Documentation for networkFVMsims: finite volume method implementation of evolving reaction-diffusion fields on a network

E. F. Koslover

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The code in this package can be used to simulate dynamics of buffered calcium ions or of other particles spreading across a tubular network. In its default mode, it treats networks as one-dimensional edges of constant radius, connected at point-like nodes. Some extensions are provided for inputting 3D meshes of more complicated structures (eg: sheets), but these are not yet documented. Alternately, rapidly-equilibrating 'reservoirs' can be included in the networks.

The code stores information about network structure (edge connectivity, edge lengths) in a NETWORK object. Note that while network edges can be curved (and thus longer than the node-to-node distance), the spatial embedding of the edges is not stored. The network structure is input by the user in a .net file (see networktools project on github for Matlab code to manipulate, visualize, and output network structures). The current code builds its own MESH structure consisting of linked mesh elements. These can be loaded and visualized with the matlab code provided in scripts.

NOTE: this documentation is far from complete and currently only includes the most basic of the features implemented.

1 Dependencies

- This project code: https://github.com/lenafabr/networkSpreadFVM
- Network manipulation code: https://github.com/lenafabr/networktools

2 Compilation Instructions

To compile and run the program, you will need the following:

- a compiler capable of handling Fortran 90. The code has been tested with the gfortran compiler.
- Optionally: Matlab to visualize output data

The code has been tested on Ubuntu Linux 20.04.

To compile with gfortran, go into the source directory. Type make. To compile with any other compiler that can handle Fortran 90, type

make FC=compiler

substituting in the command you usually use to call the compiler.

If the compilation works properly, the executable netmeshdynamicsFVM.exe will appear in the main directory.

3 Usage Instructions

To run the program in the main directory, type:

./netmeshdynamicsFVM.exe suffix > stdout.suffix

Here, suffix can be any string up to 100 characters in length. The program reads in all input information from a file named param.suffix where suffix is the command-line argument. If no argument is supplied, it will look for a file named param. If the desired parameter file does not exist, the program will exit with an error.

The parameters in the input file are given in the format "KEYWORD value" where the possible keywords and values are described in Section 5. Each keyword goes on a separate line. Any line that starts with "#" is treated as a comment and ignored. Any blank line is also ignored. The keywords in the parameter file are not case sensitive. For the most part, the order in which the keywords are given does not matter. Most of the parameters have default values, so you need only specify keywords and values when you want to change something from the default.

4 Example for a Quick Start

A few example parameter files are provided

4.1 Ca²⁺ release from a honeycomb network

The parameters for this example run are stored in examples/param.example1_ca). Within the examples directory, run the code as:

../netmeshdynamicsFVM.exe example1_ca

This will run a simulation of calcium ions and buffer proteins diffusing on a tubular network, with a local region that is releasing free calcium.

Output files:

- example1_ca.out contains the total flux of calcium out of the network at each time point
- example1_ca.snap.txt contains the concentration of free calcium and total buffer protein at all mesh cells, at each snapshot time

To parse this data and generate plots, an example matlab script is provided in scripts/example1_ca.m | TODO: insert example output figure here|

4.2 Spreading of photoactivated particles

The parameters for this example run are stored in examples/param.example_PA). Within the examples directory, run the code as:

../netmeshdynamicsFVM.exe example_PA

This will run a simulation of diffusive particles on a network spreading from an initial bolus of photoactivated particles.

Output files:

- example_PA.out contains the total flux of calcium out of the network at each time point
- example_PA.snap.txt contains the concentration of free calcium and total buffer protein at all mesh cells, at each snapshot time

To parse this data and generate plots, an example matlab script is provided in scripts/example_PA.m [TODO: insert example output figure here]

4.3 Continuous photoactivation

[TODO: make an example simulation with a continuously photoactivated region]

4.4 Ca²⁺ refill

[TODO: make an example simulation where a reservoir is refilled with calcium]

5 Keyword Index

The code will attempt to read parameters out of a file named param.suffix where "suffix" is the command line argument. If no command line arguments are supplied, it will look for a file named param. If multiple arguments are supplied, it will read multiple parameter files in sequence.

The parameter file should have one keyword per line and must end with a blank line. All blank lines and all lines beginning with # are ignored. For the most part, the order of the lines and the capitalization of the keywords does not matter. All keywords except ACTION are optional. The default values for each parameter are listed below. If a keyword is supplied, then values may or may not be needed as well. The required and optional value types are listed below.

Keywords and multiple values are separated by spaces.

When reading the parameter file, lines longer than 500 characters will be truncated. To avoid this and continue onto the next line, add "+++" at the end of the line to be continued. No individual keyword or value should be longer than 100 characters.

Floating point numbers can be formated as 1.0, 1.1D0, 10e-1, -1.0E+01, etc., where the exponential notation specifier must be D or E (case insensitive). Integer numbers can also be specified in exponential notation without decimal points (eg: 1000 or 1E3). Logical values can be specified as T, F, TRUE, FALSE, 1, or 0 (with 1 corresponding to true and 0 to false).

List of Keyword inputs

[This is currently a very incomplete list. Some lucky student is going to get the task of filling in all the implemented keywords eventually.] In the meantime, look in source/readkey.f90 for comments and default values of most of the keyword parameters. The variables set by keywords are declared in source/keys.f90

By default, the simulations work with concentrations in units of $mM \times \pi a^2$ where a is the tubule radius, length units of μ m, and time units of seconds.

• ACTION

- value: 1 string of at most 20 characters; default NONE
- This keyword sets the overall calculation performed by the program
- Possible values are: RUNDYNAMICS (the only one currently implemented)

• DELT

- value: 1 float; default 1D-4
- Time-step for forward Euler stepping in the dynamics.
- Not used

• DOFLOW

- value: 1 optional logical; default: true
- Include advective flow along edges

• GLOBALRESERVOIR

- values: 5 floats, 1 logical; defaults: -, -, -, -, F
- Include a special 'global reservoir' that interacts with all (or most) of the mesh cells.
 By default, there is no such reservoir.
- This reservoir gets its own concentration fields (stored as part of the mesh structure)
- Interchange between the global reservoir and the mesh cells is in terms of Michaelis-Menten kinetics and is not based on transport coefficients (ie: unrelated to diffusion or flow across boundaries). Assume only the first field interacts with the global reservoir
- The values supplied (in order) are:
 - * Volume V_g . If working with 1D concentrations, instead supply $V_g^{(1D)}/(pia^2)$ where a is the tubule radius.
 - * Recovery rate constant k_r in units of per area per time. If working with 1D concentrations, instead supply $k_r^{(1D)}(2\pi a)$
 - * K_{Mr} = saturation concentration for recovery.
 - * Rate constant k_{out} for pumping out of the global reservoir. Units of time⁻¹.
 - * $K_{M,\text{out}}$ = saturation concentration for pumping out.

* PERMTOGLOBALRES. If set to True, permeable nodes release their particles into the global reservoir rather than the extracellular environment.

• MESHFILE

- value: 1 string, up to 100 characters; default: *.mesh.txt
- File name in which to output mesh structure.
- * is replaced by the command-line suffix

• MESHSIZE

- values: 1 float, 1 optional integer; defaults: 0.1, 2
- Float: limit for maximum length of an edge mesh cell
- Integer: minimum number of internal mesh cells along an edge (does not include nodes). Gives smaller mesh cells when edges are short.

• NETWORKDIM

- value: 1 integer; default: 0
- Dimensionality of space in which network is embedded
- If positive value: explicitly defines the spatial dimension in which the network is embedded
- If 0, use the .net file (# items in a NODE row 2) to set the dimension

• NETFILE

- value: 1 string; default: *.net
- Input network file used to start the simulation
- Any * in the file name will be replaced by the command-line argument (suffix)

• OUTFILE

- value: 1 string (up to 100 characters) and 1 optional integer
- default: *.out 1000
- String: output file for the flux of released calcium over time
- Any * in the file name will be replaced by the command-line argument (suffix)
- The integer N indicates the flux will be written to the file every N timesteps of the simulation. This can also be set separately with keyword OUTPUTEVERY

• PRINTEVERY

- value: 1 integer; default: 1
- Print simulation progress to screen every so many steps

• PERMNODE

- value: 1 integer n, 1 or more floats p; defaults: 0
- Make a specific node permeable to one or more of the concentration fields. By default, there are no permeable nodes.
- -n is the node index. p is the permeability or permeability prefactor, scaled as described below, for each of the fields.
- If P is the membrane permeability in units of length/time, the current out of each mesh cell is $I = AP(c_i c_{\text{ext}}) = \frac{AP}{\pi a^2}(\rho_i \rho_{\text{ext}})$ where c_i are 3D concentrations and ρ_i are 1D concentrations. A_i is the mesh cell surface area.
- If working with 1D concentrations (the default) then:
 - * If PERMPREFACTOR is true, then p = 2P/(aD) where D is the particle diffusivity, a is the tubule radius. The current is $I = [p(\frac{A}{2\pi a})D](\rho_i \rho_{\text{ext}})$
 - * If PERMPREFACTOR is false, then $p = AP/(\pi a^2)$ where A is the area of a mesh cell (in units of length², assumed the same for all mesh cells). The current is $p(\rho_i \rho_{\text{ext}})$
- If working with 3D concentrations then:
 - * If PERMPREFACTOR is true, then p = P/D. The current is $I = [pAD](c_i c_{\text{ext}})$
 - * If PERMPREFACTOR is false, then p = AP. The current is $I = [p](c_i c_{\text{ext}})$.

• PERMPREFACTOR

- value: 1 optional logical
- If this is true, then the permeability value set by PERMNODE is treated as a prefactor, to be scaled by $A \cdot D$ when incorporated into simulations
- If keyword is not supplied, it is set to F
- If keyword is supplied but there is no value, it is set to T
- Strongly recommended: set this to true (larger mesh elements should have higher permeability)

• RNGSEED

- 1 integer; default: 0
- seed for random number generator
- value of 0 will seed with system time in milliseconds
- value of -1 will use the last 5 characters in the suffix
- value of -2 will use the last 4 charactes in the suffix and the millisecond time. This is useful when launching many iterations nearly simultaneously on a cluster.
- Other values: the seed is used directly for repeatable simulations.

• SNAPSHOTFILE

- value: 1 string; default: *.snap.out

 File for dumping out snapshots (concentration fields on all mesh elements). Can also be specified within SNAPSHOTS keyword.

• SNAPSHOTS

- 1 optional integer, 1 optional string, 1 optional logical; defaults: 1, *.snap.out, false
- Dump snapshots over the course of the calculation
- integer: how often to dump snapshots; string: snapshot file (* is replaced with suffix); logical: append rather than rewriting the snapshot file
- Snapshot file contains multiple snapshots of concentration profiles on all the mesh elements. Should be read with loadSnapshotFVM.m script

• VERBOSE

- value: 1 optional logical; default if not present: false; default if value unspecified: true
- Print extra output. Not really implemented in a useful way right now.