## NeuroField User Manual

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NeuroField is a C++ program (accompanied with helper scripts) that solves the neural field model of Robinson et al., where each of the simultaneous equations are handled by an object:

$$P=
u_{ab}\phi_{ab},$$
 Couple  $D_{ab}V_{ab}=P,$  Dendrite  $Q_a=S_aigl[\sum_b V_{ab}igr],$  QResponse  $\mathcal{D}_{ab}\phi_{ab}=Q_b.$ 

NeuroField generalizes the neural field theory by allowing users to:

- 1. Specify an arbitrary population model: an arbitrary number of populations and connections them may be specified;
- 2. Choose different types of populations, including neural or stimulus populations. For each neural population, the type of firing response, dendritic response type may be specified.
- 3. Choose different types of connections, including the type of axonal propagation, and synaptic coupling.
- 4. For each object, specify the parameter values.

This users guide covers the obtaining and setting up (Sec. 1), configuring (Sec. 3) and launching of NeuroField (Sec. 2), as well as postprocessing (Sec. 4) and tips and tricks (Sec. 5).

Within this documentation, specific terminology as appeared in the computer is in typewriter font. Commands are denoted as

Command to put in computer

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# 1 Obtaining and setting up NeuroField

# 1.1 Obtaining NeuroField

The code for NeuroField is managed by the version control system git, and is presently hosted on GitHub together with documentation, bug reports and feature requests. To obtain access to the repository, please send an email to Romesh Abeysuriya (r.abeysuriya@physics.usyd.edu.au).

To set up the latest version of NeuroField, execute

```
git clone git@github.com:RomeshA/neurofield.git
```

after having added your SSH key to your GitHub account. Alternatively, you can install the GitHub desktop client and click the 'Clone in Desktop' on the GitHub website. For additional instructions, check the GitHub tutorials for downloading repositories - the NeuroField repository can be downloaded following GitHub standard procedures.

## 1.2 Directory layout

Within the main directory, the user can find:

\*.h, \*.cpp C++ source code.

Configs/ Stores configuration files for NeuroField.

Documentation/ Contains documentations Documentation/user.pdf and

Documentation/developer.pdf . Running make doc generates these doc-

umentations.

Helper\_scripts/ Stores helper scripts, including plotting routines and other post-processing of

data procedures.

Launch A launcher script that handles compilation and launching of NeuroField,

capable of automating parameter sweeps and submitting jobs on the Complex

Systems cluster yossarian.

Output/ When using the launcher script to sweep over parameters, the launcher script

produces this directory, which stores all output files neurofield.\* in indepen-

dent subdirectories.

Release/ All compiled files, including the object files and the NeuroField executable is

stored here. This directory will be deleted by make clean, so user data should

not be stored here.

Test/ Directory for unit testing and is irrelevant for users.

Scrap/ Directory containing development scraps and is irrelevant for users.

neurofield.\* Output files generated by NeuroField. See Sec. 4.

## 1.3 Compiling NeuroField

You can compile NeuroField simply by running

make

from the root directory containing all of the source files. The compiler command is specified in Makefile and should be edited if you wish to use a different compiler, or if your compiler does not support some of the compiler flags.

If you are compiling on Windows, the suggested route is to cross-compile using MinGW on a Unix-like system, and then copy across any missing DLLs from the Unix system into the same directory as the executable file on the Windows machine. Compiling with the Visual C++ compiler has not been tested.

# 2 Launching NeuroField

The Launch script is used to compile and launch the NeuroField program. Users are encouraged to use this script rather than calling the NeuroField executable directly. To launch NeuroField, do:

- 1. Edit Makefile: Identify the platform to run NeuroField, and comment/uncomment the appropriate COMP directions. Generally, this step is not needed, but users are encouraged to check.
- 2. To execute with only one set of parameters, edit your configuration file in ./Configs , then run

```
./Launch Configs/config_file
```

For example,

```
./Launch Configs/cortex.conf
```

The output will be stored in the current directory.<sup>1</sup>

3. To ease parameter exploration, the launcher script is capable of doing parameter sweeps. The launcher script supports sweeping over an arbitrary number of objects, parameters and runs. For example, if the user wants to launch NeuroField 3 times, with parameters varying according to the following table,

Object(s)	Parameter	$1^{\rm st}$ run	$2^{\rm nd}$ run	$3^{\rm rd}$ run
Propag 1 , Propag 2	gamma	10	20	30
Couple 1	nu	1	2	3

simply list the above table entries into the argument list:

```
./Launch Configs/cortex.conf 'Propag 1' 'Propag 2' gamma 10 20 30 'Couple 1' nu 1 2 3
```

In terms of syntactic restriction, each row in the table can have an arbitary number of objects, but only one parameter; every row must have the same number of runs.<sup>2</sup>

- 4. Switches are accepted for choosing options. To turn on the switches, put them in the argument list to the launcher script. The following switches are accepted:
  - --restart run NeuroField in restart mode.
  - -i specify an configuration file name.
  - -o specify an output file name.
  - -h prints out a list of these switches.

<sup>&</sup>lt;sup>1</sup>Tips for vi users: you can launch NeuroField in vi with :!./Launch % [optional params] 
<sup>2</sup>Tips: the UNIX seq program allows shortening parameter value listing from 10 20 30 40 50 60 into 'seq 10 10 60', e.g. Launch Configs/cortex.conf 'Propag 1' gamma 'seq 10 10 60'

- 5. In case the script is executed on computer cluster yossarian, the launcher script tries to find a file called pbs. If no such file is found, the user would be prompted for a job name, expected computational time and email to generate pbs. This file is then submitted to the PBS system.
- 6. To clean up the directory, delete or store your neurofield.\* output files and the Output/ directory and subdirectories. Then run

```
make clean
```

to delete ./Release/ and the files LATEX produced in Documentation/.

7. This documentation can be generated by running

```
make doc
```

which generates ./Documentation/user.pdf and ./Documentation/developer.pdf . make clean deletes the files (excluding the .pdf files) created by LATEX.

# 3 Writing a configuration file

NeuroField allows an arbitrary number of populations and connections between them, with all objects taking arbitrary parameter values. These are all configured via a configuration file. This section documents the specifications of configuration files, where we use Configs/example.conf as an illustrative example.

To write a configuration file, a user can follow these steps:

- 1. Determine your population model by drawing a schematic diagram, thereby constructing a connection matrix. An example is shown in Fig. 1.
- 2. Look up existing configuration files in <code>Configs/</code>. By checking the comment located at the top of a configuration file, and also the connection matrix, a user should find the most suitable existing file to construct his own. This is less tedious (and less error prone) than writing a new one from scratch.
- 3. Specify the global parameters and connectivity matrix (Sec. 3.1).
- 4. Specify all populations (Sec. 3.2).
- 5. Specify all propagators (Sec. 3.3).
- 6. Specify all couples (Sec. 3.4).
- 7. Specify all output requests (Sec. 3.5).

General rules on the entries within a configuration file:

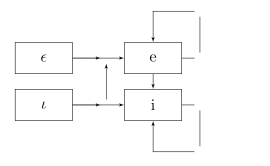
- 1. The structure of a configuration file is: 1) comment 2) global information 3) object specification 4) output specification.
- 2. Object specification involves first specifying the object type. The syntax is the object identifier, followed by its type, then a hyphen:

```
Object 1: Type -
```

Then the object parameters are specified, following this pattern:

```
Object parameter: value
```

- 3. Most parameters are essential. Failure to provide these parameters would result in NeuroField terminating with an error message. A minority of the parameters are optional.
- 4. The ordering of the parameters are important. Wrong parameter ordering results in NeuroField terminating with an error message.
- 5. The configuration file is white-space independent, e.g., there can be either no spaces, many spaces, or new lines between parameters.
- 6. For readability, users are encouraged to arrange parameter entries for different objects (via new lines and indentations) and aligning corresponding parameters between different objects.
- 7. Tip for vi users: ./helper\_scripts/neurofield.vim implements syntax highlighting for configuration files in vi . See comments within for installation instructions.



From:  $\epsilon$   $\iota$  e i To  $\epsilon$ : 0 0 0 0 0 To  $\iota$ : 0 0 0 0 0 To e: 1 0 2 3 To i: 0 4 5 6

Figure 1: Left: schematic diagram of a purely cortical population model comprising excitatory and inhibitory populations, as well as two stimulus populations; each arrow indicates a connection between populations, so that each stimulus connects to a cortical population, and each cortical population connects to all cortical populations. Right: connection matrix indicating the connections between populations; zero indicates no connection, and a connection is indicated by a nonzero number, ordered top to bottom, left to right.

```
Example config file of cortical model with excitatory and
   inhibitory neurons
Time: 1 Deltat: 1e-4
Nodes: 4 Longside: 2
Connection matrix:
From: 1 2 3 4
To 1: 0 0 0 0
To 2: 0 0 0 0
To 3: 1 0 2 3
To 4: 0 4 5 6
Population 1: Stimulation
Length: .5
Stimulus: Const - Onset: O Mean: 5
Population 2: Stimulation
Length: .5
 Stimulus: Const - Onset: O Mean: 5
Population 3: Excitatory neurons
Length: .5
Q: 8.87145
Firing: Sigmoid - Theta: 13e-3 Sigma: 3.8e-3 Qmax: 340
 Dendrite 1: V: Steady alpha: 83 beta: 769
 Dendrite 2: V: Steady alpha: 83 beta: 769
 Dendrite 3: V: Steady alpha: 83 beta: 769
Population 4: Inhibitory neurons
Length: .5
 Q: 8.87145
 Firing: Sigmoid - Theta: 13e-3 Sigma: 3.8e-3 Qmax: 340
 Dendrite 4: V: Steady alpha: 83 beta: 769
 Dendrite 5: V: Steady alpha: 83 beta: 769
 Dendrite 6: V: Steady alpha: 83 beta: 769
Propag 1: Map - phi: Steady Tau: 0
Propag 2: Wave - phi: Steady Tau: O Range: 80e-3 gamma: 116
Propag 3: Map - phi: Steady Tau: 0
Propag 4: Map - phi: Steady Tau: 0
Propag 5: Wave - phi: Steady Tau: O Range: 80e-3 gamma: 116
Propag 6: Map - phi: Steady Tau: 0
Couple 1: Map - nu: .15e-3
Couple 2: Map - nu: 1.5e-3
Couple 3: Map - nu:-1.8e-3
Couple 4: Map - nu: .15e-3
Couple 5: Map - nu: 1.5e-3
Couple 6: Map - nu:-1.8e-3
Output: Node: 1 2 Start: O Interval: 1e-4
Population: 4.V
                                  7
Dendrite: 5
Propag: 1 4.phi
Couple: 3.nu
```

### 3.1 Global information

Example config file of cortical model with excitatory and inhibitory neurons

Any text before the entry, Time: , is disregarded by NeuroField and serves as comment, which is strongly recommended for all configuration files.

```
Time: 10 Deltat: 1e-4
```

Time is the simulation duration in seconds.

Deltat is the time increment for each time step.

```
Nodes: 4 Longside: 2
```

Nodes is the number of grid points in the spatial dimension per population of neurons. The code has been explicitly designed to have equal number of neurons per population.

Longside is an optional parameter, specifying the longside of the rectangular grid. If it is not supplied, it is assumed to be a square.

Both spatial dimensions have periodic boundary conditions, so that populations have the topology of a torus.

```
Connection matrix:
From: 1 2 3 4
To 1: 0 0 0 0
To 2: 0 0 0 0
To 3: 1 0 2 3
To 4: 0 4 5 6
```

We specify an arbitrarily sized square connection matrix, where each entry is the connection from the column population to the row population.

Zero indicates no connection.

A nonzero number indicates connection. This number must be indexed from top to bottom, left to right.

## 3.2 Population data

This section contains population information sections. There are two types of neural populations: ordinary populations and stimulus populations:

### Stimulus populations

NeuroField identifies stimulus populations as populations which have no dendrites, i.e., the row for that population contains no nonzero elements. Each stimulus population information section is as follows.

```
Population 1: Stimulation
```

The identifier Population 1 is required for cross-checking.

The descriptor Stimulation is not parsed by NeuroField, but it is strongly recommended for human referencing.

```
Length: .5
```

The physical (1D) length of the population (which is a 2D sheet), in mitres. This is used in Wave propagators and the Psd of White stimulus.

```
Stimulus: Const - Onset: 0
```

The identifier Stimulus is required for cross-checking.

This is followed by the type of stimulus, to be further elaborated below.

Optional parameter Onset specifies the time onset for the stimulus to begin. If unspecified, stimulus starts at time 0.

Either Cease or Duration can be an optional parameter to specify the end time of the stimulus. If unspecified, stimulus ends at 1000 seconds.

If optional parameter Node is specified, only the specified node indices will receive stimulation.

Possible stimulus patterns:

#### Constant

```
Const - Mean: 5
```

### Pulse

```
Pulse - Amplitude: 1 Width: 2e-2 Frequency: 1 Pulses: 1
```

#### White

Gaussian noise, characterized by the mean and standard deviation:

```
White - Mean: 1 Std: 20 Ranseed: 10
```

Alternatively, the power spectral density (PSD) may be specified instead of the standard deviation. The advantage is that the PSD is invariant to change in <code>Deltat</code>, population <code>Length</code> and spatial <code>Nodes</code>. Given the PSD, <code>NeuroField</code> correctly calculates the standard deviation:

```
White - Mean: 1 Psd: 20 Ranseed: 10
```

In general, it is preferable to specify the noise using PSD rather than Std. The magnitude of nonlinear effects and the total power in the spectrum both depend on the PSD rather than the Std.

The random number generator may be specified in Ranseed. If a seed is not specified, an automatically-incremented seed will be used instead, so that multiple stimulus populations will have independent sequences. In general it is not necessary to set the seed manually unless different random numbers are required for otherwise identical runs.

### Superimposing stimuli

To superimpose 2 or more stimuli, begin with the keyword Superimpose, followed by the number of stimuli. Then list all the stimulus patterns and their parameters, with each stimulus pattern preceded by the keyword Stimulus.

```
Stimulus: Superimpose: 2
Stimulus: White - Mean: 1 Psd: 1
Stimulus: Pulse - Onset: 0.5 Width: 2e-2 Frequency 1
Pulses: 1
```

### **Ordinary populations**

Any non-stimulus population is an ordinary population.

```
Population 3: Excitory neurons
```

The identifier Population 3 is required for cross-checking.

The descriptor Excitatory neurons is not parsed by NeuroField, but it is strongly recommended for human referencing.

```
Q: 8.87145
```

The initial firing rate.

```
Firing: Sigmoid - Theta: 13e-3 Sigma: 3.8e-3 Qmax: 340
```

Specify the sigmoidal firing response of the population.

Sigma is sometimes known as  $\tilde{\sigma}$ . It is already scaled by  $\pi/\sqrt{3}$ .

Alternatively, you can specify a linear firing response by using

```
Firing: Linear - Gradient: 1 Intercept: 1
```

```
Dendrite 1: V: Steady alpha: 83 beta: 769
```

The identifier Dendrite 1, where the number 1 is the presynaptic connection index, is required for cross-checking. Users should find that these indices are simply ordered as 1, 2, 3, 4, ...

Optional parameter V may be used to specify the initial depolarization contribution from presynaptic activity. If unspecified, or set to Steady, NeuroField calculates the initial value by  $V_{ab} = \nu_{ab}\phi_{ab}$ .

alpha and beta are the parameters for the depolarization response.

## 3.3 Propagation data

Propag 1:

This identifier is required for cross-checking.

• A propagator type is required at this point. Choices are Map, Wave, and Harmonic.

### Map

Map - phi: Steady Tau: 0

This propagator is the mapping propagator where spatial spreading is negligible. Its form is given by

$$\phi_{ab}(\mathbf{r},t) = Q_b(\mathbf{r},t-\tau_{ab}).$$

Optional parameter, Tau, is the axonal delay term. If unspecified, it is taken as zero. Since all propagators contain this object, its description is given below.

Optional parameter phi may be used to specify the initial axonal firing rate. If unspecified, or set to Steady, NeuroField calculates the initial value by  $\phi_{ab} = Q_b$ .

#### Wave

This propagator is the wave equation propagator governed by the equation

$$\left[\frac{1}{\gamma_{ab}^2}\frac{d^2}{dt^2} + \frac{2}{\gamma_{ab}}\frac{d}{dt} + 1 - r_{ab}^2\nabla^2\right]\phi_{ab}(\mathbf{r}, t) = Q_b(\mathbf{r}, t - \tau_{ab}).$$

NeuroField checks whether the Courant condition must be satisfied, i.e.

$$\Delta t/\Delta x < \sqrt{2}/r_e \gamma_e$$

where  $\Delta x$  is the population length per node.

The propagator input is given by

Optional parameter phi may be used to specify the initial axonal firing rate. If unspecified, or set to Steady, NeuroField calculates the initial value by  $\phi_{ab} = Q_b$ . Range is  $r_{ab}$  in the wave equation.

gamma is the damping coefficient. Alternatively, velocity may be specified, and gamma is calculated via  $\gamma_{ab} = v_{ab}/r_{ab}$ .

In case there is only one node, this degenerates into a Harmonic propagator.

**Harmonic** This is a harmonic oscillator implementation of the damped wave equation, with no spatial variations:

$$\left[\frac{1}{\gamma_{ab}^2}\frac{d^2}{dt^2} + \frac{2}{\gamma_{ab}}\frac{d}{dt} + 1\right]\phi_{ab}(\mathbf{r}, t) = Q_b(\mathbf{r}, t - \tau_{ab}).$$

The input form is given by

```
Harmonic - phi: Steady Tau: O gamma: 116
```

#### Tau

The axonal time delay between populations. If it is spatially homogeneous, then it is a number with units of seconds. If it is spatially inhomogeneous, then input n numbers, where n =Nodes .

## 3.4 Coupling data

```
• Couple 1:
```

Identifier for cross-checking.

• A couple type is required at this point. Choices are Map, CaDP, BCM and Matrix.

### Map

Nonplastic synaptic coupling with a single constant parameter nu,

```
Map - nu: 0.0012
```

nu is the synaptic coupling parameter. It corresponds to the product of the mean synaptic strength  $s_{ab}$  and  $N_{ab}$ , the mean number of connections from cells of type b to cells of type a.

#### Matrix

Coupling becomes connection matrix, where connection strength does *not* change with time. The format of the nu matrix is the same as the population connection matrix, each row is to the same node, each column is from the same node. When outputting, each specified outputting node output the indexed row.

```
nu:
13e-6 0
0 13e-6
```

### CaDP

Calcium dependent plasticity according to Fung and Robinson.

```
CaDP - nu: 13e-6 nu_max: 80e-6 Dth: .25e-6 Pth: .45e-6 xyth: 1e-4 x: 2.3e-2 y: 2e-2 B: 30e3 glu_0: 200e-6 gNMDA: 2e-3
```

Dth and Pth are the calcium-plasticity thresholds; xyth, x and y are the plasticity rates; B, glu\_0 and gNMDA are NMDA receptor parameters.

To use CaDP, glutamate dynamics must be specified for the postsynaptic population. In the end of the relevant population entry, append

```
Glutamate dynamics - Lambda: 150e-6 tGlu: 30e-3
```

Lambda is the glutamate concentration rise per presynaptic spike; tGlu is the decay timescale for glutamate dynamics.

### BCM

Extends CaDP with metaplasticity according to Fung and Robinson. it has an additional parameter t\_BCM, the timescale of metaplasticity.

```
CaDP - nu: 13e-6 nu_max: 80e-6 Dth: .25e-6 Pth: .45e-6 xyth: 1e-4 x: 2.3e-2 y: 2e-2 B: 30e3 glu_0: 200e-6 gNMDA: 2e-3 t_BCM: 7
```

## 3.5 Output data

NeuroField outputs field quantities (i.e. a neurodynamic quantity which takes a value for each node) with respect to nodes and time. By default, the output file is neurofield.output, which can be changed by launching the program with the -o switch.

```
Output:
```

Begin with the Output declaration.

```
Node: 1 2
```

Enumerate all nodes to be outputted. If outputting all nodes, use shorthand All. If no nodes are specified, no nodes will be outputted.

```
Start: 0 Interval: 1e-4
```

Optional parameters for the time to start output, and optional parameter for time interval between outputs.

If undefined, defaults, to 0 and Deltat, respectively.

```
Population: 4.V
Dendrite: 5
Propag: 1 4.phi
Couple: 3.nu
```

NeuroField allows the user to specify which objects to output, by entering the appropriate object indices after the labels. For each object, it has some intrinsic fields that will be outputted; for example, Couple outputs nu, whereas CaDP outputs nu and Ca.

For each entry, a field name may be appended after the index with a dot, so that only that field of the object is outputted. If no field name is specified for that entry, then all fields of that object is outputted.

If a specific field of an object is specified, but that field does not exist, NeuroField checks and returns an error.

# 4 Postprocessing

NeuroField produces 3 files:

neurofield.conf When using the launcher script, this file is created to store the running configu-

ration file.

neurofield.output The result of the simulation is stored here for postprocessing.

neurofield.pbs If NeuroField is run in yossarian, then this file stores the output of the

queueing system.

When the launcher script runs with only one set of parameters, all output files are also in the present working directory. However, if the launcher script sweeps over parameters, each parameter set has its own subdirectory inside <code>Output/</code>, and each set of <code>neurofield.\*</code> files are stored in its subdirectory.

The output file starts with a copy of the input file, to enable the output file to serve as a complete representation of the simulation. The simulation results follow a series of = characters. Example content in neurofield.output is

Time	Propag.2.phi				
	0				
1.00000000000000e-04	8.87145000000000e+00				

Each column is a time series with its name indicated in the first line. The first column is always time, and in this example, the second column is Propag.2.phi, indicating that it is  $\phi_{ee}$  (when checked against connection matrix). The delimiter | indicates that the two columns are different fields, rather than different nodes of the same field. The node number is indicated in the second line.

It is also worth noting that traces will be written in the order that they are specified. For example, if you write Population: 3 1 then the columns in the output file will be arranged in this order.

# 4.1 Quickplot

The data in neurofield.output may be plotted by ./Helper\_script/quickplot.pl or in MatLab via the MatLab scripts within ./Helper\_script/.

To use ./Helper\_scriptquickplot.pl , execute

```
./Helper_script/quickplot.pl [output.file] [field] [node index]
```

where an example is

```
./Helper_script/quickplot.pl neurofield.output Propag.2.phi 1
```

or a shorthand to plot all fields and nodes is

```
./Helper_script/quickplot.pl all
```

All these commands launches gnuplot plotting sessions.

### 4.2 Matlab

A number of MatLab functions are provided to make it easy to manipulate NeuroField data from within MatLab. The functions are generally self-documenting with comments at the start of the file.

Essentially, an output file from NeuroField is read into a nf struct object in MatLab which simply contains all of the output from NeuroField in memory for easy access. Here is an example of a nf object:

```
fields: {'Propag.1.phi' 'Propag.3.phi'}
  nodes: {[1] [1]}
  data: {[300000x1 double] [300000x1 double]}
  time: [300000x1 double]
  deltat: 1.0000e-04
  npoints: 300000
```

- fields stores a record of which traces from NeuroField are present in the output file
- nodes is a cell with the same size as fields, and records for each field present, the number of the node in the output.
- data is a matrix storing the actual values of the traces
- time is a vector of time values, so that you can plot any of the data traces directly again nf.time
- deltat stores the temporal sampling rate
- npoints stores the total number of points in the output. The total duration is nf.deltat\*nf.npoints (or nf.time(end))

There are two ways to create the **nf** object. You can read the output file directly after executing NeuroField elsewhere

```
nf = nf_read('neurofield.output')
```

or you can use the nf\_run helper script to run the config file using NeuroField and automatically parse the output

```
nf = nf_run('neurofield.conf')
```

Several helper files are provided to manipulate the <code>nf</code> object. The two most important helpers are <code>nf\_extract</code> and <code>nf\_grid</code>. Often you want to extract a particular field from the <code>nf</code> object, for example, to examine the output from <code>Propag.3.phi</code>. To do this directly with the <code>nf</code> object, you would need to check the <code>fields</code> variable to find the index of the trace you wanted, and then extract it from the <code>data</code> field. In the previous example, <code>Propag.3.phi</code> is the second trace. These expressions are identical:

```
trace = nf.data{2};
trace = nf_extract(nf,'Propag.3.phi')
```

nf\_extract quickly becomes useful when there are many different fields. It is not case sensitive (so propag.3.phi works as well). You can also specify using additional arguments to extract only a portion of the time series, and also to select a subset of nodes. Finally, you can also provide multiple traces to concatenate them into a single matrix. For example,

```
trace = nf_extract(nf,'propag.1.phi,propag.3.phi');
```

will create a 300000×2 matrix with both of the traces.

Finally, if you run NeuroField with multiple nodes, it typically solves the system of equations on a square grid. Therefore, if you have output for notes 1-400, this corresponds to a 20×20 grid. nf\_grid allows you to request a trace from the nf object and have it reshaped into a square grid. This lets you easily make surface plots of the data, or perform tasks that are spatially dependent.

One important task is computing the power spectrum as predicted by the linearized analytic equations. This can be achieved using  $nf\_spatial\_spectrum$  which takes in an nf object and computes the power spectrum integrated over k taking into account volume conduction.

# 5 Tips and tricks