids.

All grids must be uniform. The results will from a *nirfasterff.base.meshvol* object stored in field *.vol*

If field *.vol* already exists, it will be calculated again, and a warning will be thrown

#### Parameters

- **xgrid** (*double NumPy array*) – x grid in mm.
- **ygrid** (*double NumPy array*) – y grid in mm.
- **zgrid** (*double NumPy array, optional*) – z grid in mm. Leave empty for 2D meshes. The default is *[]*.

#### Raises

**ValueError** – if grids not uniform, or *zgrid* empty for 3D mesh

#### Return type

*None*.

#### **isvol** ()

Check if conversion matrices between mesh and volumetric spaces are calculated

#### Returns

True if attribute *.vol* is calculate, False if not.

#### Return type

bool

#### **jacobian** (*tvec, normalize=True, solver=utils.get\_solver(), opt=utils.SolverOptions()*)

Calculates the Jacobian matrix

Returns the Jacobian, direct field data, and the adjoint data

One Jacobian is calculated at each time point in *tvec*, and the derivative is taken with regard to *aDb*

**Parameters**

- **tvec** (*double NumPy vector*) – time vector used.
- **normalize** (*bool, optional*) – if True, Jacobians are normalized to the measured boundary amplitude. The default is True.
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

**Returns**

- **J** (*double NumPy array*) – The Jacobian matrix. Size (NChannel, NVoxel, NTime)
- **data1** (*nirfasterff.base.FLdata*) – The calculated direct field. The same as directly calling `mesh.femdata(tvec)`
- **data2** (*nirfasterff.base.FLdata*) – The calculated adjoint field. The same as calling `mesh.femdata(tvec)` AFTER swapping the locations of sources and detectors

**See also:**

[jacobian\\_DCS\(\)](#)

**save\_nirfast**(*filename*)

Save mesh in the classic NIRFASTer ASCII format, which is directly compatible with the Matlab version

**Parameters**

**filename** (*str*) – name of the file to be saved as. Should have no extensions.

**Return type**

None.

**set\_prop**(*prop*)

Set optical properties of the whole mesh, using information provided in prop.

**Parameters**

**prop** (*double NumPy array*) – optical property info, similar to the MCX format:

```
[region mua(mm-1) musp(mm-1) ri alpha Db]
[region mua(mm-1) musp(mm-1) ri alpha Db]
[...]
```

where ‘region’ is the region label, and they should match exactly with `unique(mesh.region)`. The order doesn’t matter.

**Return type**

None.

**touch\_optodes**()

Moves all optodes (if non fixed) and recalculate the integration functions (i.e. barycentric coordinates).

See [touch\\_sources\(\)](#) and [touch\\_detectors\(\)](#) for details

**Return type**

None.

**nirfasterff.base.fluor\_mesh**

Define the fluorescence mesh class

**Classes**

---

*fluormesh()*Main class for fluorescence mesh.

---

**nirfasterff.base.fluor\_mesh.fluormesh****class** nirfasterff.base.fluor\_mesh.**fluormesh**

Bases: object

Main class for fluorescence mesh. The methods should cover most of the commonly-used functionalities

**name**

name of the mesh. Default: 'EmptyMesh'

**Type**

str

**nodes**

locations of nodes in the mesh. Unit: mm. Size (NNodes, dim)

**Type**

double NumPy array

**bndvtx**

indicator of whether a node is at boundary (1) or internal (0). Size (NNodes,)

**Type**

double NumPy array

**type**

type of the mesh. It is always 'fluor'.

**Type**

str

**muax**intrinsic absorption coefficient (mm<sup>-1</sup>) at excitation wavelength at each node. Size (NNodes,)**Type**

double NumPy array

**muam**intrinsic absorption coefficient (mm<sup>-1</sup>) at emission wavelength at each node. Size (NNodes,)**Type**

double NumPy array

**muaf**intrinsic absorption coefficient (mm<sup>-1</sup>) of the fluorophores at each node. Size (NNodes,)**Type**

double NumPy array

**kappax**

intrinsic diffusion coefficient (mm) at excitation wavelength at each node. Size (NNodes,). Defined as  $1/(3*(\mu_{ax} + \mu_{sx}))$

**Type**

double NumPy array

**kappam**

intrinsic diffusion coefficient (mm) at emission wavelength at each node. Size (NNodes,). Defined as  $1/(3*(\mu_{am} + \mu_{sm}))$

**Type**

double NumPy array

**ri**

refractive index at each node. Size (NNodes,)

**Type**

double NumPy array

**musx**

intrinsic reduced scattering coefficient ( $\text{mm}^{-1}$ ) at excitation wavelength at each node. Size (NNodes,)

**Type**

double NumPy array

**musm**

intrinsic reduced scattering coefficient ( $\text{mm}^{-1}$ ) at emission wavelength at each node. Size (NNodes,)

**Type**

double NumPy array

**eta**

quantum efficiency (a.u.) of the fluorophores. Size (NNodes,)

**Type**

double NumPy array

**tau**

time coefficient (second) of the fluorophores. Size (NNodes,)

**Type**

double NumPy array

**elements**

triangulation (tetrahedrons or triangles) of the mesh, Size (NElements, dim+1)

Row i contains the indices of the nodes that form tetrahedron/triangle i

One-based indexing for direct interoperability with the Matlab version

**Type**

double NumPy array

**region**

region labeling of each node. Starting from 1. Size (NNodes,)

**Type**

double NumPy array

**source**

information about the sources

**Type**

nirfasterff.base.optode

**meas**

information about the the detectors

**Type**

nirfasterff.base.optode

**link**

list of source-detector pairs, i.e. channels. Size (NChannels,3)

First column: source; Second column: detector; Third column: active (1) or not (0)

**Type**

int32 NumPy array

**c**

light speed (mm/sec) at each node. Size (NNodes,). Defined as  $c_0/r_i$ , where  $c_0$  is the light speed in vacuum

**Type**

double NumPy array

**ksi**

photon fluence rate scale factor on the mesh-outside\_mesh boundary as derived from Fresnel's law. Size (NNodes,)

**Type**

double NumPy array

**element\_area**

volume/area ( $\text{mm}^3$  or  $\text{mm}^2$ ) of each element. Size (NElements,)

**Type**

double NumPy array

**support**

total volume/area of all the elements each node belongs to. Size (NNodes,)

**Type**

double NumPy array

**vol**

object holding information for converting between mesh and volumetric space.

**Type**

nirfasterff.base.meshvol

**\_\_init\_\_()**

## Methods

<code>__init__()</code>	
<code>change_prop(idx, prop)</code>	Change optical properties (muax, musxp, ri, muam, musmp, muaf, eta, and tau) at nodes specified in idx, and automatically change fields kappax, kappam, c, and ksi as well
<code>femdata(freq[, solver, opt, xflag, mmflag, ...])</code>	Calculates fluences for each source using a FEM solver, and then the boudary measurables for each channel
<code>femdata_moments([max_moments, savefield, ...])</code>	Calculates TR moments at each location in the mesh for each source directly using Mellin transform, and then the boudary measurables for each channel
<code>femdata_tpsf(tmax, dt[, savefield, ...])</code>	Calculates TPSF at each location in the mesh for each source using a FEM solver, and then the boudary measurables for each channel
<code>from_copy(mesh)</code>	Deep copy all fields from another mesh.
<code>from_file(file)</code>	Read from classic NIRFAST mesh (ASCII) format, not checking the correctness of the loaded integration functions.
<code>from_mat(matfile[, varname])</code>	Read from Matlab .mat file that contains a NIRFASTer mesh struct.
<code>from_solid(ele, nodes[, prop, src, det, link])</code>	Construct a NIRFASTer mesh from a 3D solid mesh generated by a mesher.
<code>from_volume(vol[, param, prop, src, det, link])</code>	Construct mesh from a segmented 3D volume using the built-in CGAL mesher.
<code>gen_intmat(xgrid, ygrid[, zgrid])</code>	Calculate the information needed to convert data between mesh and volumetric space, specified by x, y, z (if 3D) grids.
<code>isvol()</code>	Check if conversion matrices between mesh and volumetric spaces are calculated
<code>jacobian([freq, normalize, solver, opt])</code>	Calculates the Jacobian matrix
<code>save_nirfast(filename)</code>	Save mesh in the classic NIRFASTer ASCII format, which is directly compatible with the Matlab version
<code>set_prop(prop)</code>	Set optical properties of the whole mesh, using information provided in prop.
<code>touch_optodes()</code>	Moves all optodes (if non fixed) and recalculate the integration functions (i.e. barycentric coordinates).

### `change_prop(idx, prop)`

Change optical properties (muax, musxp, ri, muam, musmp, muaf, eta, and tau) at nodes specified in idx, and automatically change fields kappax, kappam, c, and ksi as well

#### Parameters

- **idx** (*list or NumPy array or -1*) – zero-based indices of nodes to change. If *idx* == -1, function changes all the nodes
- **prop** (*list or NumPy array of length 8*) – new optical properties to be assigned to the specified nodes. [muax(mm-1) musxp(mm-1) ri muam(mm-1) musmp(mm-1) muaf(mm-1) eta tau(sec)].

#### Return type

None.

**femdata**(*freq*, *solver*=*utils.get\_solver()*, *opt*=*utils.SolverOptions()*, *xflag*=*True*, *mmflag*=*True*, *fflag*=*True*)

Calculates fluences for each source using a FEM solver, and then the boudary measurables for each channel

If *mesh.vol* is set, fluence data will be represented in volumetric space

See [femdata\\_fl\\_CW\(\)](#) and [femdata\\_fl\\_FD\(\)](#) for details

#### Parameters

- **freq** (*double*) – modulation frequency in Hz. If CW, set to zero and a more efficient CW solver will be used.
- **solver** (*str*, *optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

- **xflag** (*bool*, *optional*) – if intrinsic excitation field is calculated. The default is True.
- **mmflag** (*bool*, *optional*) – if intrinsic emission field is calculated. The default is True.
- **fflag** (*bool*, *optional*) – if fluorescence field is calculated. If set True, xflag must also be True. The default is True.

#### Returns

- **data** ([nirfasterff.base.FLdata](#)) – fluence and boundary measurables given the mesh and optodes.

See [FLdata\(\)](#) for details.

- **infox** ([nirfasterff.utils.ConvergenceInfo](#)) – convergence information of the solver, for intrinsic excitation.

See [ConvergenceInfo\(\)](#) for details

- **infom** ([nirfasterff.utils.ConvergenceInfo](#)) – convergence information of the solver, for intrinsic emission.

See [ConvergenceInfo\(\)](#) for details

- **infofl** ([nirfasterff.utils.ConvergenceInfo](#)) – convergence information of the solver, for fluorescence emission.

See [ConvergenceInfo\(\)](#) for details

**femdata\_moments**(*max\_moments*=3, *savefield*=*False*, *solver*=*utils.get\_solver()*, *opt*=*utils.SolverOptions()*)

Calculates TR moments at each location in the mesh for each source directly using Mellin transform, and then the boudary measurables for each channel

This is done for both excitation and fluorescence emission

If *mesh.vol* is set and *savefield* is set to *True*, internal moments data will be represented in volumetric space

See [femdata\\_fl\\_TR\\_moments\(\)](#) for details

#### Parameters

- **max\_moments** (*int32*, *optional*) – max order of moments to calculate. That is, 0th, 1st, 2nd, ..., max\_moments-th will be calculated. The default is 3.



- **savefield**(*bool*, *optional*) – If True, the internal moments are also returned. If False, only boundary moments are returned and `data.phix` and `data.phifl` will be empty.

The default is False.

- **solver**(*str*, *optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt**(`nirfasterff.utils.SolverOptions`, *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See `SolverOptions()` for details

### Returns

- **data**(`nirfasterff.base.flTRMomentsdata`) – internal and boundary moments given the mesh and optodes, both excitation and fluorescence emission.

See `flTRMomentsdata()` for details.

- **infox**(`nirfasterff.utils.ConvergenceInfo`) – convergence information of the solver, excitation.

Only the convergence info of highest order moments is returned.

See `ConvergenceInfo()` for details

- **infom**(`nirfasterff.utils.ConvergenceInfo`) – convergence information of the solver, fluorescence emission.

Only the convergence info of highest order moments is returned.

See `ConvergenceInfo()` for details

**femdata\_tpsf**(*tmax*, *dt*, *savefield=False*, *beautify=True*, *solver=utils.get\_solver()*, *opt=utils.SolverOptions()*)

Calculates TPSF at each location in the mesh for each source using a FEM solver, and then the boundary measurables for each channel

This is done for both excitation and fluorescence emission

If *mesh.vol* is set and *savefield* is set to *True*, internal TPSF data will be represented in volumetric space

See `femdata_fl_TR()` for details

### Parameters

- **tmax**(*double*) – maximum time simulated, in seconds.
- **dt**(*double*) – size of each time step, in seconds.
- **savefield**(*bool*, *optional*) – If True, the internal TPSFs are also returned. If False, only boundary TPSFs are returned and `data.phix` and `data.phifl` will be empty.

The default is False.

- **beautify**(*bool*, *optional*) – If true, zeros the initial unstable parts of the boundary TPSFs. The default is True.
- **solver**(*str*, *optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt**(`nirfasterff.utils.SolverOptions`, *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See `SolverOptions()` for details

**Returns**

- **data** (*nirfasterff.base.flTPSFdata*) – internal and boundary TPSFs given the mesh and op-todes, both excitation and fluorescence emission.

See [flTPSFdata\(\)](#) for details.

- **infox** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver, excitation.

Only the convergence info of the last time step is returned.

See [ConvergenceInfo\(\)](#) for details

- **infom** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver, fluorescence emission.

Only the convergence info of the last time step is returned.

See [ConvergenceInfo\(\)](#) for details

**from\_copy(mesh)**

Deep copy all fields from another mesh.

**Parameters**

**mesh** (*nirfasterff.base.fluormesh*) – the mesh to copy from.

**Return type**

None.

**from\_file(file)**

Read from classic NIRFAST mesh (ASCII) format, not checking the correctness of the loaded integration functions.

All fields after loading should be directly compatible with Matlab version.

**Parameters**

**file** (*str*) – name of the mesh. Any extension will be ignored.

**Return type**

None.

**Examples**

```
>>> mesh = nirfasterff.base.fluormesh()
>>> mesh.from_file('meshname')
```

**from\_mat(matfile, varname=None)**

Read from Matlab .mat file that contains a NIRFASTer mesh struct. All fields copied as is without error checking.

**Parameters**

- **matfile** (*str*) – name of the .mat file to load. Use of extension is optional.
- **varname** (*str, optional*) – if your .mat file contains multiple variables, use this argument to specify which one to load. The default is None.

When *varname==None*, *matfile* should contain exactly one structure, which is a NIRFASTer mesh, or the function will do nothing

**Return type**

None.

**from\_solid**(*ele, nodes, prop=None, src=None, det=None, link=None*)

Construct a NIRFASTer mesh from a 3D solid mesh generated by a mesher. Similar to the `solidmesh2nirfast` function in Matlab version.

Can also set the optical properties and optodes if supplied

**Parameters**

- **ele** (*int/double NumPy array*) – element list in one-based indexing. If four columns, all nodes will be labeled as region 1  
If five columns, the last column will be used for region labeling.
- **nodes** (*double NumPy array*) – node locations in the mesh. Unit: mm. Size (NNodes,3).
- **prop** (*double NumPy array, optional*) – If not *None*, calls `fluormesh.set_prop()` and sets the optical properties in the mesh. The default is *None*.

See `set_prop()` for details.

- **src** (*nirfasterff.base.optode, optional*) – If not *None*, sets the sources and moves them to the appropriate locations. The default is *None*.

See `touch_sources()` for details.

- **det** (*nirfasterff.base.optode, optional*) – If not *None*, sets the detectors and moves them to the appropriate locations. The default is *None*.

See `touch_detectors()` for details.

- **link** (*int32 NumPy array, optional*) – If not *None*, sets the channel information. Uses one-based indexing. The default is *None*.

Each row represents a channel, in the form of `[src, det, active]`, where *active* is 0 or 1

If *link* contains only two columns, all channels are considered active.

**Return type**

None.

**from\_volume**(*vol, param=utils.MeshingParams(), prop=None, src=None, det=None, link=None*)

Construct mesh from a segmented 3D volume using the built-in CGAL mesher. Calls `fluormesh.from_solid` after meshing step.

**Parameters**

- **vol** (*uint8 NumPy array*) – 3D segmented volume to be meshed. 0 is considered as outside. Regions labeled using unique integers.
- **param** (*nirfasterff.utils.MeshingParams, optional*) – parameters used in the CGAL mesher. If not specified, uses the default parameters defined in `nirfasterff.utils.MeshingParams()`.

Please modify fields `xPixelSpacing`, `yPixelSpacing`, and `SliceThickness` if your volume doesn't have `[1,1,1]` resolution

See `MesgingParams()` for details.

- **prop** (*double NumPy array, optional*) – If not *None*, calls `fluormesh.set_prop()` and sets the optical properties in the mesh. The default is *None*.

See `set_prop()` for details.

- **src** (*nirfasterff.base.optode*, *optional*) – If not *None*, sets the sources and moves them to the appropriate locations. The default is *None*.

See [touch\\_sources\(\)](#) for details.

- **det** (*nirfasterff.base.optode*, *optional*) – If not *None*, sets the detectors and moves them to the appropriate locations. The default is *None*.

See [touch\\_detectors\(\)](#) for details.

- **link** (*int32 NumPy array*, *optional*) – If not *None*, sets the channel information. Uses one-based indexing. The default is *None*.

Each row represents a channel, in the form of [*src*, *det*, *active*], where *active* is 0 or 1

If *link* contains only two columns, all channels are considered active.

#### Return type

*None*.

**gen\_intmat** (*xgrid*, *ygrid*, *zgrid*=[*]*)

Calculate the information needed to convert data between mesh and volumetric space, specified by x, y, z (if 3D) grids.

All grids must be uniform. The results will from a *nirfasterff.base.meshvol* object stored in field *.vol*

If field *.vol* already exists, it will be calculated again, and a warning will be thrown

#### Parameters

- **xgrid** (*double NumPy array*) – x grid in mm.
- **ygrid** (*double NumPy array*) – y grid in mm.
- **zgrid** (*double NumPy array*, *optional*) – z grid in mm. Leave empty for 2D meshes. The default is [*]*.

#### Raises

**ValueError** – if grids not uniform, or *zgrid* empty for 3D mesh

#### Return type

*None*.

**isvol**()

Check if conversion matrices between mesh and volumetric spaces are calculated

#### Returns

True if attribute *.vol* is calculate, False if not.

#### Return type

bool

**jacobian** (*freq*=0, *normalize*=True, *solver*=*utils.get\_solver()*, *opt*=*utils.SolverOptions()*)

Calculates the Jacobian matrix

Returns the Jacobian, direct field data, and the adjoint data

Structure of the Jacobian is detailed in [jacobian\\_fl\\_CW\(\)](#) and [jacobian\\_fl\\_FD\(\)](#)

#### Parameters

- **freq** (*double*) – modulation frequency in Hz.
- **normalize** (*bool*, *optional*) – whether normalize the Jacobian to the amplitudes of boundary measurements at excitation wavelength ('Born ratio').

The default is True.

- **solver** (*str*, *optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

#### Raises

**ValueError** – if modulation frequency is negative.

#### Returns

- **J** (*double NumPy array*) – The Jacobian matrix. Size (NChannel, NVoxel\*2) or (NChannel, NVoxel)
- **data1** (*nirfasterff.base.FLdata*) – The calculated direct field. The same as directly calling `mesh.femdata(freq)`
- **data2** (*nirfasterff.base.FLdata*) – The calculated adjoint field. The same as calling `mesh.femdata(freq)` AFTER swapping the locations of sources and detectors

#### **save\_nirfast**(*filename*)

Save mesh in the classic NIRFASTer ASCII format, which is directly compatible with the Matlab version

##### Parameters

**filename** (*str*) – name of the file to be saved as. Should have no extensions.

##### Return type

None.

#### **set\_prop**(*prop*)

Set optical properties of the whole mesh, using information provided in prop.

##### Parameters

**prop** (*double NumPy array*) – optical property info, similar to the MCX format:

```
[region muax(mm-1) musxp(mm-1) ri muam(mm-1) musmp(mm-1) muaf(mm-1)
  eta tau]
[region muax(mm-1) musxp(mm-1) ri muam(mm-1) musmp(mm-1) muaf(mm-1)
  eta tau]
[...]
```

where ‘region’ is the region label, and they should match exactly with `unique(mesh.region)`. The order doesn’t matter.

##### Return type

None.

#### **touch\_optodes**()

Moves all optodes (if non fixed) and recalculate the integration functions (i.e. barycentric coordinates).

See [touch\\_sources\(\)](#) and [touch\\_detectors\(\)](#) for details

##### Return type

None.

## nirfasterff.base.optodes

Define the optode class, an instance of which can be either a source or a detector

### Classes

---

<code>optode([coord])</code>	Class for NIRFASTer optodes, which can be either a group of sources or a group of detectors.
------------------------------	--

---

## nirfasterff.base.optodes.optode

**class** nirfasterff.base.optodes.optode(coord=[])

Bases: object

Class for NIRFASTer optodes, which can be either a group of sources or a group of detectors.

Note: The field fwhm for sources in the Matlab version has been dropped.

### **fixed**

whether an optode is fixed.

If not, it will be moved to one scattering length inside the surface (source) or on the surface (detector).

Default: 0

### **Type**

bool like

### **num**

indexing of the optodes, starting from one (1,2,3,...)

### **Type**

double NumPy vector

### **coord**

each row is the location of an optode. Unit: mm

### **Type**

double NumPy array

### **int\_func**

First column is the index (one-based) of the element each optode is in.

The subsequent columns are the barycentric coordinates (i.e. integration function) in the corresponding elements. Size (N, dim+2).

### **Type**

double NumPy array

**\_\_init\_\_**(coord=[])

## Methods

<code>__init__([coord])</code>	
<code>move_detectors(mesh)</code>	Moves detector to the appropriate locations in the mesh.
<code>move_sources(mesh)</code>	Moves sources to the appropriate locations in the mesh.
<code>touch_detectors(mesh)</code>	Recalculate/fill in all other fields based on 'fixed' and 'coord'.
<code>touch_sources(mesh)</code>	Recalculate/fill in all other fields based on 'fixed' and 'coord'.

### `move_detectors(mesh)`

Moves detector to the appropriate locations in the mesh.

For each detector, first move it to the closest point on the surface of the mesh.

Integration functions are NOT calculated after moving, to be consistent with the Matlab version.

#### Parameters

**mesh** (*NIRFASTer mesh type*) – The mesh on which the detectors are installed. Can be a 'stndmesh', 'fluormesh', or 'dcsmesh', either 2D or 3D

#### Raises

**ValueError** – If mesh.elements does not have 3 or 4 columns, or mesh.dimension is not 2 or 3.

#### Return type

None.

### `move_sources(mesh)`

Moves sources to the appropriate locations in the mesh.

For each source, first move it to the closest point on the surface of the mesh, and then move inside by one scattering length along surface normal.

where scattering length is  $1/\mu'_s$  for stnd and dcs mesh, and  $1/\mu'_{sx}$  for fluor mesh

Integration functions are also calculated after moving.

#### Parameters

**mesh** (*NIRFASTer mesh type*) – The mesh on which the sources are installed. Can be a 'stndmesh', 'fluormesh', or 'dcsmesh', either 2D or 3D

#### Raises

- **TypeError** – If mesh type is not recognized.
- **ValueError** – If mesh.elements does not have 3 or 4 columns, or mesh.dimension is not 2 or 3.

#### Return type

None.

### `touch_detectors(mesh)`

Recalculate/fill in all other fields based on 'fixed' and 'coord'.

This is useful when a set of detectors are manually added and only the locations are specified.

For non-fixed detectors, function ‘move\_detectors’ is first called, and integration functions are calculated subsequently.

For fixed detectors, recalculates integration functions directly.

If no detector locations are specified, the function does nothing

**Parameters**

**mesh** (*NIRFASTer mesh type*) – The mesh on which the sources are installed. Can be a ‘stndmesh’, ‘fluormesh’, or ‘dcsmesh’, either 2D or 3D

**Return type**

None.

**touch\_sources**(*mesh*)

Recalculate/fill in all other fields based on ‘fixed’ and ‘coord’.

This is useful when a set of sources are manually added and only the locations are specified.

For non-fixed sources, function ‘move\_sources’ is called, otherwise recalculates integration functions directly

If no source locations are specified, the function does nothing

**Parameters**

**mesh** (*NIRFASTer mesh type*) – The mesh on which the sources are installed. Can be a ‘stndmesh’, ‘fluormesh’, or ‘dcsmesh’, either 2D or 3D

**Return type**

None.

## nirfasterff.base.stnd\_mesh

Define the standard mesh class

## Classes

---

*stndmesh*()

Main class for standard mesh.

---

## nirfasterff.base.stnd\_mesh.stndmesh

**class** nirfasterff.base.stnd\_mesh.stndmesh

Bases: object

Main class for standard mesh. The methods should cover most of the commonly-used functionalities

**name**

name of the mesh. Default: ‘EmptyMesh’

**Type**

str

**nodes**

locations of nodes in the mesh. Unit: mm. Size (NNodes, dim)

**Type**

double NumPy array



**bndvtx**

indicator of whether a node is at boundary (1) or internal (0). Size (NNodes,)

**Type**

double NumPy array

**type**

type of the mesh. It is always 'std'.

**Type**

str

**mua**

absorption coefficient ( $\text{mm}^{-1}$ ) at each node. Size (NNodes,)

**Type**

double NumPy array

**kappa**

diffusion coefficient (mm) at each node. Size (NNodes,). Defined as  $1/(3*(\text{mua} + \text{mus}))$

**Type**

double NumPy array

**ri**

refractive index at each node. Size (NNodes,)

**Type**

double NumPy array

**mus**

reduced scattering coefficient ( $\text{mm}^{-1}$ ) at each node. Size (NNodes,)

**Type**

(double NumPy array

**elements**

triangulation (tetrahedrons or triangles) of the mesh, Size (NElements, dim+1)

Row i contains the indices of the nodes that form tetrahedron/triangle i

One-based indexing for direct interoperability with the Matlab version

**Type**

double NumPy array

**region**

region labeling of each node. Starting from 1. Size (NNodes,)

**Type**

double NumPy array

**source**

information about the sources

**Type**

nirfasterff.base.optode

**meas**

information about the the detectors

**Type**

nirfasterff.base.optode

**link**

list of source-detector pairs, i.e. channels. Size (NChannels,3)

First column: source; Second column: detector; Third column: active (1) or not (0)

**Type**

int32 NumPy array

**c**light speed (mm/sec) at each node. Size (NNodes,). Defined as  $c_0/r_i$ , where  $c_0$  is the light speed in vacuum**Type**

double NumPy array

**ksi**

photon fluence rate scale factor on the mesh-outside\_mesh boundary as derived from Fresnel's law. Size (NNodes,)

**Type**

double NumPy array

**element\_area**volume/area ( $\text{mm}^3$  or  $\text{mm}^2$ ) of each element. Size (NElements,)**Type**

double NumPy array

**support**

total volume/area of all the elements each node belongs to. Size (NNodes,)

**Type**

double NumPy array

**vol**

object holding information for converting between mesh and volumetric space.

**Type**

nirfasterff.base.meshvol

**\_\_init\_\_()**

## Methods

<code>__init__()</code>	
<code>change_prop(idx, prop)</code>	Change optical properties (mua, musp, and ri) at nodes specified in idx, and automatically change fields kappa, c, and ksi as well
<code>femdata(freq[, solver, opt])</code>	Calculates fluences for each source using a FEM solver, and then the boudary measurables for each channel
<code>femdata_moments([max_moments, savefield, ...])</code>	Calculates TR moments at each location in the mesh for each source directly using Mellin transform, and then the boudary measurables for each channel
<code>femdata_tpsf(tmax, dt[, savefield, ...])</code>	Calculates TPSF at each location in the mesh for each source using a FEM solver, and then the boudary measurables for each channel
<code>from_copy(mesh)</code>	Deep copy all fields from another mesh.
<code>from_file(file)</code>	Read from classic NIRFAST mesh (ASCII) format, not checking the correctness of the loaded integration functions.
<code>from_mat(matfile[, varname])</code>	Read from Matlab .mat file that contains a NIRFASTer mesh struct.
<code>from_solid(ele, nodes[, prop, src, det, link])</code>	Construct a NIRFASTer mesh from a 3D solid mesh generated by a mesher.
<code>from_volume(vol[, param, prop, src, det, link])</code>	Construct mesh from a segmented 3D volume using the built-in CGAL mesher.
<code>gen_intmat(xgrid, ygrid[, zgrid])</code>	Calculate the information needed to convert data between mesh and volumetric space, specified by x, y, z (if 3D) grids.
<code>isvol()</code>	Check if conversion matrices between mesh and volumetric spaces are calculated
<code>jacobian([freq, normalize, mus, solver, opt])</code>	Calculates the Jacobian matrix
<code>save_nirfast(filename)</code>	Save mesh in the classic NIRFASTer ASCII format, which is directly compatible with the Matlab version
<code>set_prop(prop)</code>	Set optical properties of the whole mesh, using information provided in prop.
<code>touch_optodes()</code>	Moves all optodes (if non fixed) and recalculate the integration functions (i.e. barycentric coordinates).

### `change_prop(idx, prop)`

Change optical properties (mua, musp, and ri) at nodes specified in idx, and automatically change fields kappa, c, and ksi as well

#### Parameters

- **idx** (*list or NumPy array or -1*) – zero-based indices of nodes to change. If *idx==-1*, function changes all the nodes
- **prop** (*list or NumPy array of length 3*) – new optical properties to be assigned to the specified nodes. [mua(mm-1) musp(mm-1) ri].

#### Return type

None.

**femdata**(*freq*, *solver*=*utils.get\_solver()*, *opt*=*utils.SolverOptions()*)

Calculates fluences for each source using a FEM solver, and then the boudary measurables for each channel

If *mesh.vol* is set, fluence data will be represented in volumetric space

See [femdata\\_stnd\\_CW\(\)](#) and [femdata\\_stnd\\_FD\(\)](#) for details

#### Parameters

- **freq** (*double*) – modulation frequency in Hz. If CW, set to zero and a more efficient CW solver will be used.
- **solver** (*str*, *optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

#### Returns

- **data** ([nirfasterff.base.FDdata](#)) – fluence and boundary measurables given the mesh and optodes.

See [FDdata\(\)](#) for details.

- **info** ([nirfasterff.utils.ConvergenceInfo](#)) – convergence information of the solver.

See [ConvergenceInfo\(\)](#) for details

**femdata\_moments**(*max\_moments*=3, *savefield*=False, *solver*=*utils.get\_solver()*, *opt*=*utils.SolverOptions()*)

Calculates TR moments at each location in the mesh for each source directly using Mellin transform, and then the boudary measurables for each channel

If *mesh.vol* is set and *savefield* is set to *True*, internal moments data will be represented in volumetric space

See [femdata\\_stnd\\_TR\\_moments\(\)](#) for details

#### Parameters

- **max\_moments** (*int32*, *optional*) – max order of moments to calculate. That is, 0th, 1st, 2nd, ..., max\_moments-th will be calculated. The default is 3.
- **savefield** (*bool*, *optional*) – If True, the internal moments are also returned. If False, only boundary moments are returned and *data.phi* will be empty.

The default is False.

- **solver** (*str*, *optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

#### Returns

- **data** ([nirfasterff.base.TRMomentsdata](#)) – internal and boundary moments given the mesh and optodes.

See [TRMomentsdata\(\)](#) for details.

- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.

Only the convergence info of highest order moments is returned.

See [ConvergenceInfo\(\)](#) for details

**femdata\_tpsf**(*tmax*, *dt*, *savefield=False*, *beautify=True*, *solver=utils.get\_solver()*,  
*opt=utils.SolverOptions()*)

Calculates TPSF at each location in the mesh for each source using a FEM solver, and then the boundary measurables for each channel

If *mesh.vol* is set and *savefield* is set to *True*, internal TPSF data will be represented in volumetric space

See [femdata\\_std\\_TR\(\)](#) for details

#### Parameters

- **tmax** (*double*) – maximum time simulated, in seconds.
- **dt** (*double*) – size of each time step, in seconds.
- **savefield** (*bool*, *optional*) – If *True*, the internal TPSFs are also returned. If *False*, only boundary TPSFs are returned and *data.phi* will be empty.

The default is *False*.

- **beautify** (*bool*, *optional*) – If *true*, zeros the initial unstable parts of the boundary TPSFs. The default is *True*.
- **solver** (*str*, *optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

#### Returns

- **data** (*nirfasterff.base.TPSFdata*) – internal and boundary TPSFs given the mesh and optodes.

See [TPSFdata\(\)](#) for details.

- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.

Only the convergence info of the last time step is returned.

See [ConvergenceInfo\(\)](#) for details

**from\_copy**(*mesh*)

Deep copy all fields from another mesh.

#### Parameters

**mesh** (*nirfasterff.base.stndmesh*) – the mesh to copy from.

#### Return type

None.

**from\_file**(*file*)

Read from classic NIRFAST mesh (ASCII) format, not checking the correctness of the loaded integration functions.

All fields after loading should be directly compatible with Matlab version.

**Parameters**

**file** (*str*) – name of the mesh. Any extension will be ignored.

**Return type**

None.

**Examples**

```
>>> mesh = nirfasterff.base.stndmesh()
>>> mesh.from_file('meshname')
```

**from\_mat** (*matfile*, *varname=None*)

Read from Matlab .mat file that contains a NIRFASTer mesh struct. All fields copied as is without error checking.

**Parameters**

- **matfile** (*str*) – name of the .mat file to load. Use of extension is optional.
- **varname** (*str*, *optional*) – if your .mat file contains multiple variables, use this argument to specify which one to load. The default is None.

When *varname==None*, *matfile* should contain exactly one structure, which is a NIRFASTer mesh, or the function will do nothing

**Return type**

None.

**from\_solid** (*ele*, *nodes*, *prop=None*, *src=None*, *det=None*, *link=None*)

Construct a NIRFASTer mesh from a 3D solid mesh generated by a mesher. Similar to the `solidmesh2nirfast` function in Matlab version.

Can also set the optical properties and optodes if supplied

**Parameters**

- **ele** (*int/double NumPy array*) – element list in one-based indexing. If four columns, all nodes will be labeled as region 1

If five columns, the last column will be used for region labeling.

- **nodes** (*double NumPy array*) – node locations in the mesh. Unit: mm. Size (Nnodes,3).

- **prop** (*double NumPy array*, *optional*) – If not *None*, calls `stndmesh.set_prop()` and sets the optical properties in the mesh. The default is None.

See [set\\_prop\(\)](#) for details.

- **src** (*nirfasterff.base.optode*, *optional*) – If not *None*, sets the sources and moves them to the appropriate locations. The default is None.

See [touch\\_sources\(\)](#) for details.

- **det** (*nirfasterff.base.optode*, *optional*) – If not *None*, sets the detectors and moves them to the appropriate locations. The default is None.

See [touch\\_detectors\(\)](#) for details.

- **link** (*int32 NumPy array*, *optional*) – If not *None*, sets the channel information. Uses one-based indexing. The default is None.

Each row represents a channel, in the form of *[src, det, active]*, where *active* is 0 or 1

If *link* contains only two columns, all channels are considered active.

#### Return type

None.

**from\_volume**(*vol*, *param*=*utils.MeshingParams()*, *prop*=*None*, *src*=*None*, *det*=*None*, *link*=*None*)

Construct mesh from a segmented 3D volume using the built-in CGAL mesher. Calls *stndmesh.from\_solid* after meshing step.

#### Parameters

- **vol** (*uint8 NumPy array*) – 3D segmented volume to be meshed. 0 is considered as outside. Regions labeled using unique integers.
- **param** (*nirfasterff.utils.MeshingParams*, *optional*) – parameters used in the CGAL mesher. If not specified, uses the default parameters defined in *nirfasterff.utils.MeshingParams()*.

Please modify fields *xPixelSpacing*, *yPixelSpacing*, and *SliceThickness* if your volume doesn't have [1,1,1] resolution

See *MeshingParams()* for details.

- **prop** (*double NumPy array*, *optional*) – If not *None*, calls *stndmesh.set\_prop()* and sets the optical properties in the mesh. The default is *None*.

See *set\_prop()* for details.

- **src** (*nirfasterff.base.optode*, *optional*) – If not *None*, sets the sources and moves them to the appropriate locations. The default is *None*.

See *touch\_sources()* for details.

- **det** (*nirfasterff.base.optode*, *optional*) – If not *None*, sets the detectors and moves them to the appropriate locations. The default is *None*.

See *touch\_detectors()* for details.

- **link** (*int32 NumPy array*, *optional*) – If not *None*, sets the channel information. Uses one-based indexing. The default is *None*.

Each row represents a channel, in the form of *[src, det, active]*, where *active* is 0 or 1

If *link* contains only two columns, all channels are considered active.

#### Return type

None.

**gen\_intmat**(*xgrid*, *ygrid*, *zgrid*=[*1*])

Calculate the information needed to convert data between mesh and volumetric space, specified by x, y, z (if 3D) grids.

All grids must be uniform. The results will from a *nirfasterff.base.meshvol* object stored in field *.vol*

If field *.vol* already exists, it will be calculated again, and a warning will be thrown

#### Parameters

- **xgrid** (*double NumPy array*) – x grid in mm.
- **ygrid** (*double NumPy array*) – y grid in mm.
- **zgrid** (*double NumPy array*, *optional*) – z grid in mm. Leave empty for 2D meshes. The default is [*1*].

**Raises**

**ValueError** – if grids not uniform, or zgrid empty for 3D mesh

**Return type**

None.

**isvol()**

Check if conversion matrices between mesh and volumetric spaces are calculated

**Returns**

True if attribute `.vol` is calculate, False if not.

**Return type**

bool

**jacobian**(*freq=0, normalize=True, mus=False, solver=utils.get\_solver(), opt=utils.SolverOptions()*)

Calculates the Jacobian matrix

Returns the Jacobian, direct field data, and the adjoint data

Structure of the Jacobian is detailed in [jacobian\\_stnd\\_CW\(\)](#) and [jacobian\\_stnd\\_FD\(\)](#)

**Parameters**

- **freq** (*double*) – modulation frequency, in Hz.
- **normalize** (*bool, optional*) – whether normalize the Jacobian to the amplitudes of boundary measurements, i.e. use Rytov approximation.  
The default is True.
- **mus** (*bool, optional*) – whether derivates wrt mus (left half of the ‘full’ Jacobian) is calculated. Only has effect when `freq=0`. The default is True.
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

**Raises**

**ValueError** – if `freq` is negative.

**Returns**

- **J** (*double or complex double NumPy array*) – The Jacobian matrix. Size (NChannel\*2, NVoxel\*2) if `freq>0`, (NChannel, NVoxel\*2) if `freq=0` and `mus=True`, (NChannel, NVoxel) if `freq=0` and `mus=False`
- **data1** ([nirfasterff.base.FDdata](#)) – The calculated direct field. The same as directly calling `mesh.femdata(freq)`
- **data2** ([nirfasterff.base.FDdata](#)) – The calculated adjoint field. The same as calling `mesh.femdata(freq)` AFTER swapping the locations of sources and detectors

**save\_nirfast**(*filename*)

Save mesh in the classic NIRFASTer ASCII format, which is directly compatible with the Matlab version

**Parameters**

**filename** (*str*) – name of the file to be saved as. Should have no extensions.

**Return type**

None.



**set\_prop(prop)**

Set optical properties of the whole mesh, using information provided in prop.

**Parameters**

**prop** (*double NumPy array*) – optical property info, similar to the MCX format:

```
[region mua(mm-1) musp(mm-1) ri]
[region mua(mm-1) musp(mm-1) ri]
[...]
```

where ‘region’ is the region label, and they should match exactly with `unique(mesh.region)`.  
The order doesn’t matter.

**Return type**

None.

**touch\_optodes()**

Moves all optodes (if non fixed) and recalculate the integration functions (i.e. barycentric coordinates).

See [touch\\_sources\(\)](#) and [touch\\_detectors\(\)](#) for details

**Return type**

None.

## 4.1.2 nirfasterff.forward

Functions for forward data calculation

### Modules

<code>nirfasterff.forward.analytical</code>	Analytical solutions to the diffusion equation in semi-infinite media
<code>nirfasterff.forward.femdata</code>	The FEM solvers.

### nirfasterff.forward.analytical

Analytical solutions to the diffusion equation in semi-infinite media

### Functions

<code>semi_infinite_CW(mua, musp, n, rho[, z, ...])</code>	Calculates the continuous-wave fluence in space using the analytical solution to the diffusion equation in semi-infinite media.
<code>semi_infinite_DCS(mua, musp, n, rho, aDb, ...)</code>	Calculates DCS G1 curve using the analytical solution to the correlation diffusion equation in semi-infinite media
<code>semi_infinite_FD(mua, musp, n, freq, rho[, ...])</code>	Calculates the frequency-domain fluence in space using the analytical solution to the diffusion equation in semi-infinite media.
<code>semi_infinite_TR(mua, musp, n, rho, T, dt[, ...])</code>	Calculates TPSF at a given location using the analytical solution to the diffusion equation in semi-infinite media

### nirfasterff.forward.analytical.semi\_infinite\_CW

`nirfasterff.forward.analytical.semi_infinite_CW(mua, musp, n, rho, z=0, boundary='exact', n_air=1.0)`

Calculates the continuous-wave fluence in space using the analytical solution to the diffusion equation in semi-infinite media.

Internally calls the FD version with freq set to zero

#### Parameters

- **mua** (*double*) – absorption coefficient of the medium, in  $\text{mm}^{-1}$ .
- **musp** (*double*) – reduced scattering coefficient of the medium, in  $\text{mm}^{-1}$ .
- **n** (*double*) – refractive index of the medium.
- **rho** (*double NumPy vector or scalar*) – distance to the light source, projected to the x-y (i.e. boundary plane of the semi-infinite space) plane, in mm.  
Can be a vector, in which case fluences calculated at multiple locations will be returned
- **z** (*double NumPy vector or scalar, optional*) – depth of the location(s) of interest. 0 for boundary measurement.  
If a vector, it must have the same length as rho  
The default is 0.
- **boundary** (*str, optional*) – type of the boundary condition, which can be 'robin', 'approx', or 'exact'. The default is 'exact'.  
See [boundary\\_attenuation\(\)](#) for details.
- **n\_air** (*double, optional*) – refractive index outside of the semi-infinite space, which is typically assumed to be air. The default is 1.0.

#### Returns

calculated fluence, where each element corresponds to a location, as specified by rho and z. Size (NLocation,)

#### Return type

double NumPy vector

### References

Durduran et al, 2010, Rep. Prog. Phys. doi:10.1088/0034-4885/73/7/076701

### nirfasterff.forward.analytical.semi\_infinite\_DCS

`nirfasterff.forward.analytical.semi_infinite_DCS(mua, musp, n, rho, aDb, wlength, tvec, z=0, boundary='exact', n_air=1.0, normalize=0)`

Calculates DCS G1 curve using the analytical solution to the correlation diffusion equation in semi-infinite media

Function assumes Brownian motion, that is,  $\langle \Delta r^2 \rangle = 6\alpha D b \tau$

#### Parameters

- **mua** (*double*) – absorption coefficient of the medium, in  $\text{mm}^{-1}$ .
- **musp** (*double*) – reduced scattering coefficient of the medium, in  $\text{mm}^{-1}$ .

- **n** (*double*) – refractive index of the medium.
- **rho** (*scalar*) – distance to the light source, projected to the x-y (i.e. boundary plane of the semi-infinite space) plane, in mm.
- **aDb** (*double*) – The lumped flow parameter  $\alpha Db$  in Brownian motion.
- **wavelength** (*double or int*) – wavelength used, in nm.
- **tvec** (*double Numpy vector*) – time vector used to calculate the G1 curve. It is usually a good idea to use log space.
- **z** (*double, optional*) – depth of the location of interest. 0 for boundary measurement. The default is 0.
- **boundary** (*str, optional*) – type of the boundary condition, which can be ‘robin’, ‘approx’, or ‘exact’. The default is ‘exact’.

See [boundary\\_attenuation\(\)](#) for details.

- **n\_air** (*double, optional*) – refractive index outside of the semi-infinite space, which is typically assumed to be air. The default is 1.0.
- **normalize** (*bool, optional*) – if true, returns the normalized g1 curve, instead of G1. The default is 0.

#### Returns

G1 or g1 curve calculated at the given location and time points.

#### Return type

double NumPy vector

## References

Durduran et al, 2010, Rep. Prog. Phys. doi:10.1088/0034-4885/73/7/076701

### nirfasterff.forward.analytical.semi\_infinite\_FD

`nirfasterff.forward.analytical.semi_infinite_FD(mua, musp, n, freq, rho, z=0, boundary='exact', n_air=1.0)`

Calculates the frequency-domain fluence in space using the analytical solution to the diffusion equation in semi-infinite media.

#### Parameters

- **mua** (*double*) – absorption coefficient of the medium, in  $\text{mm}^{-1}$ .
- **musp** (*double*) – reduced scattering coefficient of the medium, in  $\text{mm}^{-1}$ .
- **n** (*double*) – refractive index of the medium.
- **freq** (*double NumPy vector or scalar*) – modulation frequency, in Hz.
- **rho** (*double NumPy vector or scalar*) – distance to the light source, projected to the x-y (i.e. boundary plane of the semi-infinite space) plane, in mm.

Can be a vector, in which case fluences calculated at multiple locations will be returned

- **z** (*double NumPy vector or scalar, optional*) – depth of the location(s) of interest. 0 for boundary measurement.

If a vector, it must have the same length as rho

The default is 0.

- **boundary** (*str*, *optional*) – type of the boundary condition, which can be ‘robin’, ‘approx’, or ‘exact’. The default is ‘exact’.

See [boundary\\_attenuation\(\)](#) for details.

- **n\_air** (*double*, *optional*) – refractive index outside of the semi-infinite space, which is typically assumed to be air. The default is 1.0.

### Returns

calculated complex fluence, where each row (or element in vector) corresponds to a location, as specified by rho and z,

and each column corresponds to a modulation frequency. Size (NLocation,), or (NLocation, NFreq)

### Return type

complex double NumPy array

### References

Durduran et al, 2010, Rep. Prog. Phys. doi:10.1088/0034-4885/73/7/076701

### nirfasterff.forward.analytical.semi\_infinite\_TR

`nirfasterff.forward.analytical.semi_infinite_TR(mua, musp, n, rho, T, dt, z=0, boundary='EBC-Robin')`

Calculates TPSF at a given location using the analytical solution to the diffusion equation in semi-infinite media

### Parameters

- **mua** (*double*) – absorption coefficient of the medium, in  $\text{mm}^{-1}$ .
- **musp** (*double*) – reduced scattering coefficient of the medium, in  $\text{mm}^{-1}$ .
- **n** (*double*) – refractive index of the medium.
- **rho** (*scalar*) – distance to the light source, projected to the x-y (i.e. boundary plane of the semi-infinite space) plane, in mm.
- **T** (*double NumPy vector or scalar*) – if a scalar, it is the total amount of time and time vector will be generated also based on dt (see below).

if a vector, it is directly used as the time vector, and the argument dt will be ignored. Unit: seconds

- **dt** (*double*) – step size of the time vector (in seconds). If the argument T is a vector, it will be ignored.
- **z** (*double, optional*) – depth of the location of interest. 0 for boundary measurement. The default is 0.

Note that when  $z=0$ , the function returns reflectance, instead of fluence. Please refer to the References for detail.

- **boundary** (*str, optional*) – type of the boundary condition, which can be (case insensitive),

‘PCB-exact’ - partial current boundary condition, with exact internal reflectance

'PCB-approx' - partial current boundary condition, with Groenhuis internal reflectance approximation

'PCB-Robin' - partial current boundary condition, with internal reflectance derived from Fresnel's law

'EBC-exact' - extrapolated boundary condition, with exact internal reflectance

'EBC-approx' - extrapolated boundary condition, with Groenhuis internal reflectance approximation

'EBC-Robin' - extrapolated boundary condition, with internal reflectance derived from Fresnel's law

'ZBC' - zero boundary condition

The default is 'EBC-Robin'.

See [boundary\\_attenuation\(\)](#) for the differences between 'exact', 'approx' and 'robin'

#### Raises

**ValueError** – if boundary condition is not of a recognized kind.

#### Returns

**phi** – Column 0: the time vector; Column 1: calculated TPSF at the given location. Size (NTime, 2)

#### Return type

double Numpy array

#### References

Hielscher et al., 1995, Phys. Med. Biol. doi:10.1088/0031-9155/40/11/013

Kienle and Patterson, 1997, JOSA A. doi:10.1364/JOSAA.14.000246

#### nirfasterff.forward.femdata

The FEM solvers. It is usually recommended to use the high-level wrappers in the mesh classes by calling the mesh.femdata\* functions.

## Functions

<code>femdata_DCS(mesh, tvec[, solver, opt])</code>	Forward modeling calculating steady-state fluences and G1/g1 curves by solving the correlation diffusion equation.
<code>femdata_fl_CW(mesh[, solver, opt, xflag, ...])</code>	Forward modeling for CW in fluorescence meshes.
<code>femdata_fl_FD(mesh, freq[, solver, opt, ...])</code>	Forward modeling for FD in fluorescence meshes.
<code>femdata_fl_TR(mesh, tmax, dt[, savefield, ...])</code>	Forward modeling calculating TPSF on a fluorescence mesh.
<code>femdata_fl_TR_moments(mesh[, max_moments, ...])</code>	Forward modeling calculating TR moments using Mellin transform on a fluorescence mesh.
<code>femdata_std_CW(mesh[, solver, opt])</code>	Forward modeling for CW.
<code>femdata_std_FD(mesh, freq[, solver, opt])</code>	Forward modeling for FD.
<code>femdata_std_TR(mesh, tmax, dt[, savefield, ...])</code>	Forward modeling calculating TPSF on a standard mesh.
<code>femdata_std_TR_moments(mesh[, max_moments, ...])</code>	Forward modeling calculating TR moments using Mellin transform on a standard mesh.

### nirfasterff.forward.femdata.femdata\_DCS

`nirfasterff.forward.femdata.femdata_DCS(mesh, tvec, solver=utils.get_solver(), opt=utils.SolverOptions())`

Forward modeling calculating steady-state fluences and G1/g1 curves by solving the correlation diffusion equation. Please consider using `mesh.femdata(tvec)` instead.

The fluences as well as its boundary amplitudes are exactly the same as what the CW solver would give when  $\tau=0$ .

g1 curve is simple G1 curve normalized by the boundary amplitudes.

The function calculates the MASS matrices, the source vectors, and calls the G1 solver, which internally utilizes the CW solver.

When calculating the flow-related term, the function assumes Brownian motion and uses only `mesh.aDb` (that is, `mesh.a` and `mesh.Db` are ignored).

This is to say, we assume  $\langle \Delta r^2 \rangle = 6\alpha Db\tau$

#### Parameters

- **mesh** (`nirfasterff.base.dcsmesh`) – the mesh used to calculate the forward data.
- **tvec** (`double NumPy array`) – time vector (i.e. :math:`\tau` au) for the G1 curve, in seconds. It is usually a good idea to use log scale
- **solver** (`str, optional`) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (`nirfasterff.utils.SolverOptions, optional`) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See `SolverOptions()` for details

#### Raises

**TypeError** – if mesh is not a dcs mesh.

#### Returns

- **data** (`nirfasterff.base.DCSdata`) – contains fluence, G1 curve, and g1 curve calculated at each spatial location, and also the boundary data.

If `mesh.vol` is set, internal G1 curves will be returned in volumetric space

See [DCSdata\(\)](#) for details.

- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver when calculating the fluence field.

See [ConvergenceInfo\(\)](#) for details

See also:

[gen\\_mass\\_matrix\(\)](#), [gen\\_sources\(\)](#), and [get\\_field\\_CW\(\)](#)

## References

Durduran et al, 2010, Rep. Prog. Phys. doi:10.1088/0034-4885/73/7/076701

### nirfasterff.forward.femdata.femdata\_fl\_CW

`nirfasterff.forward.femdata.femdata_fl_CW(mesh, solver=utils.get_solver(), opt=utils.SolverOptions(), xflag=True, mmflag=True, flflag=True)`

Forward modeling for CW in fluorescence meshes. Please consider using `mesh.femdata(0)` instead.

The function calculates the MASS matrices, the source vectors, and calls the CW solver (preconditioned conjugated gradient).

The optional flags can be used to determine which fields are calculated. By default all true.

Note that when `flflag` is set to True, `xflag` must also be True.

#### Parameters

- **mesh** (*nirfasterff.base.fluormesh*) – the mesh used to calculate the forward data.
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

- **xflag** (*bool, optional*) – if intrinsic excitation field is calculated. The default is True.
- **mmflag** (*bool, optional*) – if intrinsic emission field is calculated. The default is True.
- **flflag** (*bool, optional*) – if fluorescence field is calculated. If set True, `xflag` must also be True. The default is True.

#### Raises

- **TypeError** – If mesh is not a fluor mesh.
- **ValueError** – If `flflag` is set True but `xflag` is not.

#### Returns

- **data** (*nirfasterff.base.FLdata*) – fluence and boundary measurables given the mesh and optodes.

If `mesh.vol` is defined, the returned fluences will be in volumetric space

See [FLdata\(\)](#) for details.

- **infox** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver, for intrinsic excitation.

See [ConvergenceInfo\(\)](#) for details

- **infom** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver, for intrinsic emission.

See [ConvergenceInfo\(\)](#) for details

- **infofl** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver, for fluorescence emission.

See [ConvergenceInfo\(\)](#) for details

See also:

[get\\_field\\_CW\(\)](#), [gen\\_mass\\_matrix\(\)](#), [gen\\_sources\(\)](#), and [gen\\_sources\\_fl\(\)](#)

### **nirfasterff.forward.femdata.femdata\_fl\_FD**

```
nirfasterff.forward.femdata.femdata_fl_FD(mesh, freq, solver=utils.get_solver(),
                                           opt=utils.SolverOptions(), xflag=True, mmflag=True,
                                           flflag=True)
```

Forward modeling for FD in fluorescence meshes. Please consider using `mesh.femdata(frequency)` instead.

The function calculates the MASS matrix, the source vectors, and calls the FD solver (preconditioned BiCGStab).

The optional flags can be used to determine which fields are calculated. By default all true.

Note that when `flflag` is set to `True`, `xflag` must also be `True`.

#### **Parameters**

- **mesh** (*nirfasterff.base.stndmesh*) – the mesh used to calculate the forward data.
- **freq** (*double*) – modulation frequency in Hz.  
When it is 0, function continues with the BiCGstab solver, but generates a warning that the CW solver should be used for better performance
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.  
See [SolverOptions\(\)](#) for details
- **xflag** (*bool, optional*) – if intrinsic excitation field is calculated. The default is `True`.
- **mmflag** (*bool, optional*) – if intrinsic emission field is calculated. The default is `True`.
- **flflag** (*bool, optional*) – if fluorescence field is calculated. If set `True`, `xflag` must also be `True`. The default is `True`.

#### **Raises**

- **TypeError** – If mesh is not a fluor mesh.
- **ValueError** – If `flflag` is set `True` but `xflag` is not.

#### **Returns**



- **data** (*nirfasterff.base.FLdata*) – fluence and boundary measurables given the mesh and optodes.

If mesh.vol is defined, the returned fluences will be in volumetric space

See [FLdata\(\)](#) for details.

- **infox** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver, for intrinsic excitation.

See [ConvergenceInfo\(\)](#) for details

- **infom** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver, for intrinsic emission.

See [ConvergenceInfo\(\)](#) for details

- **infofl** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver, for fluorescence emission.

See [ConvergenceInfo\(\)](#) for details

See also:

[get\\_field\\_FD\(\)](#), [gen\\_mass\\_matrix\(\)](#), [gen\\_sources\(\)](#), and [gen\\_sources\\_fl\(\)](#)

## nirfasterff.forward.femdata.femdata\_fl\_TR

`nirfasterff.forward.femdata.femdata_fl_TR(mesh, tmax, dt, savefield=False, beautify=True, solver=utils.get_solver(), opt=utils.SolverOptions())`

Forward modeling calculating TPSF on a fluorescence mesh. Please consider using `mesh.femdata_tpsf(tmax, dt)` instead.

The function calculates the MASS matrices, the source vectors, and calls two separate TPSF solvers (both pre-conditioned conjugated gradient):

First time calculates the TPSF for the excitation field, the result of which is consequently convolved with the decay.

The second solver is called with the convolved excitation field as its input to calculate the TPSF for fluorescence emission.

### Parameters

- **mesh** (*nirfasterff.base.fluormesh*) – the mesh used to calculate the forward data.
- **tmax** (*double*) – maximum time simulated, in seconds.
- **dt** (*double*) – size of each time step, in seconds.
- **savefield** (*bool, optional*) – If True, the internal TPSFs are also returned. If False, only boundary TPSFs are returned and `data.phix` and `data.phifl` will be empty.

The default is False.

- **beautify** (*bool, optional*) – If true, zeros the initial unstable parts of the boundary TPSFs. The default is True.
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

**Raises**

**TypeError** – if mesh is not a fluor mesh.

**Returns**

- **data** (*nirfasterff.base.flTPSFdata*) – internal and boundary TPSFs given the mesh and optodes, both excitation and fluorescence emission.

If *mesh.vol* is set and *savefield* is set to *True*, internal TPSF data will be represented in volumetric space

See [flTPSFdata\(\)](#) for details.

- **infox** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver, excitation.

Only the convergence info of the last time step is returned.

See [ConvergenceInfo\(\)](#) for details

- **infom** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver, fluorescence emission.

Only the convergence info of the last time step is returned.

See [ConvergenceInfo\(\)](#) for details

See also:

[gen\\_mass\\_matrix\(\)](#), [get\\_field\\_TR\(\)](#), and [get\\_field\\_TRFL\(\)](#)

**nirfasterff.forward.femdata.femdata\_fl\_TR\_moments**

```
nirfasterff.forward.femdata.femdata_fl_TR_moments(mesh, max_moments=3, savefield=False,
                                                    solver=utils.get_solver(),
                                                    opt=utils.SolverOptions())
```

Forward modeling calculating TR moments using Mellin transform on a fluorescence mesh. Please consider using `mesh.femdata_moments()` instead.

The function calculates the MASS matrix, the source vectors, and calls the two Mellin moments solver (both preconditioned conjugated gradient):

First time calculates the moments for the excitation field, the result of which is consequently used as the input of the second solver,

which calculates the moments of fluorescence emission based on the excitation moments

Calculates 0th, 1st, 2nd, ..., *max\_moments*-th moments directly without calculating TPSF first, and this is done for both excitation and fluorescence emission.

This is more efficient, if the time series are not of concern.

**Parameters**

- **mesh** (*nirfasterff.base.fluormesh*) – the mesh used to calculate the forward data.
- **max\_moments** (*int32, optional*) – max order of moments to calculate. That is, 0th, 1st, 2nd, ..., *max\_moments*-th will be calculated. The default is 3.
- **savefield** (*bool, optional*) – If True, the internal moments are also returned. If False, only boundary moments are returned and `data.phix` and `data.phifl` will be empty.

The default is False.

- **solver** (*str*, *optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

#### Raises

- **TypeError** – if mesh is not a fluor mesh.
- **ValueError** – if max\_moments is negative.

#### Returns

- **data** ([nirfasterff.base.flTRMomentsdata](#)) – internal and boundary moments given the mesh and optodes, both excitation and fluorescence emission.

If *mesh.vol* is set and *savefield* is set to *True*, internal moments will be represented in volumetric space

See [flTRMomentsdata\(\)](#) for details.

- **infox** ([nirfasterff.utils.ConvergenceInfo](#)) – convergence information of the solver, excitation.

Only the convergence info of highest order moments is returned.

See [ConvergenceInfo\(\)](#) for details

- **infom** ([nirfasterff.utils.ConvergenceInfo](#)) – convergence information of the solver, fluorescence emission.

Only the convergence info of highest order moments is returned.

See [ConvergenceInfo\(\)](#) for details

See also:

[gen\\_mass\\_matrix\(\)](#), [get\\_field\\_TRmoments\(\)](#), and [get\\_field\\_TRFLmoments\(\)](#)

### nirfasterff.forward.femdata.femdata\_stnd\_CW

`nirfasterff.forward.femdata.femdata_stnd_CW(mesh, solver=utils.get_solver(), opt=utils.SolverOptions())`

Forward modeling for CW. Please consider using `mesh.femdata(0)` instead.

The function calculates the FEM MASS matrix, the source vectors, and calls the CW solver (preconditioned conjugated gradient).

#### Parameters

- **mesh** ([nirfasterff.base.stndmesh](#)) – the mesh used to calculate the forward data.
- **solver** (*str*, *optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

#### Raises

**TypeError** – If mesh is not a stnd mesh.

#### Returns

- **data** (*nirfasterff.base.FDdata*) – fluence and boundary measurables given the mesh and optodes.

If mesh.vol is defined, the returned fluence will be in volumetric space

See [FDdata\(\)](#) for details.

- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.

See [ConvergenceInfo\(\)](#) for details

See also:

[get\\_field\\_CW\(\)](#), [gen\\_mass\\_matrix\(\)](#), and [gen\\_sources\(\)](#)

### nirfasterff.forward.femdata.femdata\_stnd\_FD

```
nirfasterff.forward.femdata.femdata_stnd_FD(mesh, freq, solver=utils.get_solver(),  
                                             opt=utils.SolverOptions())
```

Forward modeling for FD. Please consider using mesh.femdata(freq) instead. freq in Hz

The function calculates the MASS matrix, the source vectors, and calls the FD solver (preconditioned BiCGStab).

#### Parameters

- **mesh** (*nirfasterff.base.stndmesh*) – the mesh used to calculate the forward data.
- **freq** (*double*) – modulation frequency in Hz.  
When it is 0, function continues with the BiCGstab solver, but generates a warning that the CW solver should be used for better performance
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

#### Raises

**TypeError** – If mesh is not a stnd mesh.

#### Returns

- **data** (*nirfasterff.base.FDdata*) – fluence and boundary measurables given the mesh and optodes.

If mesh.vol is defined, the returned fluence will be in volumetric space

See [FDdata\(\)](#) for details.

- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.

See [ConvergenceInfo\(\)](#) for details

See also:

[get\\_field\\_FD\(\)](#), [gen\\_mass\\_matrix\(\)](#), and [gen\\_sources\(\)](#)

**nirfasterff.forward.femdata.femdata\_stnd\_TR**

```
nirfasterff.forward.femdata.femdata_stnd_TR(mesh, tmax, dt, savefield=False, beautify=True,
                                             solver=utils.get_solver(), opt=utils.SolverOptions())
```

Forward modeling calculating TPSF on a standard mesh. Please consider using `mesh.femdata_tpsf(tmax, dt)` instead.

The function calculates the MASS matrices, the source vectors, and calls the standard TPSF solver (preconditioned conjugated gradient).

**Parameters**

- **mesh** (*nirfasterff.base.stndmesh*) – the mesh used to calculate the forward data.
- **tmax** (*double*) – maximum time simulated, in seconds.
- **dt** (*double*) – size of each time step, in seconds.
- **savefield** (*bool, optional*) – If True, the internal TPSFs are also returned. If False, only boundary TPSFs are returned and `data.phi` will be empty.  
The default is False.
- **beautify** (*bool, optional*) – If true, zeros the initial unstable parts of the boundary TPSFs. The default is True.
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

**Raises**

- **TypeError** – If mesh is not a stnd mesh.
- **ValueError** – If tmax is smaller than dt

**Returns**

- **data** (*nirfasterff.base.TPSFdata*) – internal and boundary TPSFs given the mesh and optodes.

If `mesh.vol` is defined and `savefield==True`, the returned internal TPSFs will be in volumetric space

See [TPSFdata\(\)](#) for details.

- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.

Only the convergence info of the last time step is returned.

See [ConvergenceInfo\(\)](#) for details

**See also:**

[gen\\_mass\\_matrix\(\)](#), [gen\\_sources\(\)](#), and [get\\_field\\_TR\(\)](#)

## nirfasterff.forward.femdata.femdata\_stnd\_TR\_moments

```
nirfasterff.forward.femdata.femdata_stnd_TR_moments(mesh, max_moments=3, savefield=False,
                                                    solver=utils.get_solver(),
                                                    opt=utils.SolverOptions())
```

Forward modeling calculating TR moments using Mellin transform on a standard mesh. Please consider using `mesh.femdata_moments()` instead.

The function calculates the MASS matrices, the source vectors, and calls the Mellin moments solver (preconditioned conjugated gradient).

Calculates 0th, 1st, 2nd, ..., `max_moments`-th moments directly without calculating TPSF first.

This is more efficient, if the actual TPSFs are not of concern.

### Parameters

- **mesh** (*nirfasterff.base.stndmesh*) – the mesh used to calculate the forward data.
- **max\_moments** (*int32, optional*) – max order of moments to calculate. That is, 0th, 1st, 2nd, ..., `max_moments`-th will be calculated. The default is 3.
- **savefield** (*bool, optional*) – If True, the internal moments are also returned. If False, only boundary moments are returned and `data.phi` will be empty.

The default is False.

- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

### Raises

- **TypeError** – If mesh is not a stnd mesh.
- **ValueError** – If `max_moments` is negative.

### Returns

- **data** (*nirfasterff.base.TRMomentsdata*) – internal and boundary moments given the mesh and optodes.

If `mesh.vol` is defined and `savefield==True`, the returned internal moments will be in volumetric space

See [TRMomentsdata\(\)](#) for details.

- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.

Only the convergence info of highest order moments is returned.

See [ConvergenceInfo\(\)](#) for details

See also:

[gen\\_mass\\_matrix\(\)](#), [gen\\_sources\(\)](#), and [get\\_field\\_TRmoments\(\)](#)

## References

Arridge and Schweiger, Applied Optics, 1995. doi:10.1364/AO.34.002683

### 4.1.3 nirfasterff.inverse

Calculation of the Jacobian matrices and a basic Tikhonov regularization function

#### Functions

<code>jacobian_DCS(mesh, tvec[, normalize, ...])</code>	Calculates the Jacobian matrix for a DCS mesh using the adjoint method
<code>jacobian_fl_CW(mesh[, normalize, solver, opt])</code>	Calculates the continuous-wave fluorescence Jacobian matrix using the adjoint method
<code>jacobian_fl_FD(mesh, freq[, normalize, ...])</code>	Calculates the frequency-domain fluorescence Jacobian matrix using the adjoint method
<code>jacobian_stnd_CW(mesh[, normalize, mus, ...])</code>	Calculates the continuous-wave Jacobian matrix using the adjoint method
<code>jacobian_stnd_FD(mesh, freq[, normalize, ...])</code>	Calculates the frequency-domain Jacobian matrix using the adjoint method
<code>tikhonov(A, reg, y)</code>	Solves Tikhonov regularization (ie ridge regression)

#### nirfasterff.inverse.jacobian\_DCS

`nirfasterff.inverse.jacobian_DCS(mesh, tvec, normalize=True, solver=utils.get_solver(),  
opt=utils.SolverOptions())`

Calculates the Jacobian matrix for a DCS mesh using the adjoint method

One Jacobian is calculated at each time point in tvec, and the derivative is taken with regard to aDb

##### Parameters

- **mesh** (*nirfasterff.base.dcsmesh*) – mesh on which the Jacobian is calculated.
- **tvec** (*double NumPy vector*) – time vector used.
- **normalize** (*bool, optional*) – if True, Jacobians are normalized to the measured boundary amplitude. The default is True.
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See `SolverOptions()` for details

##### Raises

**TypeError** – if mesh is not a DCS mesh, or mesh.vol is not defined.

##### Returns

- **J** (*double NumPy array*) – The Jacobian matrix. Size (NChannel, NVoxel, NTime)
- **data1** (*nirfasterff.base.FLdata*) – The calculated direct field. The same as directly calling `mesh.femdata(tvec)`

- **data2** (*nirfasterff.base.FLdata*) – The calculated adjoint field. The same as calling `mesh.femdata(tvec)` AFTER swapping the locations of sources and detectors

### **nirfasterff.inverse.jacobian\_fl\_CW**

`nirfasterff.inverse.jacobian_fl_CW(mesh, normalize=True, solver=utils.get_solver(),  
opt=utils.SolverOptions())`

Calculates the continuous-wave fluorescence Jacobian matrix using the adjoint method

$$J_{ij} = dA_i / d_{\gamma_j}$$

where  $A$  is fluorescence amplitude if normalization is False, fluorescence amplitude divided by excitation amplitude ('Born ratio') if True

$\gamma_j = \text{mesh.eta}[j] * \text{mesh.muaf}[j]$

#### **Parameters**

- **mesh** (*nirfasterff.base.fluormesh*) – mesh on which the Jacobian is calculated.
- **normalize** (*bool, optional*) – whether normalize the Jacobian to the amplitudes of boundary measurements at excitation wavelength ('Born ratio').  
The default is True.
- **solver** (*str, optional*) – Choose between 'CPU' or 'GPU' solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [\*SolverOptions\(\)\*](#) for details

#### **Raises**

**TypeError** – if mesh is not a fluor mesh, or mesh.vol is not defined.

#### **Returns**

- **J** (*double NumPy array*) – The Jacobian matrix. Size (NChannel, NVoxel)
- **data1** (*nirfasterff.base.FLdata*) – The calculated direct field. The same as directly calling `mesh.femdata(0)`
- **data2** (*nirfasterff.base.FLdata*) – The calculated adjoint field. The same as calling `mesh.femdata(0)` AFTER swapping the locations of sources and detectors

### **References**

Milstein et al., JOSA A, 2004. doi:10.1364/JOSAA.21.001035



## nirfasterff.inverse.jacobian\_fl\_FD

```
nirfasterff.inverse.jacobian_fl_FD(mesh, freq, normalize=True, solver=utils.get_solver(),
                                   opt=utils.SolverOptions())
```

Calculates the frequency-domain fluorescence Jacobian matrix using the adjoint method

The Jacobian is structured as, suppose we have M channels and N voxels:

```
[d_real{A_1}/d_gamma_1, d_real{A_1}/d_gamma_2, ..., d_real{A_1}/d_gamma_{N}, d_real
↪{A_1}/d_tau_1, d_real{A_1}/d_tau_2, ..., d_real{A_1}/d_tau_{N}]
[d_imag{A_1}/d_gamma_1, d_imag{A_1}/d_gamma_2, ..., d_imag{A_1}/d_gamma_{N}, d_imag
↪{A_1}/d_tau_1, d_imag{A_1}/d_tau_2, ..., d_imag{A_1}/d_tau_{N}]
[d_real{A_2}/d_gamma_1, d_real{A_2}/d_gamma_2, ..., d_real{A_2}/d_gamma_{N}, d_real
↪{A_2}/d_tau_1, d_real{A_2}/d_tau_2, ..., d_real{A_2}/d_tau_{N}]
[d_imag{A_2}/d_gamma_1, d_imag{A_2}/d_gamma_2, ..., d_imag{A_2}/d_gamma_{N}, d_imag
↪{A_2}/d_tau_1, d_imag{A_2}/d_tau_2, ..., d_imag{A_2}/d_tau_{N}]
...
[d_real{A_M}/d_gamma_1, d_real{A_M}/d_gamma_2, ..., d_real{A_M}/d_gamma_{N}, d_real
↪{A_M}/d_tau_1, d_real{A_M}/d_tau_2, ..., d_real{A_M}/d_tau_{N}]
[d_imag{A_M}/d_gamma_1, d_imag{A_M}/d_gamma_2, ..., d_imag{A_M}/d_gamma_{N}, d_imag
↪{A_M}/d_tau_1, d_imag{A_M}/d_tau_2, ..., d_imag{A_M}/d_tau_{N}]
```

where A is fluorescence amplitude if normalization is False, fluorescence amplitude divided by excitation amplitude ('Born ratio') if True  $\gamma_j = \text{mesh.eta}[j] * \text{mesh.muaf}[j]$

### Parameters

- **mesh** (*nirfasterff.base.fluormesh*) – mesh on which the Jacobian is calculated.
- **freq** (*double*) – modulation frequency in Hz.
- **normalize** (*bool, optional*) – whether normalize the Jacobian to the amplitudes of boundary measurements at excitation wavelength ('Born ratio').

The default is True.

- **solver** (*str, optional*) – Choose between 'CPU' or 'GPU' solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

### Raises

**TypeError** – if mesh is not a fluor mesh, or mesh.vol is not defined.

### Returns

- **J** (*double NumPy array*) – The Jacobian matrix. Size (NChannel\*2, NVoxel\*2)
- **data1** (*nirfasterff.base.FLdata*) – The calculated direct field. The same as directly calling `mesh.femdata(freq)`
- **data2** (*nirfasterff.base.FLdata*) – The calculated adjoint field. The same as calling `mesh.femdata(freq)` AFTER swapping the locations of sources and detectors

## References

Milstein et al., JOSA A, 2004. doi:10.1364/JOSAA.21.001035

## nirfasterff.inverse.jacobian\_std\_CW

`nirfasterff.inverse.jacobian_std_CW(mesh, normalize=True, mus=False, solver=utils.get_solver(), opt=utils.SolverOptions())`

Calculates the continuous-wave Jacobian matrix using the adjoint method

Calculates spatial distributions of sensitivity of field registered on the mesh boundary to changes of optical properties per voxel.

When mus is set to True, the Jacobian is structured as, suppose we have M channels and N voxels:

```
[dA_1/dmusp_1, dA_1/dmusp_2, ..., dA_1/dmusp_{N}, dA_1/dmua_1, dA_1/dmua_2, ..., dA_
↪ 1/dmua_{N}]
[dA_2/dmusp_1, dA_2/dmusp_2, ..., dA_2/dmusp_{N}, dA_2/dmua_1, dA_2/dmua_2, ..., dA_
↪ 2/dmua_{N}]
...
[dA_M/dmusp_1, dA_M/dmusp_2, ..., dA_M/dmusp_{N}, dA_M/dmua_1, dA_M/dmua_2, ..., dA_
↪ M/dmua_{N}]
```

where A and Phi denote the measured amplitude and the phase if *normalize=False*, and the log of them if *normalize=True*

When mus is set to False, the returned Jacobian is only the right half of the above. That is, only derivatives wrt mua

Note that the calculation is only done in the volumetric space

### Parameters

- **mesh** (*nirfasterff.base.stndmesh*) – mesh on which the Jacobian is calculated.
- **normalize** (*bool, optional*) – whether normalize the Jacobian to the amplitudes of boundary measurements, i.e. use Rytov approximation.  
The default is True.
- **mus** (*bool, optional*) – whether derivatives wrt mus (left half of the ‘full’ Jacobian) is calculated. The default is False.
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

### Raises

**TypeError** – if mesh is not a stnd mesh, or mesh.vol is not defined.

### Returns

- **J** (*double NumPy array*) – The Jacobian matrix. Size (NChannel, NVoxel\*2) if mus=True, (NChannel, NVoxel) if mus=False
- **data1** (*nirfasterff.base.FDdata*) – The calculated direct field. The same as directly calling `mesh.femdata(0)`

- **data2** (*nirfasterff.base.FDdata*) – The calculated adjoint field. The same as calling `mesh.femdata(0)` AFTER swapping the locations of sources and detectors

### nirfasterff.inverse.jacobian\_stnd\_FD

`nirfasterff.inverse.jacobian_stnd_FD(mesh, freq, normalize=True, mus=True, solver=utils.get_solver(), opt=utils.SolverOptions())`

Calculates the frequency-domain Jacobian matrix using the adjoint method

Calculates spatial distributions of sensitivity of field registered on the mesh boundary to changes of optical properties per voxel.

When `freq=0`, see `jacobian_stnd_CW()` for the structure of the Jacobian

When `freq>0`, the Jacobian is structured as, suppose we have `M` channels and `N` voxels:

```
[dA_1/dmusp_1, dA_1/dmusp_2, ..., dA_1/dmusp_{N}, dA_1/dmua_1, dA_1/dmua_2, ..., dA_
↪1/dmua_{N}]
[dPhi_1/dmusp_1, dPhi_1/dmusp_2, ..., dPhi_1/dmusp_{N}, dPhi_1/dmua_1, dPhi_1/dmua_
↪2, ..., dPhi_1/dmua_{N}]
[dA_2/dmusp_1, dA_2/dmusp_2, ..., dA_2/dmusp_{N}, dA_2/dmua_1, dA_2/dmua_2, ..., dA_
↪2/dmua_{N}]
[dPhi_2/dmusp_1, dPhi_2/dmusp_2, ..., dPhi_2/dmusp_{N}, dPhi_2/dmua_1, dPhi_2/dmua_
↪2, ..., dPhi_2/dmua_{N}]
...
[dA_M/dmusp_1, dA_M/dmusp_2, ..., dA_M/dmusp_{N}, dA_M/dmua_1, dA_M/dmua_2, ..., dA_
↪M/dmua_{N}]
[dPhi_M/dmusp_1, dPhi_M/dmusp_2, ..., dPhi_M/dmusp_{N}, dPhi_M/dmua_1, dPhi_M/dmua_
↪2, ..., dPhi_M/dmua_{N}]
```

where `A` and `Phi` denote the measured amplitude and the phase if `normalize=False`, and the log of them if `normalize=True`

Note that the calculation is only done in the volumetric space

#### Parameters

- **mesh** (*nirfasterff.base.stndmesh*) – mesh on which the Jacobian is calculated.
- **freq** (*double*) – modulation frequency, in Hz.
- **normalize** (*bool, optional*) – whether normalize the Jacobian to the amplitudes of boundary measurements, i.e. use Rytov approximation.

The default is `True`.

- **mus** (*bool, optional*) – whether derivates wrt `mus` (left half of the ‘full’ Jacobian) is calculated. Only has effect when `freq=0`. The default is `True`.
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See `SolverOptions()` for details

#### Raises

**TypeError** – if `mesh` is not a `stnd mesh`, or `mesh.vol` is not defined.

#### Returns

- **J** (*double or complex double NumPy array*) – The Jacobian matrix. Size (NChannel\*2, NVoxel\*2) if freq>0, (NChannel, NVoxel\*2) if freq=0 and mus=True, (NChannel, NVoxel) if freq=0 and mus=False
- **data1** (*nirfasterff.base.FDdata*) – The calculated direct field. The same as directly calling mesh.femdata(freq)
- **data2** (*nirfasterff.base.FDdata*) – The calculated adjoint field. The same as calling mesh.femdata(freq) AFTER swapping the locations of sources and detectors

## References

Arridge, Applied Optics, 1995. doi:10.1364/AO.34.007395

### nirfasterff.inverse.tikhonov

nirfasterff.inverse.tikhonov(*A, reg, y*)

Solves Tikhonov regularization (ie ridge regression)

That is, given a linear system  $y = Ax$ , it solves  $\arg \min_x ||Ax - y||_2^2 + ||\Gamma x||_2^2$

where  $A$  is the forward matrix,  $y$  is the recording, and  $\Gamma$  is the regularization matrix

#### Parameters

- **A** (*double NumPy array*) – forward matrix, e.g. the Jacobian in DOT.
- **reg** (*double NumPy array or scalar*) – if a scalar, the same regularization is applied to all elements of  $x$ , i.e.  $\Gamma = reg * I$ .  
if a vector, the regularization matrix is assumed to be diagonal, with the diagonal elements specified in  $reg$ , i.e.  $\Gamma = diag(reg)$ .  
if a matrix, it must be symmetric. In this case,  $\Gamma = reg$ .
- **y** (*double NumPy vector*) – measurement vector, e.g. dOD at each channel in DOT.

#### Raises

**ValueError** – if  $reg$  is of incompatible size or not symmetric (if matrix).

#### Returns

**result** – Tikhonov regularized solution to the linear system.

#### Return type

double NumPy vector

### 4.1.4 nirfasterff.io

Some functions for reading/writing certain data types.

As of now, they are only used by the CGAL mesher, and there should be no need for the user to directly call them.

## Functions

<code>readMEDIT(fname)</code>	Read a mesh generated by the CGAL mesher, which is saved in MEDIT format
<code>saveinr(vol, fname[, xPixelSpacing, ...])</code>	Save a volume in the INRIA format.

### nirfasterff.io.readMEDIT

`nirfasterff.io.readMEDIT(fname)`

Read a mesh generated by the CGAL mesher, which is saved in MEDIT format

Directly translated from the Matlab version

#### Parameters

**fname** (*str*) – name of the file to be loaded.

#### Returns

- **elements** (*NumPy array*) – list of elements in the mesh. Zero-based
- **nodes** (*NumPy array*) – node locations of the mesh, in mm.
- **faces** (*NumPy array*) – list of faces in the mesh. In case of 2D, it's the same as elements. Zero-based
- **nnpe** (*int*) – size of dimension 1 of elements, i.e. 4 for 3D mesh and 3 for 2D mesh.

### nirfasterff.io.saveinr

`nirfasterff.io.saveinr(vol, fname, xPixelSpacing=1., yPixelSpacing=1., SliceThickness=1.)`

Save a volume in the INRIA format. This is for the CGAL mesher.

Directly translated from the Matlab version

#### Parameters

- **vol** (*NumPy array*) – the volume to be saved.
- **fname** (*str*) – file name to be saved as.
- **xPixelSpacing** (*double, optional*) – volume resolution in x direction. The default is 1..
- **yPixelSpacing** (*double, optional*) – volume resolution in y direction. The default is 1..
- **SliceThickness** (*double, optional*) – volume resolution in z direction. The default is 1..

#### Return type

None.

### 4.1.5 nirfasterff.lib

Low-level functions implemented in C/C++, on both GPU and CPU

You should NOT directly call these functions. Please use the wrapper functions provided instead

#### Modules

---

*nirfasterff.lib.nirfasterff\_cpu**nirfasterff.lib.nirfasterff\_cuda*

---

#### nirfasterff.lib.nirfasterff\_cpu

#### Functions

---

*IntGradGrid*(\*args, \*\*kwargs) Overloaded function.*IntGrid*(\*args, \*\*kwargs) Overloaded function.*ele\_area*(arg0, arg1)*gen\_mass\_matrix*(arg0, arg1, arg2, arg3, ...)*gen\_source\_fl*(\*args, \*\*kwargs) Overloaded function.*get\_field\_CW*(\*args, \*\*kwargs) Overloaded function.*get\_field\_FD*(\*args, \*\*kwargs) Overloaded function.*get\_field\_TR*(\*args, \*\*kwargs) Overloaded function.*get\_field\_TRFL*(arg0, arg1, arg2, arg3, arg4, ...)*get\_field\_TRFL\_moments*(arg0, arg1, arg2, ...)*get\_field\_TR\_moments*(\*args, \*\*kwargs) Overloaded function.*gradientIntfunc*(arg0, arg1, arg2)*gradientIntfunc2*(arg0, arg1, arg2)*isCUDA*()*mesh\_support*(arg0, arg1, arg2)*pointLocation*(arg0, arg1, arg2)

---

**nirfasterff.lib.nirfasterff\_cpu.IntGradGrid****nirfasterff.lib.nirfasterff\_cpu.IntGradGrid(\*args, \*\*kwargs)**

Overloaded function.

1. IntGradGrid(arg0: numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c\_contiguous],  
arg1: numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c\_contiguous],  
arg2: numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c\_contiguous],  
arg3: numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c\_contiguous],  
arg4: numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c\_contiguous],  
arg5: numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c\_contiguous], arg6:  
numpy.ndarray[numpy.int32[m, n], flags.writeable, flags.c\_contiguous], arg7: int) ->  
numpy.ndarray[numpy.complex128[m, n]]
2. IntGradGrid(arg0: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous],  
arg1: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous],  
arg2: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous],  
arg3: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous],  
arg4: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous],  
arg5: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous], arg6:  
numpy.ndarray[numpy.int32[m, n], flags.writeable, flags.c\_contiguous], arg7: int) ->  
numpy.ndarray[numpy.float64[m, n]]

**nirfasterff.lib.nirfasterff\_cpu.IntGrid****nirfasterff.lib.nirfasterff\_cpu.IntGrid(\*args, \*\*kwargs)**

Overloaded function.

1. IntGrid(arg0: numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c\_contiguous],  
arg1: numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c\_contiguous],  
arg2: numpy.ndarray[numpy.int32[m, n], flags.writeable, flags.c\_contiguous]) ->  
numpy.ndarray[numpy.complex128[m, n]]
2. IntGrid(arg0: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous],  
arg1: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous],  
arg2: numpy.ndarray[numpy.int32[m, n], flags.writeable, flags.c\_contiguous]) ->  
numpy.ndarray[numpy.float64[m, n]]

**nirfasterff.lib.nirfasterff\_cpu.ele\_area**

**nirfasterff.lib.nirfasterff\_cpu.ele\_area**(arg0: numpy.ndarray[numpy.float64[m, n], flags.writeable,  
flags.c\_contiguous], arg1: numpy.ndarray[numpy.float64[m,  
n], flags.writeable, flags.c\_contiguous]) →  
numpy.ndarray[numpy.float64[m, 1]]

**nirfasterff.lib.nirfasterff\_cpu.gen\_mass\_matrix**

`nirfasterff.lib.nirfasterff_cpu.gen_mass_matrix`(arg0: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg1: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg2: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg3: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg4: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg5: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg6: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg7: *float*) → *tuple[numpy.ndarray[numpy.int32[m, 1]], numpy.ndarray[numpy.int32[m, 1]], numpy.ndarray[numpy.complex128[m, 1]]]*

**nirfasterff.lib.nirfasterff\_cpu.gen\_source\_fl**

`nirfasterff.lib.nirfasterff_cpu.gen_source_fl`(\*args, \*\*kwargs)

Overloaded function.

1. `gen_source_fl`(arg0: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg1: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg2: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*) → *numpy.ndarray[numpy.float64[m, n]]*
2. `gen_source_fl`(arg0: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg1: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg2: *numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c\_contiguous]*) → *numpy.ndarray[numpy.complex128[m, n]]*

**nirfasterff.lib.nirfasterff\_cpu.get\_field\_CW**

`nirfasterff.lib.nirfasterff_cpu.get_field_CW`(\*args, \*\*kwargs)

Overloaded function.

1. `get_field_CW`(arg0: *numpy.ndarray[numpy.int32[m, 1], flags.writeable]*, arg1: *numpy.ndarray[numpy.int32[m, 1], flags.writeable]*, arg2: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg3: *scipy.sparse.csc\_matrix[numpy.float64]*, max\_iter: *int = 1000*, AbsoluteTolerance: *float = 1e-12*, RelativeTolerance: *float = 1e-12*, divergence: *float = 100000000.0*) → *tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff\_cpu.ConvergenceInfoCPU]]*
2. `get_field_CW`(arg0: *numpy.ndarray[numpy.int32[m, 1], flags.writeable]*, arg1: *numpy.ndarray[numpy.int32[m, 1], flags.writeable]*, arg2: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg3: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, max\_iter: *int = 1000*, AbsoluteTolerance: *float = 1e-12*, RelativeTolerance: *float = 1e-12*, divergence: *float = 100000000.0*) → *tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff\_cpu.ConvergenceInfoCPU]]*



**nirfasterff.lib.nirfasterff\_cpu.get\_field\_FD**

`nirfasterff.lib.nirfasterff_cpu.get_field_FD(*args, **kwargs)`

Overloaded function.

1. `get_field_FD(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.complex128[m, 1], flags.writeable], arg3: scipy.sparse.csc_matrix[numpy.complex128], max_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0) -> tuple[numpy.ndarray[numpy.complex128[m, n]], list[nirfasterff.lib.nirfasterff_cpu.ConvergenceInfoCPU]]`
2. `get_field_FD(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.complex128[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c_contiguous], max_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0) -> tuple[numpy.ndarray[numpy.complex128[m, n]], list[nirfasterff.lib.nirfasterff_cpu.ConvergenceInfoCPU]]`

**nirfasterff.lib.nirfasterff\_cpu.get\_field\_TR**

`nirfasterff.lib.nirfasterff_cpu.get_field_TR(*args, **kwargs)`

Overloaded function.

1. `get_field_TR(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg4: scipy.sparse.csc_matrix[numpy.float64], arg5: int, max_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0) -> tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff_cpu.ConvergenceInfoCPU]]`
2. `get_field_TR(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg4: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c_contiguous], arg5: int, max_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0) -> tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff_cpu.ConvergenceInfoCPU]]`

**nirfasterff.lib.nirfasterff\_cpu.get\_field\_TRFL**

`nirfasterff.lib.nirfasterff_cpu.get_field_TRFL(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg4: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c_contiguous], arg5: int, max_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0) -> tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff_cpu.ConvergenceInfoCPU]]`

**nirfasterff.lib.nirfasterff\_cpu.get\_field\_TRFL\_moments**

```
nirfasterff.lib.nirfasterff_cpu.get_field_TRFL_moments(arg0: numpy.ndarray[numpy.int32[m, 1],
                                                    flags.writeable], arg1:
                                                    numpy.ndarray[numpy.int32[m, 1],
                                                    flags.writeable], arg2:
                                                    numpy.ndarray[numpy.float64[m, 1],
                                                    flags.writeable], arg3:
                                                    numpy.ndarray[numpy.float64[m, 1],
                                                    flags.writeable], arg4:
                                                    numpy.ndarray[numpy.float64[m, n],
                                                    flags.writeable, flags.c_contiguous], arg5:
                                                    numpy.ndarray[numpy.float64[m, 1],
                                                    flags.writeable], arg6:
                                                    numpy.ndarray[numpy.float64[m, 1],
                                                    flags.writeable], arg7: int, max_iter: int =
                                                    1000, AbsoluteTolerance: float = 1e-12,
                                                    RelativeTolerance: float = 1e-12,
                                                    divergence: float = 100000000.0) →
                                                    tuple[numpy.ndarray[numpy.float64[m, n]],
                                                    list[nirfasterff.lib.nirfasterff_cpu.ConvergenceInfoCPU]]
```

**nirfasterff.lib.nirfasterff\_cpu.get\_field\_TR\_moments**

```
nirfasterff.lib.nirfasterff_cpu.get_field_TR_moments(*args, **kwargs)
```

Overloaded function.

1. `get_field_TR_moments`(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg4: scipy.sparse.csc\_matrix[numpy.float64], arg5: int, max\_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0) -> tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff\_cpu.ConvergenceInfoCPU]]
2. `get_field_TR_moments`(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg4: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous], arg5: int, max\_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0) -> tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff\_cpu.ConvergenceInfoCPU]]

**nirfasterff.lib.nirfasterff\_cpu.gradientIntfunc**

```
nirfasterff.lib.nirfasterff_cpu.gradientIntfunc(arg0: numpy.ndarray[numpy.float64[m, n],
                                                    flags.writeable, flags.c_contiguous], arg1:
                                                    numpy.ndarray[numpy.float64[m, n], flags.writeable,
                                                    flags.c_contiguous], arg2:
                                                    numpy.ndarray[numpy.float64[m, n], flags.writeable,
                                                    flags.c_contiguous]) →
                                                    tuple[numpy.ndarray[numpy.int32[m, 1]],
                                                    numpy.ndarray[numpy.float64[m, n]]]
```

**nirfasterff.lib.nirfasterff\_cpu.gradientIntfunc2**

**nirfasterff.lib.nirfasterff\_cpu.gradientIntfunc2**(arg0: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg1: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg2: *numpy.ndarray[numpy.int32[m, 1], flags.writeable]*)  
 → *numpy.ndarray[numpy.float64[m, n]]*

**nirfasterff.lib.nirfasterff\_cpu.isCUDA**

**nirfasterff.lib.nirfasterff\_cpu.isCUDA**() → bool

**nirfasterff.lib.nirfasterff\_cpu.mesh\_support**

**nirfasterff.lib.nirfasterff\_cpu.mesh\_support**(arg0: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg1: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg2: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*) →  
*numpy.ndarray[numpy.float64[m, 1]]*

**nirfasterff.lib.nirfasterff\_cpu.pointLocation**

**nirfasterff.lib.nirfasterff\_cpu.pointLocation**(arg0: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg1: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg2: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*) →  
*tuple[numpy.ndarray[numpy.int32[m, 1]], numpy.ndarray[numpy.float64[m, n]]]*

**Classes**


---

*ConvergenceInfoCPU*


---

**nirfasterff.lib.nirfasterff\_cpu.ConvergenceInfoCPU****class** nirfasterff.lib.nirfasterff\_cpu.ConvergenceInfoCPU

Bases: pybind11\_object

**\_\_init\_\_**(self: nirfasterff.lib.nirfasterff\_cpu.ConvergenceInfoCPU) → None**Methods**

<code>__init__(self)</code>
-----------------------------

**Attributes**

<code>isConverged</code>
--------------------------

<code>isConvergedToAbsoluteTolerance</code>
---

<code>iteration</code>
------------------------

<code>residual</code>
-----------------------

**nirfasterff.lib.nirfasterff\_cuda****Functions**

<code>gen_mass_matrix(arg0, arg1, arg2, arg3, ...)</code>	
---	--

<code>gen_source_fl(*args, **kwargs)</code>	Overloaded function.
---	----------------------

<code>get_field_CW(*args, **kwargs)</code>	Overloaded function.
--	----------------------

<code>get_field_FD(*args, **kwargs)</code>	Overloaded function.
--	----------------------

<code>get_field_TR(*args, **kwargs)</code>	Overloaded function.
--	----------------------

<code>get_field_TRFL(arg0, arg1, arg2, arg3, arg4, ...)</code>	
--	--

<code>get_field_TRFL_moments(arg0, arg1, arg2, ...)</code>	
--	--

<code>get_field_TR_moments(*args, **kwargs)</code>	Overloaded function.
--	----------------------

**nirfasterff.lib.nirfasterff\_cuda.gen\_mass\_matrix**

**nirfasterff.lib.nirfasterff\_cuda.gen\_mass\_matrix**(arg0: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg1: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg2: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg3: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg4: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg5: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg6: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg7: *float, GPU: int = -1*) → *tuple[numpy.ndarray[numpy.int32[m, 1]], numpy.ndarray[numpy.int32[m, 1]], numpy.ndarray[numpy.complex128[m, 1]]]*

**nirfasterff.lib.nirfasterff\_cuda.gen\_source\_fl**

**nirfasterff.lib.nirfasterff\_cuda.gen\_source\_fl**(\*args, \*\*kwargs)

Overloaded function.

1. **gen\_source\_fl**(arg0: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg1: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg2: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, GPU: *int = -1*) → *numpy.ndarray[numpy.float64[m, n]]*
2. **gen\_source\_fl**(arg0: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg1: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, arg2: *numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c\_contiguous]*, GPU: *int = -1*) → *numpy.ndarray[numpy.complex128[m, n]]*

**nirfasterff.lib.nirfasterff\_cuda.get\_field\_CW**

**nirfasterff.lib.nirfasterff\_cuda.get\_field\_CW**(\*args, \*\*kwargs)

Overloaded function.

1. **get\_field\_CW**(arg0: *numpy.ndarray[numpy.int32[m, 1], flags.writeable]*, arg1: *numpy.ndarray[numpy.int32[m, 1], flags.writeable]*, arg2: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg3: *scipy.sparse.csc\_matrix[numpy.float64]*, max\_iter: *int = 1000*, AbsoluteTolerance: *float = 1e-12*, RelativeTolerance: *float = 1e-12*, divergence: *float = 100000000.0*, GPU: *int = -1*) → *tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff\_cuda.ConvergenceInfoGPU]]*
2. **get\_field\_CW**(arg0: *numpy.ndarray[numpy.int32[m, 1], flags.writeable]*, arg1: *numpy.ndarray[numpy.int32[m, 1], flags.writeable]*, arg2: *numpy.ndarray[numpy.float64[m, 1], flags.writeable]*, arg3: *numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous]*, max\_iter: *int = 1000*, AbsoluteTolerance: *float = 1e-12*, RelativeTolerance: *float = 1e-12*, divergence: *float = 100000000.0*, GPU: *int = -1*) → *tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff\_cuda.ConvergenceInfoGPU]]*

**nirfasterff.lib.nirfasterff\_cuda.get\_field\_FD****nirfasterff.lib.nirfasterff\_cuda.get\_field\_FD**(\*args, \*\*kwargs)

Overloaded function.

1. `get_field_FD`(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.complex128[m, 1], flags.writeable], arg3: scipy.sparse.csc\_matrix[numpy.complex128], max\_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0, GPU: int = -1) -> tuple[numpy.ndarray[numpy.complex128[m, n]], list[nirfasterff.lib.nirfasterff\_cuda.ConvergenceInfoGPU]]
2. `get_field_FD`(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.complex128[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.complex128[m, n], flags.writeable, flags.c\_contiguous], max\_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0, GPU: int = -1) -> tuple[numpy.ndarray[numpy.complex128[m, n]], list[nirfasterff.lib.nirfasterff\_cuda.ConvergenceInfoGPU]]

**nirfasterff.lib.nirfasterff\_cuda.get\_field\_TR****nirfasterff.lib.nirfasterff\_cuda.get\_field\_TR**(\*args, \*\*kwargs)

Overloaded function.

1. `get_field_TR`(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg4: scipy.sparse.csc\_matrix[numpy.float64], arg5: int, max\_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0, GPU: int = -1) -> tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff\_cuda.ConvergenceInfoGPU]]
2. `get_field_TR`(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg4: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous], arg5: int, max\_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0, GPU: int = -1) -> tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff\_cuda.ConvergenceInfoGPU]]

**nirfasterff.lib.nirfasterff\_cuda.get\_field\_TRFL****nirfasterff.lib.nirfasterff\_cuda.get\_field\_TRFL**(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg4: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c\_contiguous], arg5: int, max\_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0, GPU: int = -1) -> tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff\_cuda.ConvergenceInfoGPU]]

**nirfasterff.lib.nirfasterff\_cuda.get\_field\_TRFL\_moments**

```
nirfasterff.lib.nirfasterff_cuda.get_field_TRFL_moments(arg0: numpy.ndarray[numpy.int32[m, 1],
                                                    flags.writeable], arg1:
                                                    numpy.ndarray[numpy.int32[m, 1],
                                                    flags.writeable], arg2:
                                                    numpy.ndarray[numpy.float64[m, 1],
                                                    flags.writeable], arg3:
                                                    numpy.ndarray[numpy.float64[m, 1],
                                                    flags.writeable], arg4:
                                                    numpy.ndarray[numpy.float64[m, n],
                                                    flags.writeable, flags.c_contiguous], arg5:
                                                    numpy.ndarray[numpy.float64[m, 1],
                                                    flags.writeable], arg6:
                                                    numpy.ndarray[numpy.float64[m, 1],
                                                    flags.writeable], arg7: int, max_iter: int =
1000, AbsoluteTolerance: float = 1e-12,
RelativeTolerance: float = 1e-12,
divergence: float = 100000000.0, GPU: int
= -1) →
tuple[numpy.ndarray[numpy.float64[m, n]],
list[nirfasterff.lib.nirfasterff_cuda.ConvergenceInfoGPU]]
```

**nirfasterff.lib.nirfasterff\_cuda.get\_field\_TR\_moments**

```
nirfasterff.lib.nirfasterff_cuda.get_field_TR_moments(*args, **kwargs)
```

Overloaded function.

1. `get_field_TR_moments(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg4: scipy.sparse.csc_matrix[numpy.float64], arg5: int, max_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0, GPU: int = -1) -> tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff_cuda.ConvergenceInfoGPU]]`
2. `get_field_TR_moments(arg0: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg1: numpy.ndarray[numpy.int32[m, 1], flags.writeable], arg2: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg3: numpy.ndarray[numpy.float64[m, 1], flags.writeable], arg4: numpy.ndarray[numpy.float64[m, n], flags.writeable, flags.c_contiguous], arg5: int, max_iter: int = 1000, AbsoluteTolerance: float = 1e-12, RelativeTolerance: float = 1e-12, divergence: float = 100000000.0, GPU: int = -1) -> tuple[numpy.ndarray[numpy.float64[m, n]], list[nirfasterff.lib.nirfasterff_cuda.ConvergenceInfoGPU]]`

## Classes

---

*ConvergenceInfoGPU*

---

### `nirfasterff.lib.nirfasterff_cuda.ConvergenceInfoGPU`

**class** `nirfasterff.lib.nirfasterff_cuda.ConvergenceInfoGPU`

Bases: `pybind11_object`

`__init__`(*self*: `nirfasterff.lib.nirfasterff_cuda.ConvergenceInfoGPU`) → `None`

## Methods

---

`__init__`(*self*)

---

## Attributes

---

`isConverged``isConvergedToAbsoluteTolerance``iteration``residual`

---

### 4.1.6 `nirfasterff.math`

Some low-level functions used by the forward solvers.

It is usually unnecessary to use them directly and caution must be exercised, as many of them interact closely with the C++ libraries and can cause crashes if used incorrectly



## Functions

<code>gen_mass_matrix(mesh, freq[, solver, GPU])</code>	Calculate the MASS matrix, and return the coordinates in CSR format.
<code>gen_sources(mesh)</code>	Calculate the source vectors (point source only) for the sources in mesh.source field
<code>gen_sources_fl(mesh, phix[, frequency, ...])</code>	Calculates FEM sources vector for re-emission.
<code>get_boundary_data(mesh, phi)</code>	Calculates boundary data given the field data in mesh
<code>get_field_CW(csrI, csrJ, csrV, qvec[, opt, ...])</code>	Call the Preconditioned Conjugate Gradient solver with FSAI preconditioner.
<code>get_field_FD(csrI, csrJ, csrV, qvec[, opt, ...])</code>	Call the Preconditioned BiConjugate Stablized solver with FSAI preconditioner.
<code>get_field_TR(csrI, csrJ, csrV, qvec, dt, ...)</code>	Call the Preconditioned Conjugate Gradient solver with FSAI preconditioner.
<code>get_field_TRFL(csrI, csrJ, csrV, phix, dt, ...)</code>	Call the Preconditioned Conjugate Gradient solver with FSAI preconditioner.
<code>get_field_TRFLmoments(csrI, csrJ, csrV, mx, ...)</code>	Call the Preconditioned Conjugate Gradient solver with FSAI preconditioner.
<code>get_field_TRmoments(csrI, csrJ, csrV, qvec, ...)</code>	Call the Preconditioned Conjugate Gradient solver with FSAI preconditioner.

### nirfasterff.math.gen\_mass\_matrix

`nirfasterff.math.gen_mass_matrix(mesh, freq, solver=utils.get_solver(), GPU=-1)`

Calculate the MASS matrix, and return the coordinates in CSR format.

The current Matlab version outputs COO format, so the results are NOT directly compatible

If calculation fails on GPU (if chosen), it will generate a warning and automatically switch to CPU

#### Parameters

- **mesh** (*nirfasterff.base.stndmesh*) – the mesh used to calculate the MASS matrix.
- **freq** (*double*) – modulation frequency, in Hz.
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **GPU** (*int, optional*) – GPU selection. -1 for automatic, 0, 1, ... for manual selection on multi-GPU systems. The default is -1.

#### Raises

- **RuntimeError** – if both CUDA and CPU versions fail.
- **TypeError** – if ‘solver’ is not ‘CPU’ or ‘GPU’.

#### Returns

- **csrI** (*int32 NumPy vector, zero-based*) – I indices of the MASS matrix, in CSR format. Size (NNodes,)
- **csrJ** (*int32 NumPy vector, zero-based*) – J indices of the MASS matrix, in CSR format. Size (nnz(MASS),)
- **csrV** (*float64 or complex128 NumPy vector*) – values of the MASS matrix, in CSR format. Size (nnz(MASS),)

### nirfasterff.math.gen\_sources

nirfasterff.math.gen\_sources(*mesh*)

Calculate the source vectors (point source only) for the sources in mesh.source field

**Parameters**

**mesh** (*NIRFASTer mesh type*) – mesh used to calculate the source vectors. Source information is also defined here.

**Returns**

**qvec** – source vectors, where each column corresponds to one source. Size (NNodes, Nsources).

**Return type**

complex double NumPy array

### nirfasterff.math.gen\_sources\_fl

nirfasterff.math.gen\_sources\_fl(*mesh, phix, frequency=0., solver=utils.get\_solver(), GPU=-1*)

Calculates FEM sources vector for re-emission.

If calculation fails on GPU (if chosen), it will generate a warning and automatically switch to CPU.

**Parameters**

- **mesh** (*nirfasterff.base.fluormesh*) – mesh used to calculate the source vectors. Source information is also defined here.
- **phix** (*double NumPy array*) – excitation fluence calculated at each node for each source. Size (NNodes, NSources)
- **frequency** (*double, optional*) – modulation frequency, in Hz. The default is 0..
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **GPU** (*int, optional*) – GPU selection. -1 for automatic, 0, 1, ... for manual selection on multi-GPU systems. The default is -1.

**Raises**

- **RuntimeError** – if both CUDA and CPU versions fail.
- **TypeError** – if ‘solver’ is not ‘CPU’ or ‘GPU’.

**Returns**

**qvec** – calculated fluence emission source vectors. Size (NNodes, NSources)

**Return type**

double or complex double NumPy array

### nirfasterff.math.get\_boundary\_data

nirfasterff.math.get\_boundary\_data(*mesh*, *phi*)

Calculates boundary data given the field data in mesh

The field data can be any of the supported type: fluence, TPSF, or moments

#### Parameters

- **mesh** (*nirfasterff mesh type*) – the mesh whose boundary and detectors are used for the calculation.
- **phi** (*double or complex double NumPy array*) – field data as calculated by one of the 'get\_field\_\*' solvers. Size (NNodes, NSources)

#### Returns

**data** – measured boundary data at each channel. Size (NChannels,).

#### Return type

double or complex double NumPy array

### nirfasterff.math.get\_field\_CW

nirfasterff.math.get\_field\_CW(*csrI*, *csrJ*, *csrV*, *qvec*, *opt*=*utils.SolverOptions()*, *solver*=*utils.get\_solver()*)

Call the Preconditioned Conjugate Gradient solver with FSAI preconditioner. For CW data only.

The current Matlab version uses COO format input, so they are NOT directly compatible

If calculation fails on GPU (if chosen), it will generate a warning and automatically switch to CPU.

On GPU, the algorithm first tries to solve for all sources simultaneously, but this can fail due to insufficient GPU memory.

If this is the case, it will generate a warning and solve the sources one by one. The latter is not as fast, but requires much less memory.

On CPU, the algorithm only solves the sources one by one.

#### Parameters

- **csrI** (*int32 NumPy vector, zero-based*) – I indices of the MASS matrix, in CSR format.
- **csrJ** (*int32 NumPy vector, zero-based*) – J indices of the MASS matrix, in CSR format.
- **csrV** (*double NumPy vector*) – values of the MASS matrix, in CSR format.
- **qvec** (*double NumPy array, or Scipy CSC sparse matrix*) – The source vectors. i-th column corresponds to source i. Size (NNode, NSource)

See [gen\\_sources\(\)](#) for details.

- **solver** (*str, optional*) – Choose between 'CPU' or 'GPU' solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

#### Raises

- **TypeError** – if MASS matrix and source vectors are not both real, or if solver is not ‘CPU’ or ‘GPU’.
- **RuntimeError** – if both GPU and CPU solvers fail.

#### Returns

- **phi** (*double NumPy array*) – Calculated fluence at each source. Size (NNodes, Nsources)
- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.

See [ConvergenceInfo\(\)](#) for details

See also:

[gen\\_mass\\_matrix\(\)](#)

### nirfasterff.math.get\_field\_FD

`nirfasterff.math.get_field_FD(csrI, csrJ, csrV, qvec, opt=utils.SolverOptions(), solver=utils.get_solver())`

Call the Preconditioned BiConjugate Stabilized solver with FSAI preconditioner.

This is designed for FD data, but can also work for CW if an all-zero imaginary part is added to the MASS matrix and source vectors.

The current Matlab version uses COO format input, so they are NOT directly compatible

If calculation fails on GPU (if chosen), it will generate a warning and automatically switch to CPU.

On GPU, the algorithm first tries to solve for all sources simultaneously, but this can fail due to insufficient GPU memory.

If this is the case, it will generate a warning and solve the sources one by one. The latter is not as fast, but requires much less memory.

On CPU, the algorithm only solves the sources one by one.

#### Parameters

- **csrI** (*int32 NumPy vector, zero-based*) – I indices of the MASS matrix, in CSR format.
- **csrJ** (*int32 NumPy vector, zero-based*) – J indices of the MASS matrix, in CSR format.
- **csrV** (*complex double NumPy vector*) – values of the MASS matrix, in CSR format.
- **qvec** (*complex double NumPy array, or Scipy CSC sparse matrix*) – The source vectors. i-th column corresponds to source i. Size (NNode, NSource)

See [gen\\_sources\(\)](#) for details.

- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

#### Raises

- **TypeError** – if MASS matrix and source vectors are not both complex, or if solver is not ‘CPU’ or ‘GPU’.

- **RuntimeError** – if both GPU and CPU solvers fail.

#### Returns

- **phi** (*complex double NumPy array*) – Calculated fluence at each source. Size (NNodes, Nsources)
- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.

See [ConvergenceInfo\(\)](#) for details

See also:

[gen\\_mass\\_matrix\(\)](#)

### nirfasterff.math.get\_field\_TR

```
nirfasterff.math.get_field_TR(csrI, csrJ, csrV, qvec, dt, max_step, opt=utils.SolverOptions(),
                             solver=utils.get_solver())
```

Call the Preconditioned Conjugate Gradient solver with FSAI preconditioner. Calculates TPSF data

NOT interchangeable with the current MATLAB version

If calculation fails on GPU (if chosen), it will generate a warning and automatically switch to CPU.

On both GPU and CPU, the algorithm solves the sources one by one

#### Parameters

- **csrI** (*int32 NumPy vector, zero-based*) – I indices of the MASS matrices, in CSR format.
- **csrJ** (*int32 NumPy vector, zero-based*) – J indices of the MASS matrices, in CSR format.
- **csrV** (*complex double NumPy vector*) – values of the MASS matrices, in CSR format.  
This is calculated using [gen\\_mass\\_matrix](#) with  $\omega=1$ . The real part coincides with  $K+C$ , and the imaginary part coincides with  $-M$ .

See references for details

- **qvec** (*double NumPy array, or Scipy CSC sparse matrix*) – The source vectors. i-th column corresponds to source i. Size (NNode, NSource)

See [gen\\_sources\(\)](#) for details.

- **dt** (*double*) – time step size, in seconds.
- **max\_step** (*int32*) – total number of time steps.
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

#### Raises

- **TypeError** – if **csrV** is not complex, or if **qvec** is not real, or if **solver** is not ‘CPU’ or ‘GPU’.
- **RuntimeError** – if both GPU and CPU solvers fail.

#### Returns

- **phi** (*double NumPy array*) – Calculated TPSF at each source. Size (NNodes, Nsources\*max\_step), structured as,  
[src0\_step0, src1\_step0,...,src0\_step1, src1\_step1,...]
- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.  
Only the convergence info of the last time step is returned.  
See [ConvergenceInfo\(\)](#) for details

See also:

[gen\\_mass\\_matrix\(\)](#)

## References

Arridges et al., Med. Phys., 1993. doi:10.1118/1.597069

## nirfasterff.math.get\_field\_TRFL

`nirfasterff.math.get_field_TRFL(csrI, csrJ, csrV, phix, dt, max_step, opt=utils.SolverOptions(), solver=utils.get_solver())`

Call the Preconditioned Conjugate Gradient solver with FSAI preconditioner. Calculates the TPSFs of fluorescence emission given the TPSFs of excitation

If calculation fails on GPU (if chosen), it will generate a warning and automatically switch to CPU.

On both GPU and CPU, the algorithm solves the sources one by one

### Parameters

- **csrI** (*int32 NumPy vector, zero-based*) – I indices of the MASS matrices at emission wavelength, in CSR format.
- **csrJ** (*int32 NumPy vector, zero-based*) – J indices of the MASS matrices at emission wavelength, in CSR format.
- **csrV** (*complex double NumPy vector*) – values of the MASS matrices at emission wavelength, in CSR format.

This is calculated using `gen_mass_matrix` with `omega=1`.

- **phix** (*double NumPy array*) – TPSF of the excitation. Size (NNodes, NSources\*NTime), structured as,  
[src0\_step0, src1\_step0,...,src0\_step1, src1\_step1,...]
- **dt** (*double*) – time step size, in seconds.
- **max\_step** (*int32*) – total number of time steps. It should match exactly with the number of steps in the excitation data
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** (*nirfasterff.utils.SolverOptions, optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

### Raises

- **TypeError** – if `csrV` is not complex, or if `phix` is not real, or if solver is not ‘CPU’ or ‘GPU’.
- **RuntimeError** – if both GPU and CPU solvers fail.

#### Returns

- **phi** (*double NumPy array*) – Calculated TPSF at each source of fluorescence emission. Size (NNodes, Nsources\*max\_step), structured as,  
[src0\_step0, src1\_step0, ..., src0\_step1, src1\_step1, ...]
- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.  
Only the convergence info of the last time step is returned.  
See [ConvergenceInfo\(\)](#) for details

See also:

[gen\\_mass\\_matrix\(\)](#), [get\\_field\\_TR\(\)](#)

#### nirfasterff.math.get\_field\_TRFLmoments

`nirfasterff.math.get_field_TRFLmoments(csrI, csrJ, csrV, mx, alpha, tau, max_moment, opt=utils.SolverOptions(), solver=utils.get_solver())`

Call the Preconditioned Conjugate Gradient solver with FSAI preconditioner. Directly calculates moments of re-emission using Mellin transform, given the moments of excitation

If calculation fails on GPU (if chosen), it will generate a warning and automatically switch to CPU.

On both GPU and CPU, the algorithm solves the sources one by one

#### Parameters

- **csrI** (*int32 NumPy vector, zero-based*) – I indices of the MASS matrices at emission wavelength, in CSR format.
- **csrJ** (*int32 NumPy vector, zero-based*) – J indices of the MASS matrices at emission wavelength, in CSR format.
- **csrV** (*complex double NumPy vector*) – values of the MASS matrices at emission wavelength, in CSR format.

This is calculated using `gen_mass_matrix` with `omega=1`.

- **mx** (*double NumPy array*) – moments of the excitation. Size (NNodes, Nsources\*(max\_moment+1)), structured as,  
[src0\_m0, src1\_m0, ..., src0\_m1, src1\_m1, ...]
- **alpha** (*double NumPy array*) – defined as `mesh.eta*mesh.muaf`.
- **tau** (*double NumPy array*) – decay factor, as defined in `mesh.tau`.
- **max\_moment** (*int32*) – max order of moments to calculate. That is, 0th, 1st, 2nd, ..., max\_moments-th will be calculated.

This should match exact with the `max_moment` of the excitation

- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.

See [SolverOptions\(\)](#) for details

#### Raises

- **TypeError** – if `csrV` is not complex, or if `mx` is not real, or if solver is not ‘CPU’ or ‘GPU’.
- **RuntimeError** – if both GPU and CPU solvers fail.

#### Returns

- **phi** (*double NumPy array*) – Calculated Mellin transform of fluorescence emission at each source. Size (NNodes, Nsources\*(max\_moment+1)), structured as,  
[src0\_m0, src1\_m0, ..., src0\_m1, src1\_m1, ...]

- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.

Only the convergence info of the highest order moments is returned.

See [ConvergenceInfo\(\)](#) for details

See also:

[gen\\_mass\\_matrix\(\)](#), [get\\_field\\_TRmoments\(\)](#)

### nirfasterff.math.get\_field\_TRmoments

`nirfasterff.math.get_field_TRmoments(csrI, csrJ, csrV, qvec, max_moment, opt=utils.SolverOptions(), solver=utils.get_solver())`

Call the Preconditioned Conjugate Gradient solver with FSAI preconditioner. Directly calculates moments of TR data using Mellin transform

NOT interchangeable with the current MATLAB version

If calculation fails on GPU (if chosen), it will generate a warning and automatically switch to CPU.

On both GPU and CPU, the algorithm solves the sources one by one

#### Parameters

- **csrI** (*int32 NumPy vector, zero-based*) – I indices of the MASS matrices, in CSR format.
- **csrJ** (*int32 NumPy vector, zero-based*) – J indices of the MASS matrices, in CSR format.

- **csrV** (*complex double NumPy vector*) – values of the MASS matrices, in CSR format.

This is calculated using `gen_mass_matrix` with `omega=1`. The real part coincides with `K+C`, and the imaginary part coincides with `-B`.

See references for details

- **qvec** (*double NumPy array, or Scipy CSC sparse matrix*) – The source vectors. *i*-th column corresponds to source *i*. Size (NNode, NSource)
- **max\_moment** (*int32*) – max order of moments to calculate. That is, 0th, 1st, 2nd, ..., max\_moments-th will be calculated.
- **solver** (*str, optional*) – Choose between ‘CPU’ or ‘GPU’ solver (case insensitive). Automatically determined (GPU prioritized) if not specified
- **opt** ([nirfasterff.utils.SolverOptions](#), *optional*) – Solver options. Uses default parameters if not specified, and they should suffice in most cases.



See [SolverOptions\(\)](#) for details

#### Raises

- **TypeError** – if `csrV` is not complex, or if `qvec` is not real, or if solver is not ‘CPU’ or ‘GPU’.
- **RuntimeError** – if both GPU and CPU solvers fail.

#### Returns

- **phi** (*double NumPy array*) – Calculated Mellin transform at each source. Size (NNodes, Nsources\*(max\_moment+1)), structured as,  
[src0\_m0, src1\_m0,...,src0\_m1, src1\_m1,...]

- **info** (*nirfasterff.utils.ConvergenceInfo*) – convergence information of the solver.

Only the convergence info of the highest order moments is returned.

See [ConvergenceInfo\(\)](#) for details

See also:

[gen\\_mass\\_matrix\(\)](#)

#### References

Arridge and Schweiger, Applied Optics, 1995. doi:10.1364/AO.34.002683

### 4.1.7 nirfasterff.meshing

Functions used for mesh generation and quality check

#### Modules

<a href="#">nirfasterff.meshing.auxiliary</a>	Auxiliary functions used for mesh quality check.
<a href="#">nirfasterff.meshing.meshutils</a>	Functions used for mesh generation and mesh quality check

#### nirfasterff.meshing.auxiliary

Auxiliary functions used for mesh quality check.

They are unlikely to become useful to an ordinary user, but still documented for completeness

Use with caution: no error checking mechanisms implemented

## Functions

<code>check_facearea(nodes, ele)</code>	Calculates the areas of each face, and check if they are close to zero
<code>check_tetrahedron_faces(ele)</code>	Check for faces shared by more than two tetrahedrons
<code>checkedges(ele)</code>	Check for orphan edges and edges shared by more than two triangles
<code>quality_triangle_radius(nodes, ele)</code>	Radius ratio: $2 \cdot \text{inradius} / \text{circumradius}$
<code>simpqual(nodes, ele)</code>	For each tetrahedron, calculates the didehedral angles and returns the smallest sine of them
<code>vector_vector_angle(u, v)</code>	Calculates vector-vector angles, in radian

### nirfasterff.meshing.auxiliary.check\_facearea

nirfasterff.meshing.auxiliary.**check\_facearea**(nodes, ele)

Calculates the areas of each face, and check if they are close to zero

Close to zero defined as  $1e6$  of the max span of the mesh

#### Parameters

- **nodes** (*double NumPy array*) – node locations of the mesh.
- **ele** (*int32 NumPy array*) – element list of the mesh, zero-based.

#### Returns

- **area** (*double NumPy vector*) – areas of each face. Size (NElements,)
- **zeroflag** (*bool NumPy vector*) – flags of whether the area is close to zero, for each face. Size (NElements,)

### nirfasterff.meshing.auxiliary.check\_tetrahedron\_faces

nirfasterff.meshing.auxiliary.**check\_tetrahedron\_faces**(ele)

Check for faces shared by more than two tetrahedrons

#### Parameters

**ele** (*int32 NumPy array*) – element list of the mesh, zero-based.

#### Returns

**flag** – 0 if no faulty faces found, and 2 if faces shared by more than two tetrahedrons are found.

#### Return type

int

**nirfasterff.meshing.auxiliary.checkededges****nirfasterff.meshing.auxiliary.checkededges**(*ele*)

Check for orphan edges and edges shared by more than two triangles

**Parameters****ele** (*int32 NumPy array*) – element list of the mesh, zero-based.**Returns****flag** – 0 if no errors found; 1 if edges shared by more than two triangles found; 2 if dangling edges found; 3 if both errors found.**Return type**

int

**nirfasterff.meshing.auxiliary.quality\_triangle\_radius****nirfasterff.meshing.auxiliary.quality\_triangle\_radius**(*nodes, ele*)Radius ratio:  $2 \cdot \text{inradius} / \text{circumradius}$ 

Value between 0 and 1. Equals 1 only when a triangle is equilateral

**Parameters**

- **nodes** (*double NumPy array*) – node locations of the mesh.
- **ele** (*int32 NumPy array*) – element list of the mesh, zero-based.

**Returns**

radius ratios for each triangle. Size (NElements,)

**Return type**

double NumPy vector

**References**[https://en.wikibooks.org/wiki/Trigonometry/Circles\\_and\\_Triangles/The\\_Incircle](https://en.wikibooks.org/wiki/Trigonometry/Circles_and_Triangles/The_Incircle)**nirfasterff.meshing.auxiliary.simpqual****nirfasterff.meshing.auxiliary.simpqual**(*nodes, ele*)

For each tetrahedron, calculates the dihedral angles and returns the smallest sine of them

**Parameters**

- **nodes** (*double NumPy array*) – node locations of the mesh.
- **ele** (*int32 NumPy array*) – element list of the mesh, zero-based.

**Returns**

smallest sine of the dihedral angles for each element. Size (NElements,)

**Return type**

double NumPy vector

## References

[https://en.wikipedia.org/wiki/Dihedral\\_angle](https://en.wikipedia.org/wiki/Dihedral_angle)

### nirfasterff.meshing.auxiliary.vector\_vector\_angle

`nirfasterff.meshing.auxiliary.vector_vector_angle(u, v)`

Calculates vector-vector angles, in radian

Each row of u, v is a vector, and the angles are calculated pairwise row by row

#### Parameters

- **u** (*double NumPy array*) – first set of vectors.
- **v** (*double NumPy array*) – second set of vectors.

#### Returns

pairwise vector-vector angles, in radian. Same number of rows as u and v

#### Return type

double NumPy vector

### nirfasterff.meshing.meshutils

Functions used for mesh generation and mesh quality check

## Functions

<code>CheckMesh2D</code> (elements, nodes[, base, verbose])	Main function that calculates and checks the quality of a 2D mesh
<code>CheckMesh3D</code> (elements, nodes[, base, verbose])	Main function that calculates and checks the quality of a 3D mesh, which can be either a solid or surface mesh
<code>RunCGALMeshGenerator</code> (mask[, opt])	Generate a tetrahedral mesh from a volume using CGAL 6.0.1 mesher, where different regions are labeled used a distinct integer.
<code>boundfaces</code> (nodes, elements[, base, renumber])	Finds the boundary faces of a 3D tetrahedral mesh
<code>checkmesh3d_solid</code> (ele, nodes[, verbose])	Calculates and returns the quality metrics of the tetrahedrons in a 3D tetrahedral mesh
<code>checkmesh3d_surface</code> (ele, nodes[, verbose])	Calculates and returns the quality metrics of the triangles in a 3D surface mesh

### nirfasterff.meshing.meshutils.CheckMesh2D

`nirfasterff.meshing.meshutils.CheckMesh2D(elements, nodes, base=1, verbose=False)`

Main function that calculates and checks the quality of a 2D mesh

#### Parameters

- **elements** (*int32 NumPy array*) – element list of the mesh, zero-based.
- **nodes** (*double NumPy array*) – node locations of the mesh, in mm.

- **base** (*int*, *optional*) – one- or zero-based indexing of the element list. Can be 1, or 0. The default is 1.
- **verbose** (*bool*, *optional*) – whether print the problematic elements to stdout, if any. The default is False.

**Raises**

**TypeError** – if elements and nodes do not define a valid 2D mesh.

**Returns**

- **flag** (*int*) – flags of mesh quality, set by bits: ‘b2b1b0’.  
b1 set if faulty edges found; b2 set if triangles with small area found
- **q\_radius\_ratio** (*double NumPy array*) – radius ratio of each triangle, defined as  $2 \cdot \text{inradius} / \text{circumradius}$ .
- **area** (*double NumPy array*) – area of each triangle in mesh.

**nirfasterff.meshing.meshutils.CheckMesh3D**

`nirfasterff.meshing.meshutils.CheckMesh3D(elements, nodes, base=1, verbose=False)`

Main function that calculates and checks the quality of a 3D mesh, which can be either a solid or surface mesh

If surface mesh, `checkmesh3d_surface()` is called

If solid mesh, `checkmesh3d_solid()` is first used, and `checkmesh3d_surface()` is subsequently used to check its outer surface

**Parameters**

- **ele** (*int32 NumPy array*) – element list of the mesh, zero-based.
- **nodes** (*double NumPy array*) – node locations of the mesh, in mm.
- **base** (*int*, *optional*) – one- or zero-based indexing of the element list. Can be 1, or 0. The default is 1.
- **verbose** (*bool*, *optional*) – whether print the problematic elements to stdout, if any. The default is False.

**Raises**

**TypeError** – if elements and nodes do not define a valid 3D mesh.

**Returns**

- **vol** (*double NumPy vector*) – Volume (for solid mesh) or area (for surface mesh) of each element.
- **vol\_ratio** (*double NumPy vector*) – volume ratio for each tetrahedron in a solid mesh. Returns scalar 0.0 in case of a surface mesh.
- **q\_area** (*double NumPy vector*) – area ratio for each triangle in a surface mesh. Returns scalar 0.0 in case of a solid mesh.
- **status\_solid** (*int*) – flags of solid mesh quality, set by bits: ‘b3b2b1b0’.  
b1 set if small volumes found; b2 set if small volume ratios found; b3 set if faces shared by more than two tetrahedrons found

- **status\_surface** (*int*) – flags of surface mesh quality, set by bits: ‘b3b2b1b0’.  
b1 set if edges shared by more than two triangles found; b2 set if dangling edges found; b3 triangles with small area found

See also:

`nirfasterff.meshing.checkmesh3d_solid()`, `nirfasterff.meshing.checkmesh3d_surface()`

### **nirfasterff.meshing.meshutils.RunCGALMeshGenerator**

`nirfasterff.meshing.meshutils.RunCGALMeshGenerator`(*mask*, *opt*=*utils.MeshingParams*())

Generate a tetrahedral mesh from a volume using CGAL 6.0.1 mesher, where different regions are labeled used a distinct integer.

Internally, the function makes a system call to the mesher binary, which can also be used standalone through the command line.

Also runs a pruning steps after the mesh generation, where nodes not referred to in the element list are removed.

#### **Parameters**

- **mask** (*uint8 NumPy array*) – 3D volumetric data defining the space to mesh. Regions defined by different integers. 0 is background.
- **opt** (*nirfasterff.utils.MeshingParams*, *optional*) – meshing parameters used. Default values will be used if not specified.

See `nirfasterff.utils.MeshingParams()` for details

#### **Returns**

- **ele** (*int NumPy array*) – element list calculated by the mesher, one-based. Last column indicates the region each element belongs to
- **nodes** (*double NumPy array*) – element list calculated by the mesher, in mm.

#### **References**

[https://doc.cgal.org/latest/Mesh\\_3/index.html#Chapter\\_3D\\_Mesh\\_Generation](https://doc.cgal.org/latest/Mesh_3/index.html#Chapter_3D_Mesh_Generation)

### **nirfasterff.meshing.meshutils.boundfaces**

`nirfasterff.meshing.meshutils.boundfaces`(*nodes*, *elements*, *base*=0, *renumber*=True)

Finds the boundary faces of a 3D tetrahedral mesh

#### **Parameters**

- **nodes** (*double NumPy array*) – node locations of the mesh. Size (NNodes, 3)
- **elements** (*NumPy array*) – element list of the mesh, can be either one-based or zero-based, which must be specified using the ‘base’ argument.
- **base** (*int*, *optional*) – one- or zero-based indexing of the element list. Can be 1, or 0. The default is 0.
- **renumber** (*bool*, *optional*) – whether renumber of the node indices in the extracted surface mesh. The default is True.

#### **Returns**

- **new\_faces** (*int32 NumPy array*) – list of boundary faces of the mesh. Base of indexing is consistent with input element list

If *renumber=True*, node indices are renumbered; if not, same node indices as in ‘elements’ are used

- **new\_points** (*double NumPy array*) – point locations of the boundary nodes.

If *renumber=True*, returns the subset of node locations that are on the surface; if not, it is the same as input *nodes*

### nirfasterff.meshing.meshutils.checkmesh3d\_solid

nirfasterff.meshing.meshutils.**checkmesh3d\_solid**(*ele, nodes, verbose=False*)

Calculates and returns the quality metrics of the tetrahedrons in a 3D tetrahedral mesh

Please consider using nirfasterff.meshing.CheckMesh3D() instead.

#### Parameters

- **ele** (*int32 NumPy array*) – element list of the mesh, zero-based.
- **nodes** (*double NumPy array*) – node locations of the mesh, in mm.
- **verbose** (*bool, optional*) – whether print the problematic tetrahedrons to stdout, if any. The default is False.

#### Raises

**TypeError** – if mesh is not 3D tetrahedral, or if element list uses undefined nodes.

#### Returns

- **vol** (*double NumPy vector*) – volume of each tetrahedron, mm<sup>3</sup>.
- **vol\_ratio** (*double NumPy vector*) – volume ratio, defined as the smallest sine of dihedral angles of each tetrahedron.
- **zeroflag** (*bool NumPy vector*) – flags of whether the volume of a tetrahedron is too small.
- **faceflag** (*bool*) – whether there are faces shared by more than two tetrahedrons.

### nirfasterff.meshing.meshutils.checkmesh3d\_surface

nirfasterff.meshing.meshutils.**checkmesh3d\_surface**(*ele, nodes, verbose=False*)

Calculates and returns the quality metrics of the triangles in a 3D surface mesh

Please consider using nirfasterff.meshing.CheckMesh3D() instead.

#### Parameters

- **ele** (*int32 NumPy array*) – element list of the mesh, zero-based.
- **nodes** (*double NumPy array*) – node locations of the mesh, in mm.
- **verbose** (*bool, optional*) – whether print the problematic triangles to stdout, if any. The default is False.

#### Raises

**TypeError** – if mesh is not 3D tetrahedral, or if element list uses undefined nodes.

#### Returns

- **q\_radius\_ratio** (*double NumPy vector*) – radius ratio of each triangle, defined as  $2 \cdot \text{inradius} / \text{circumradius}$ .
- **q\_area\_ratio** (*double NumPy vector*) – face area divided by ‘ideal area’ for each triangle, where ideal area is the area of an equilateral triangle whose edge length equals the longest edge in the face.
- **area** (*double NumPy vector*) – area of each triangle, in  $\text{mm}^2$ .
- **zeroflag** (*bool NumPy vector*) – flags whether the area a triangle is close to zero.
- **edgeflag** (*int*) – flag if any problematic edges. Flag set by bits ‘b1b0’:  
b1=1 if dangling edges found, b0=1 if there exist edges shared by more than two triangles

### 4.1.8 nirfasterff.utils

Utility functions and auxiliary classes frequently used in the package.

#### Functions

<code>boundary_attenuation(n_incidence[, ...])</code>	Calculate the boundary attenuation factor between two media.
<code>check_element_orientation_2d(ele, nodes)</code>	Make sure the 2D triangular elements are oriented counter clock wise.
<code>compress_coo(coo_idx, N)</code>	Convert COO indices to compressed.
<code>gen_intmat_impl(mesh, xgrid, ygrid, zgrid)</code>	YOU SHOULD NOT USE THIS FUNCTION DIRECTLY.
<code>get_solver()</code>	Get the default solver.
<code>isCUDA()</code>	Checks if system has a CUDA device with compute capability $\geq 5.2$
<code>pointLineDistance(A, B, p)</code>	Calculate the distance between a point and a line (defined by two points), and find the projection point
<code>pointLocation(mesh, pointlist)</code>	Similar to Matlab's <code>pointLocation</code> function, queries which elements in mesh the points belong to, and also calculate the barycentric coordinates.
<code>pointTriangleDistance(TRI, P)</code>	Calculate the distance between a point and a triangle (defined by three points), and find the projection point
<code>uncompress_coo(compressed_idx)</code>	Convert compressed indices to COO.

#### nirfasterff.utils.boundary\_attenuation

`nirfasterff.utils.boundary_attenuation(n_incidence, n_transmission=1.0, method='robin')`

Calculate the boundary attenuation factor between two media.

If vectors are used as inputs, they must have the same size and calculation is done for each pair

If `n_incidence` is a vector but `n_transmission` is a scalar, code assumes `n_transmission` to be the same for each value in `n_incidence`

##### Parameters

- **n\_incidence** (*double Numpy vector or scalar*) – refractive index of the medium within the boundary, e.g. a tissue.



- **n\_transmission** (*double Numpy vector or scalar, optional*) – refractive index of the medium outside of the boundary, e.g. air. The default is 1.0.
- **method** (*str, optional*) – boundary type, which can be,
  - 'robin' - internal reflectance derived from Fresnel's law
  - 'approx' - Groenhuis internal reflectance approximation ( $1.440n^{-2} + 0.710n^{-1} + 0.668 + 0.00636n$ )
  - 'exact' - exact internal reflectance (integrals of polarised reflectances, etc.)
 The default is 'robin'.

**Raises**

**ValueError** – if n\_incidence and n\_transmission are both vectors and have difference sizes, or if method is not of a recognized kind

**Returns**

**A** – calculated boundary attenuation factor.

**Return type**

double Numpy vector or scalar

**References**

Durduran et al, 2010, Rep. Prog. Phys. doi:10.1088/0034-4885/73/7/076701

**nirfasterff.utils.check\_element\_orientation\_2d**

nirfasterff.utils.**check\_element\_orientation\_2d**(*ele, nodes*)

Make sure the 2D triangular elements are oriented counter clock wise.

This is a direct translation from the Matlab version.

**Parameters**

- **ele** (*NumPy array*) – Elements in a 2D mesh. One-based. Size: (NNodes, 3).
- **nodes** (*NumPy array*) – Node locations in a 2D mesh. Size: (NNodes, 2).

**Raises**

**TypeError** – If ele does not have three rows, i.e. not a 2D triangular mesh.

**Returns**

**ele** – Re-oriented element list.

**Return type**

NumPy array

**nirfasterff.utils.compress\_coo****nirfasterff.utils.compress\_coo**(*coo\_idx*, *N*)

Convert COO indices to compressed.

**Parameters**

- **coo\_idx** (*int NumPy array*) – Input indices in COO format, zero-based.
- **N** (*int*) – Number of rows in the sparse matrix.

**Returns**

Output indices in compressed format, zero-based. Size (N+1,)

**Return type**

int NumPy array

**nirfasterff.utils.gen\_intmat\_impl****nirfasterff.utils.gen\_intmat\_impl**(*mesh*, *xgrid*, *ygrid*, *zgrid*)

YOU SHOULD NOT USE THIS FUNCTION DIRECTLY. USE MESH.GEN\_INTMAT INSTEAD.

Heart of the gen\_intmat function, which calculates the necessary information for converting between mesh and grid space

**Parameters**

- **mesh** (*nirfasterff.base.stndmesh or nirfasterff.base.fluormesh*) – The original mesh with with FEM data is calculated.
- **xgrid** (*float64 NumPy 1-D array*) – x grid in mm. Must be regular.
- **ygrid** (*float64 NumPy 1-D array*) – y grid in mm. Must be regular.
- **zgrid** (*float64 NumPy 1-D array, or [] if 2D mesh*) – z grid in mm. Must be regular.

**Returns**

- **gridinmesh** (*int32 NumPy array*) – Col 0: indices of grid points that are in the mesh; Col 1: indices of the elements the grid point is in. Flattened in ‘F’ order, one-based.
- **meshingrid** (*int32 NumPy array*) – Indices of mesh nodes that are in the grid. One-based.
- **int\_mat\_mesh2grid** (*CSC sparse matrix, float64*) – Sparse matrix converting data from mesh space to grid space. Size (NGrid, NNodes).
- **int\_mat\_mesh2grid** (*CSC sparse matrix, float64*) – Sparse matrix converting data from grid space to mesh space. Size (NNodes, NGrid).

**nirfasterff.utils.get\_solver****nirfasterff.utils.get\_solver**()

Get the default solver.

**Returns**

If isCUDA is true, returns ‘GPU’, otherwise ‘CPU’.

**Return type**

str

## nirfasterff.utils.isCUDA

### nirfasterff.utils.isCUDA()

Checks if system has a CUDA device with compute capability  $\geq 5.2$

On a Mac machine, it automatically returns False without checking

#### Returns

True if a CUDA device with compute capability  $\geq 5.2$  exists, False if not.

#### Return type

bool

## nirfasterff.utils.pointLineDistance

### nirfasterff.utils.pointLineDistance(A, B, p)

Calculate the distance between a point and a line (defined by two points), and find the projection point

This is a direct translation from the Matlab version

#### Parameters

- **A** (*NumPy array*) – first point on the line. Size (2,) or (3,)
- **B** (*NumPy array*) – second point on the line. Size (2,) or (3,)
- **p** (*NumPy array*) – point of query. Size (2,) or (3,)

#### Returns

- **dist** (*double*) – point-line distance.
- **point** (*NumPy array*) – projection point on the line.

## nirfasterff.utils.pointLocation

### nirfasterff.utils.pointLocation(mesh, pointlist)

Similar to Matlab's pointLocation function, queries which elements in mesh the points belong to, and also calculate the barycentric coordinates.

This is a wrapper of the C++ function pointLocation, which implements an AABB tree based on Darren Engwirda's findtria package

#### Parameters

- **mesh** (*NIRFASTer mesh*) – Can be any of the NIRFASTer mesh types (stnd, fluor, dcs). 2D or 3D.
- **pointlist** (*NumPy array*) – A list of points to query. Shape (N, dim), where N is number of points.

#### Returns

- **ind** (*double NumPy array*) – i-th queried point is in element *ind[i]* of mesh (zero-based). If not in mesh, *ind[i]* = -1. Size: (N,).
- **int\_func** (*double NumPy array*) – i-th row is the barycentric coordinates of i-th queried point. If not in mesh, corresponding row is all zero. Size: (N, dim+1).

## References

<https://github.com/dengwirda/find-tria>

### nirfasterff.utils.pointTriangleDistance

nirfasterff.utils.**pointTriangleDistance**(*TRI, P*)

Calculate the distance between a point and a triangle (defined by three points), and find the projection point

#### Parameters

- **TRI** (*Numpy array*) – The three points (per row) defining the triangle. Size: (3,3)
- **P** (*Numpy array*) – point of query. Size (3,).

#### Returns

- **dist** (*double*) – point-triangle distance.
- **PP0** (*NumPy array*) – projection point on the triangular face.

## Notes

This is modified from Joshua Shaffer's code, available at: <https://gist.github.com/joshuashaffer/99d58e4ccbd37ca5d96e>

which is based on Gwendolyn Fischer's Matlab code: <https://uk.mathworks.com/matlabcentral/fileexchange/22857-distance-between-a-point-and-a-triangle-in-3d>

### nirfasterff.utils.uncompress\_coo

nirfasterff.utils.**uncompress\_coo**(*compressed\_idx*)

Convert compressed indices to COO.

#### Parameters

**compressed\_idx** (*int NumPy array*) – Input indices in compressed format, zero-based.

#### Returns

**coo\_idx** – Output indices in COO format, zero-based.

#### Return type

int NumPy array

## Classes

<i>ConvergenceInfo</i> ([info])	Convergence information of the FEM solvers.
<i>MeshingParams</i> ([xPixelSpacing, ...])	Parameters to be used by the CGAL mesher.
<i>SolverOptions</i> ([max_iter, AbsoluteTolerance, ...])	Parameters used by the FEM solvers, Equivalent to 'solver_options' in the Matlab version

**nirfasterff.utils.ConvergenceInfo**

**class** nirfasterff.utils.**ConvergenceInfo**(*info=None*)

Bases: object

Convergence information of the FEM solvers. Only used internally as a return type of functions nirfasterff.math.get\_field\_\*

Constructed using the output of the internal C++ functions

**isConverged**

if solver converged to relative tolerance, for each rhs

**Type**

bool array

**isConvergedToAbsoluteTolerance**

if solver converged to absolute tolerance, for each rhs

**Type**

bool array

**iteration**

iterations taken to converge, for each rhs

**Type**

int array

**residual**

final residual, for each rhs

**Type**

double array

**\_\_init\_\_**(*info=None*)

**Methods**

```
__init__([info])
```

**nirfasterff.utils.MeshingParams**

**class** nirfasterff.utils.**MeshingParams**(*xPixelSpacing=1., yPixelSpacing=1., SliceThickness=1., facet\_angle=25., facet\_size=3., facet\_distance=2., cell\_radius\_edge=3., general\_cell\_size=3., subdomain=np.array([0., 0.]), lloyd\_smooth=True, offset=None*)

Bases: object

Parameters to be used by the CGAL mesher. Note: they should all be double

**xPixelSpacing**

voxel distance in x direction. Default: 1.0

**Type**

double

**yPixelSpacing**

voxel distance in y direction. Default: 1.0

**Type**

double

**SliceThickness**

voxel distance in z direction. Default: 1.0

**Type**

double

**facet\_angle**

lower bound for the angle (in degrees) of surface facets. Default: 25.0

**Type**

double

**facet\_size**

upper bound for the radii of surface Delaunay balls circumscribing the facets. Default: 3.0

**Type**

double

**facet\_distance**

upper bound for the distance between the circumcenter of a surface facet and the center of its surface Delaunay ball. Default: 2.0

**Type**

double

**cell\_radius\_edge**

upper bound for the ratio between the circumradius of a mesh tetrahedron and its shortest edge. Default: 3.0

**Type**

double

**general\_cell\_size**

upper bound on the circumradii of the mesh tetrahedra, when no region-specific parameters (see below) are provided. Default: 3.0

**Type**

double

**subdomain**

Specify cell size for each region, in format:

```
[region_label1, cell_size1]
[region_label2, cell_size2]
...
```

If a region is not specified, value in “general\_cell\_size” will be used. Default: np.array([0., 0.])

**Type**

double Numpy array

**lloyd\_smooth**

Switch for Lloyd smoother before local optimization. This can take up to 120s (hard limit set) but improves mesh quality. Default: True

**Type**  
bool

#### offset

offset value to be added to the nodes after meshing. Size (3,). Default: None

**Type**  
double Numpy array

#### Notes

Refer to CGAL documentation for details of the meshing algorithm as well as its parameters

[https://doc.cgal.org/latest/Mesh\\_3/index.html#Chapter\\_3D\\_Mesh\\_Generation](https://doc.cgal.org/latest/Mesh_3/index.html#Chapter_3D_Mesh_Generation)

```
__init__(xPixelSpacing=1., yPixelSpacing=1., SliceThickness=1., facet_angle=25., facet_size=3.,
         facet_distance=2., cell_radius_edge=3., general_cell_size=3., subdomain=np.array([0., 0.]),
         lloyd_smooth=True, offset=None)
```

#### Methods

```
__init__([xPixelSpacing, yPixelSpacing, ...])
```

### nirfasterff.utils.SolverOptions

```
class nirfasterff.utils.SolverOptions(max_iter=1000, AbsoluteTolerance=1e-12,
                                       RelativeTolerance=1e-12, divergence=1e8, GPU=-1)
```

Bases: object

Parameters used by the FEM solvers, Equivalent to ‘solver\_options’ in the Matlab version

**max\_iter**  
maximum number of iterations allowed. Default: 1000

**Type**  
int

**AbsoluteTolerance**  
Absolute tolerance for convergence. Default: 1e-12

**Type**  
double

**RelativeTolerance**  
Relative (to the initial residual norm) tolerance for convergence. Default: 1e-12

**Type**  
double

**divergence**  
Stop the solver when residual norm greater than this value. Default: 1e8

**Type**  
double

**GPU**

GPU selection. -1 for automatic, 0, 1, ... for manual selection on multi-GPU systems. Default: -1

**Type**

int

`__init__(max_iter=1000, AbsoluteTolerance=1e-12, RelativeTolerance=1e-12, divergence=1e8, GPU=-1)`

**Methods**

---

`__init__([max_iter, AbsoluteTolerance, ...])`

---

### 4.1.9 nirfasterff.visualize

Functions for basic data visualization

**Functions**

<code>plot3dmesh(mesh[, data, selector, alpha])</code>	Fast preview of data within a 3D FEM mesh in the nirfasterff format.
<code>plotimage(mesh[, data])</code>	Fast preview of data within a 2D FEM mesh in the nirfasterff format.

**nirfasterff.visualize.plot3dmesh**

`nirfasterff.visualize.plot3dmesh(mesh, data=None, selector=None, alpha=0.8)`

Fast preview of data within a 3D FEM mesh in the nirfasterff format.

Plots an image of the values on the mesh at the intersection specified by “selector”.

For 2D mesh, use `plotimage()` instead

**Parameters**

- **mesh** (*nirfasterff mesh type*) – a 3D nirfasterff mesh to plot the data on.
- **data** (*double NumPy vector, optional*) – data to be plotted, with size (NNode,). If not specified, treated as all zero.
- **selector** (*str, optional*) – Specifies the intersection at which the data will be plotted, e.g. ‘x>50’, or ‘(x>50) | (y<100)’, or ‘x + y + z < 200’.

Note that “=” is not supported. When “|” or “&” are used, make sure that all conditions are put in parantheses separately

If not specified, function plots the outermost shell of the mesh.

- **alpha** (*float, optional*) – transparency, between 0-1. Default is 0.8

**Raises**

**TypeError** – if mesh not 2D.

**Returns**



- *matplotlib.figure.Figure* – the figure to be displayed
- *mpl\_toolkits.mplot3d.axes3d.Axes3D* – Current axes of the plot. Can be subsequently used for further plotting.

### Notes

This function is adapted from the ‘plotmesh’ function in the iso2mesh toolbox

<https://iso2mesh.sourceforge.net/cgi-bin/index.cgi>

### nirfasterff.visualize.plotimage

`nirfasterff.visualize.plotimage(mesh, data=None)`

Fast preview of data within a 2D FEM mesh in the nirfasterff format.

Plots an image of the values on the mesh. For 3D mesh, use `plot3dmesh()` instead

#### Parameters

- **mesh** (*nirfasterff mesh type*) – a 2D nirfasterff mesh to plot the data on.
- **data** (*double NumPy vector, optional*) – data to be plotted, with size (NNode,). If not specified, treated as all zero.

#### Raises

**TypeError** – if mesh not 2D.

#### Returns

- *matplotlib.figure.Figure* – the figure to be displayed
- *mpl\_toolkits.mplot3d.axes3d.Axes3D* – Current axes of the plot. Can be subsequently used for further plotting.



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