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Modeling neuron–glia interactions with the Brian 2 simulator

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1 Introduction

1.1 Challenges in modeling of neuron–glia interactions

Computational modeling is an important part of modern neuroscience research (Abbott, 2008). Neuronal models are available at many scales of investigation (Dayan and Abbott, 2001): highly detailed multi-compartmental models describe the ion channels that establish the membrane potential and are responsible for action potential generation; simplified models, such as the integrate-and-fire neuron, regard action potentials as stereotypical events that are not described any further; and even more abstract models, such as Poisson neurons, only aim to capture the statistics of timing of action potentials rather than the shape or the biophysical mechanism of their generation. Such neuronal models can be readily simulated by a wide range of simulator packages for neuronal simulations (Brette et al., 2007), given that they are well-established and fairly standardized and thus are often provided by libraries built in these packages (but see Brette, 2012).

The same does not hold true for glial cell models (De Pittà et al., 2013). Despite compelling emerging evidence that glial cells could crucially regulate neural network activity (Poskanzer and Yuste, 2016) and plasticity (De Pittà et al., 2016), the vast majority of available neuronal models completely ignores the possible contribution of glia to neuronal physiology. Arguably, one of the reasons for this is that standard glia models are yet to be defined (Chapter 1), and thus simulator packages generally do not provide models of glial cells as part of their libraries.

Although several popular neural simulators, such as NEURON (Carnevale and Hines, 2006) or NEST (Gewaltig and Diesmann, 2007), allow for extending their built-in library with user-defined glia models, this is generally not straightforward as it requires specific programming skills in a low-level language. Additionally, it usually involves an additional step of compilation and integration into the simulator every time the library is changed. This may ultimately discourage research that involves adding or modifying a glia model in a simulator, since iterative
improvements are very inconvenient. As a consequence, computational studies that model glia either use a standard simulator such as NEURON but are limited in usability by the specific choice made for the glial model which cannot easily be modified by the user (Aleksin et al., 2017), or use custom code written in a general-purpose language such as MATLAB (De Pittà et al., 2011; Naeem et al., 2015; Wade et al., 2011) or lower-level languages such as C/C++ (Nadkarni et al., 2008; Volman et al., 2007), in turn suffering from reduced accessibility to researchers with less technical experience. Overall, both scenarios raise potential issues in terms of portability, reproducibility and correctness of the code which are detrimental to model sharing and dissemination (Cannon et al., 2007).

In contrast to most other simulators, the Brian simulator (Goodman and Brette, 2008; Goodman et al., 2009) was created with the aim to ease definition and portability of new models. In its latest version, Brian 2, this flexibility is extended to and combined with a generic approach for code generation that allows high-performance simulations (Goodman, 2010; Stimberg et al., 2014). This chapter explores these aspects, elucidating several advantages that should encourage researchers to use the Brian 2 simulator to model glia in their work.

1.2 The Brian simulator

The Brian simulator, created in 2008, is provided as a package for the Python programming language. All aspects of the model can be defined in a single Python script and are made explicit: rather than relying on predefined “black-box” models, models can be readily and flexibly described in mathematical terms by differential equations for continuous dynamics and a series of update statements for discrete events (Brette 2012; Stimberg et al., 2014). This allows high code readability and flexibility as the user can freely change details of the model’s equations which are written in mathematical notation with only very little Brian-specific syntax. Furthermore, because the description of the model is explicit, all model details are unambiguously defined and appear in the main simulation description file.

In line with the principle of readability and simplicity, Brian 2 also comes with a system for the use of physical units. For example, it allows the user to directly specify a parameter in µm/s units, by multiplying its value by umolar/second. Brian 2 also checks the consistency of all specified units across all assignments, statements, and equations and issues an error if there is a mismatch.

Brian is open source and freely distributed under the GPL-compatible CeCILL v2.1 license. For more information see http://briansimulator.org and http://brian2.readthedocs.io.

1.3 Modeling strategy

In the following, we focus on astrocytes and their interactions with synapses, but the modeling arguments and code design principles that we present are of general validity and could be used to model other glial cells, such as microglia, oligodendrocytes or reactive astrocytes. Computational modeling of neuron–astrocyte signaling has previously been tackled both on the microscopic (molecular) level and the macroscopic (network) scale. On the microscopic level, the MCell simulator (Stiles et al., 2001) has been used to investigate specific astrocytic signals impinging on synaptic elements (Beenhakker and Huguenard, 2010). On the network level, the NEURON-based ARACHNE platform to study astrocyte functions in neural network physiology is available, but only considers astrocyte-mediated ‘volume-transmitted’ extracellular signaling (Aleksin et al., 2017; Savtchenko and Rusakov, 2014).

In the following modeling section we pursue instead a mixed strategy, considering models of astrocytes and of their interactions with synapses that lump both microscopic and macroscopic aspects. Based on this approach, we show how Brian 2 can be used to create a network of
neurons and synapses connected with a network of astrocytes that sense synaptic activity and modulate it in turn, starting from a molecular-level description of astrocytic signaling. We do so by first introducing a simple network model of only neurons and synapses (Sections 2.2 and 2.3). Then, we present modeling of individual astrocytes that respond to synaptic activation by intracellular calcium signaling (Section 2.4) and release gliotransmitters that could modulate synaptic transmission (Sections 2.5 and 2.6). Next, we discuss signaling between astrocytes in a network (Section 2.7) and finally we combine all these aspects in a recurrent network of interacting neurons, synapses, and astrocytes (Section 2.8).

For the sake of brevity, we only show excerpts of the Brian 2 code describing the models used in the presented simulations. The full code (see also Appendix B), including the code for recording and analyzing the results, as well as plotting the figures of this chapter can be downloaded from https://github.com/mdepitta/comp-glia-book. It is also part of the Brian 2 documentation at http://brian2.readthedocs.io and available in ModelDB (McDougal et al., 2017) at http://modeldb.yale.edu/233393.

2 Modeling of neuron–glia network interactions with Brian 2

2.1 General approach

In Brian 2, models of neurons, synapses, and astrocytes are defined by a set of state variables, e.g. the neuron’s membrane potential, the synaptic conductances, or the astrocyte’s intracellular calcium, and a description of their evolution over time. This description takes the form of ordinary differential equations (ODEs) for the continuous temporal dynamics between “events”, e.g. the membrane potential, postsynaptic conductances or astrocytic intracellular calcium concentration between action potentials. Discontinuities in the dynamics of state variables triggered by events such as the crossing of the firing threshold by a neuron’s membrane potential, or the arrival of an action potential at a presynaptic terminal, are described by a set of statements that update these variables.

Groups of elements that share the same description of their dynamics are represented by a single object. Thus, for example, groups of neurons, synapses, and astrocytes would each be represented by an object. However, these groups can be heterogeneous. For example, neurons or astrocytes in the same group can be stimulated by different synapses, or synapses can be characterized by different cellular parameters. State variables are updated according to the dynamics specified by the user. If the evolution of a state variable is of interest for the purpose of analysis or visualization, the user can record it by a “monitor” object either for all elements of a group or for a subset thereof.

Once all elements of a model have been specified, including initial values, constants, and model parameters, the simulation can be launched. This is done by calling the run function with a parameter that specifies the desired total simulated time in biological time units (e.g. second). Upon completion of the run call, the user can analyze the simulation results either by accessing the final values of state variables (for example to analyze the synaptic weight distribution at the end of a simulated plasticity-inducing protocol), or by accessing values stored in a monitor. All these values are readily accessible as NumPy arrays (van der Walt et al., 2011) and can be stored, analyzed and displayed by standard tools. Since simulation results are annotated with physical units, plotting them in a specific scale can easily be done by dividing them by those units. For example, when the state variable v (“membrane potential”) of a group of neurons has been recorded by monitor=StateMonitor(neurons, ’v’), the first neuron’s membrane potential may be shown in mV as function of time in ms using Matplotlib’s (Hunter, 2007) plot function by plot(monitor.t/ms, monitor[0].v/mV).
2.2 Neurons

In *Brian 2*, neurons are represented by objects of the `NeuronGroup` class (Figure 1A). Each `NeuronGroup` object models the activity of a group of neurons with identical dynamics, i.e. neurons whose state variables evolve according to the same differential equations. Consider, for example, the simple model of an integrate-and-fire neuron with conductance-based excitatory (\( g_e \)) and inhibitory synapses (\( g_i \)) and a constant input current \( I_{ex} \) [Dayan and Abbott 2001], whose equations are

\[
\begin{align*}
\frac{dv}{dt} &= \left( g_l (E_l - v) + g_e (E_e - v) + g_i (E_i - v) + I_{ex} \right) \frac{1}{C_m} \\
\frac{dg_e}{dt} &= -\frac{g_e}{\tau_e} \\
\frac{dg_i}{dt} &= -\frac{g_i}{\tau_i}
\end{align*}
\]

In *Brian 2* we closely follow the above mathematical notation, defining the neuron model by a multi-line string:

```
neuron_eqs = '''
  dv/dt = (g_l*(E_l-v) + g_e*(E_e-v) + g_i*(E_i-v) + I_ex) / C_m : volt (unless refractory)
  dg_e/dt = -g_e / tau_e : siemens # post-synaptic exc. conductance
  dg_i/dt = -g_i / tau_i : siemens # post-synaptic inh. conductance
'''
```

Each line of the `neuron_eqs` string defines a state variable of the model (\( v, g_e \) or \( g_i \)) and consists of two parts separated by a colon and an optional comment after the `#` sign. The part before the colon specifies the ODE for the dynamics of the state variable. The part that follows the colon specifies the physical units of this state variable and (optionally) additional information about it. In the case of the neuron’s membrane potential \( v \) for example, the additional specification (unless refractory) states that the differential equation for \( v \) is not to be integrated during the refractory period following the firing of an action potential by the neuron, but instead should stay at the post-spike reset value. It should be noted that the stated physical unit after the colon has to be a SI base unit, i.e. a unit such as volt, second, or siemens, and not a scaled unit such as mV, ms, or nS\(^1\). This is to emphasize that internally all variable values are stored as floating point numbers in the base unit. While users do not have to care about this most of the time – they set values using the unit system in whatever scale they prefer and receive values with the scaling information back – this become relevant when the unit information is stripped away, e.g. when quantities are passed through library functions that are not unit-aware.

The implementation of the neuron model in equations 1–3 refers to model parameters, namely the leak, excitatory, and inhibitory reversal potentials \( E_l, E_e \) and \( E_i \); the constant input current \( I_{ex} \); the membrane capacitance \( C_m \); and the time constants of excitatory and inhibitory synaptic inputs, \( \tau_e \) and \( \tau_i \). Here, these parameters are taken to be equal for all simulated neurons and can be defined by standard Python variables (one per parameter) in the script that runs the simulation\(^2\). Alternatively, neuron-specific parameters can be defined by appending lines in the form of `name : unit (constant)` to the model equations.

\(^1\)Note that molar is not a SI base unit, because it is defined as \( M = \text{mol} \text{L}^{-1} \), i.e. referring to L instead of the SI base unit m\(^3\). Since \( 1 \text{mol} \text{L}^{-1} = 1000 \text{mol} \text{m}^{-3} \), the base unit to use is nmol (mmolar).

\(^2\)Brian 2 also offers an alternative system to specify constants via a `namespace` argument that receives a Python dictionary mapping constant names to their values. Refer to *Brian 2*’s online documentation for details at [http://brian2.readthedocs.io](http://brian2.readthedocs.io).
In the scenario of a network of $N_e$ excitatory and $N_i$ inhibitory neurons (Figure 1B), we can then create a `NeuronGroup` object of $N_e + N_i$ neurons based on the above description, and further define the condition for firing of an action potential (threshold), as well as the statement(s) (if any) to be executed after an action potential (reset), and finally the refractory period (refractory), which in this case we assume to be a constant value $\tau_r$ (defined along with the other model parameters):

```python
neurons = NeuronGroup(N_e + N_i, model=neuron_eqs,
    threshold='v>V_th', reset='v=V_r',
    refractory='tau_r', method='euler')
```

The model’s state variables are exposed as attributes of the `neurons` object so that, for example, the membrane potential can be accessed by `neurons.v`, and their initial value can be assigned in the same way. Although all neurons in our example are described by the same equations, the initial values of their variables can be different. Here, we set these latter to random values using string expressions that are executed via the code generation facilities provided by Brian 2 (Stimberg et al., 2014) and refer to uniformly distributed random numbers between 0 and 1 using the predefined function `rand()`. Finally, we use Python’s slicing syntax to separate the group into subgroups of excitatory and inhibitory neurons:

```python
neurons.v = 'E_l + rand()*(V_th-E_l)'
neurons.g_e = 'rand()*w_e'
neurons.g_i = 'rand()*w_i'
exc_neurons = neurons[:N_e]
inh_neurons = neurons[N_e:]
```

### 2.3 Synapses

In most models of neuronal systems, neurons are connected by chemical synapses that are activated by action potentials fired by presynaptic neurons. In the following, we use the phenomenological description of neocortical synapses exhibiting short-term plasticity originally introduced by Tsodyks and Markram (Tsodyks, 2005; Tsodyks et al., 1998). According to this description, synaptic release is modeled by the product of two variables $u_S$ and $x_S$, where $u_S$ loosely relates to the neurotransmitter resources “docked” for release by the Ca$^{2+}$ sensor for synaptic exocytosis of neurotransmitter, and $x_S$ represents the fraction of total neurotransmitter available for release (Fuhrmann et al., 2002; Tsodyks, 2005). Between action potentials, $u_S$ decays to 0 at rate $\Omega_f$ while $x_S$ recovers to 1 at rate $\Omega_d$, i.e.

\[
\frac{du_S}{dt} = -\Omega_f u_S \tag{4}
\]
\[
\frac{dx_S}{dt} = \Omega_d (1 - x_S) \tag{5}
\]

The arrival of an action potential triggers calcium influx at the presynaptic terminal, which moves a fraction $U_0$ of the neurotransmitter resources not scheduled for release ($1 - u_S$) to the readily-releasable “docked” state ($u_S$). Subsequently, a fraction $u_S$ of the available neurotransmitter resources is released as $r_S$ while $x_S$ is reduced by the same amount, that is

\[
u_S \leftarrow u_S + U_0 (1 - u_S) \tag{6}
\]
\[
r_S \leftarrow u_S x_S \tag{7}
\]
\[
x_S \leftarrow x_S - r_S \tag{8}
\]

In Brian 2, connections between neurons are modeled by objects of the `Synapses` class (Figure 1A). Analogously to neurons of a `NeuronGroup`, we define each synapse’s state variables $x_S$ and $u_S$ by a multi-line string:
Figure 1. Modeling of neurons and synapses. A In Brian 2 neurons and synapses are modeled by two distinct classes, NeuronGroup and Synapses. As a convention, in all the schemes presented in this chapter, NeuronGroup objects are shown as rectangles, whereas Synapses objects are displayed as ellipses. B In a classic balanced network model (Brunel, 2000), neurons are separated into excitatory (exc_neurons) and inhibitory ones (inh_neurons), being recurrently connected both by excitatory (exc_synapses) and by inhibitory synapses (inh_synapses). C Raster plot of the firing activity of 25% out of all excitatory (red) and inhibitory neurons (blue) of the network in panel B and associated network-averaged firing rate (computed in 1 ms time bins). D Asynchronous network activity coexists with large fluctuations in postsynaptic excitatory (red traces) and inhibitory conductances (blue traces) and relatively sporadic firing by individual neurons (green dotted line: firing threshold, $V_{th}$). Timing of incoming presynaptic action potentials also shapes the dynamics of synaptic transmission by short-term synaptic plasticity. Synaptic release of neurotransmitter ($r_S$) is not fixed, but rather varies at each action potential, depending on the history of synaptic activity reflected in the values of the synaptic state variables $u_S$ (orange) and $x_S$ (purple) at the action potential instant. Postsynaptic conductances and membrane potential are shown for neuron 50 from the raster plot. Displayed synaptic variables are from one excitatory synapse made by this neuron. Model parameters as in Table C.1 and in addition, $I_{ex} = 150 \text{ pA}$. 
Synaptic equations are specified to be (event-driven) because we only need values of synaptic variables when an action potential arrives presynaptically. This tells Brian 2 to update synaptic variables only at new incoming action potentials using the analytical solution of their differential equations and the time interval since the last update. This is possible because the synapse’s equations are linear so that their analytical solution is known, allowing to simulate a large number of synapses efficiently. We also note that in our model implementation, synaptic equations do not include postsynaptic conductances as they were previously defined in the neuronal equations. This allows storing and updating synaptic conductances only once per neuron rather than once per synapse and is mathematically equivalent due to the assumed linear summation of postsynaptic conductances.

Discrete changes of synaptic variables on arrival of an action potential can be implemented by a series of statements in a multi-line string such as

```python
synapses_action = '''
u_S += U_0 * (1 - u_S)
r_S = u_S * x_S
x_S -= r_S
'''
```

In addition, excitatory (inhibitory) synapses will increase the excitatory (inhibitory) conductance in the postsynaptic cell whenever a presynaptic action potential arrives (Figure 1B), i.e.

\[
\begin{align*}
ge_e &\leftarrow g_e + w_e r_S \\
g_i &\leftarrow g_i + w_i r_S
\end{align*}
\]  

(9)  
(10)

This is done by extending `synapses_action` by an update assignment `+=` of the respective postsynaptic conductance, identified by the `post` suffix, i.e. `g_e_post` and `g_i_post`. Combining all of this together, we create two types of synapses, respectively originating from excitatory and inhibitory neurons, i.e.

```python
exc_syn = Synapses(exc_neurons, neurons, model=synapses_eqs, 
on_pre=synapses_action+'g_e_post += w_e*r_S')
inh_syn = Synapses(inh_neurons, neurons, model=synapses_eqs, 
on_pre=synapses_action+'g_i_post += w_i*r_S')
```

where the `on_pre` keyword argument denotes that the series of statements should be executed on arrival of a presynaptic action potential.

It must be stressed that the above code only defines the synaptic model in Brian 2, but not the connectivity. To create synapses, we have to specify what source–target neuron pairs should be connected together out of all the possible pairs specified by the first two input arguments in the Synapses initializer. One way to do this is by specifying a logical condition on neuronal connectivity and, optionally, a connection probability, provided as arguments to the `connect` method of the Synapses object (Stimberg et al., 2014). Here, we want to connect all possible neuron pairs with a probability of 20% for each pair for inhibitory neurons, and 5% for excitatory neurons. Thus we do not set any condition to be fulfilled and only specify a probability for all possible connection pairs:

```python
exc_syn.connect(p=0.05)
inh_syn.connect(p=0.2)
```
For parameters values such as those in Table C.1, simulation of the resulting network reveals the emergence of characteristic asynchronous neuronal firing activity (Brunel, 2000), as evidenced in Figure 1C by the top left raster plot of the firing activity of 25% of the neurons in the network. The network-averaged firing rate associated with this raster plot indeed presents noisy dynamics (Figure 1C, bottom panel) that coexists with large fluctuations of postsynaptic excitatory (red traces) and inhibitory conductances (blue traces) and sporadic firing of individual neurons (Figure 1D, top panels). Consideration of a sample excitatory synapse of the network allows appreciating how synaptic dynamics is modulated by short-term plasticity (Figure 1D, bottom panels). The state variables $u_S$ and $x_S$ associated with the sample synapse evolve in a characteristic exponential fashion intermingled with discontinuities triggered by action potentials arriving at the presynaptic terminal. The resulting values define the amount of synaptic neurotransmitter released by the synapse ($r_S$), setting how effectively each action potential is transmitted to the postsynaptic neuron.

2.4 Astrocytes

Intracellular Ca\(^{2+}\) concentration is unanimously regarded as a prominent readout signal of astrocyte activity (Zorec et al., 2012). Although astrocytic intracellular Ca\(^{2+}\) can be regulated by multiple mechanisms, Ca\(^{2+}\)-induced Ca\(^{2+}\) release (CICR) from the astrocyte’s endoplasmic reticulum (ER) appears to be one of the main mechanisms to regularly occur in the healthy brain (Nimmerjahn, 2009). Recall from Chapter 5 that astrocytic CICR is triggered by the intracellular second messenger inositol 1,4,5-trisphosphate (IP\(_3\)), which is produced upon astrocyte activation, and can be described, in one of its simplest formulation (De Pitta et al., 2008; Falcke, 2004), by two ordinary differential equations in the Hodgkin-Huxley form (Li and Rinzel, 1994). The first equation is a mass balance for Ca\(^{2+}\) ($C$) in terms of three fluxes $J_r$, $J_l$, $J_p$ which respectively denote CICR ($J_r$), Ca\(^{2+}\) leak from the ER ($J_l$), and Ca\(^{2+}\) uptake from the cytosol back to the ER by Ca\(^{2+}\)/ATPase pumps ($J_p$). The second equation is for the gating variable ($h$) of de-inactivation of the channels that are responsible for CICR. These channels are inside the astrocyte, on the membrane that separates the ER Ca\(^{2+}\)-rich stores from the cell’s cytosol, and are nonlinearly gated by both IP\(_3\) ($I$) and Ca\(^{2+}\). This leads to the well-known two-equation model originally introduced by Li and Rinzel (1994):

$$\frac{dC}{dt} = J_r + J_l - J_p$$
$$\frac{dh}{dt} = \frac{h_\infty - h}{\tau_h}$$

where

$$J_r = \Omega_C m_\infty^3 h^3 (C_T - (1 + g_A)C)$$
$$J_l = \Omega_L (C_T - (1 + g_A)C)$$
$$J_p = O_P H_2 (C, K_F)$$

and $H$ denotes the sigmoidal (Hill) function with $H_n (x, K) = x^n / (x^n + K^n)$.

For the sake of completeness, we also want to consider the stochastic opening and closing process of CICR-mediating channels which, as discussed in Chapter 4, can be mimicked by introducing a white noise term $\xi(t)$ into the equation governing the dynamics of the gating variable $h$, so that equation 12 becomes (Shuai and Jung, 2002):

$$\frac{dh}{dt} = \frac{h_\infty - h}{\tau_h} (1 + \xi(t)\sqrt{\tau_h})$$
Astrocytic IP₃ is regulated by the complex Ca²⁺-modulated interplay of enzymatic production by phospholipase Cβ (Jβ) and Cδ (Jδ) and degradation by IP₃ 3-kinase (J₃K) and inositol polyphosphatase 5-phosphatase (J₅P) (De Pittà et al., 2009a). To reproduce experimental observations we consider two possible ways to trigger IP₃ production. One is by synaptic stimulation of astrocytic metabotropic receptors which starts phospholipase Cβ-mediated IP₃ production, modeled by making Jβ proportional to the activated fraction of these receptors (denoted hereafter by ΓA). The other way is to include a further Jex term for constant IP₃ production by an exogenous source of stimulation such as, for example, IP₃ uncaging or intracellular diffusion from subcellular regions far from the CICR site (Goldberg et al., 2010).

In conclusion, IP₃ dynamics evolves according to the mass balance equation (De Pittà et al., 2009a)

\[
\frac{dI}{dt} = J_\beta(\Gamma_A) + J_\delta - J_3K - J_5P + J_{ex}
\]

(14)

with

\[
J_\beta(\Gamma_A) = O_\beta \Gamma_A
\]

\[
J_\delta = O_\delta \frac{K_\delta}{K_\delta + I} \mathcal{H}_2(C, K_\delta)
\]

\[
J_3K = O_3K \mathcal{H}_4(C, K_D) \mathcal{H}_1(I, K_3)
\]

\[
J_5P = \Omega_5P I
\]

\[
J_{ex} = -F_{ex}/2 \left(1 + \tanh \left(\frac{|\Delta I_{bias}| - I_\Theta}{\omega_I}\right)\right) \text{sgn}(\Delta I_{bias})
\]

where the fraction of activated astrocyte receptors ΓA depends on the neurotransmitter concentration in the periastrocytic space YS, and is given by the further equation (Wallach et al., 2014) (Chapter 5)

\[
\frac{d\Gamma_A}{dt} = O_N Y_S (1 - \Gamma_A) - \Omega_N (1 + \zeta \cdot \mathcal{H}_1(C, K_KC)) \Gamma_A
\]

(15)

For a concise description of the meaning of all the model’s parameters in the above equations, see Table C.2 in the Appendix.

Dynamics of the astrocyte’s state variables ΓA, I, C, h are governed by ODEs akin to neuronal state variables, although on a longer time scale (De Pittà et al., 2009b). Accordingly, they can be implemented by a NeuronGroup object, exactly in the same way as neuronal variables. In this fashion, the following code exemplifies how we can create two astrocytes with dynamics governed by the above equations:

```python
astro_eqs = '# Fraction of activated astrocyte receptors:
dGamma_A/dt = O_N * Y_S * (1 - Gamma_A) -
            Omega_N*(1 + zeta * C/(C + K_KC)) * Gamma_A : 1

# IP_3 dynamics:
dI/dt = J_beta + J_delta - J_3K - J_5P + J_ex : mmolar
J_beta = O_beta * Gamma_A : mmolar/second
J_delta = O_delta / (1 + I/kappa_delta) * C**2/(C**2 + K_delta**2) : mmolar/second
J_3K = O_3K * C**4/(C**4 + K_D**4) * I/(I + K_3K) : mmolar/second
J_5P = Omega_5P*I : mmolar/second
delta_I_bias = I - I_bias : mmolar
J_ex = -F_ex/2*(1 + tanh((abs(delta_I_bias) - I_Theta)/omega_I)) *
       sign(delta_I_bias) : mmolar/second
I_bias : mmolar (constant)
```
# Ca^{2+}-induced Ca^{2+} release:
\[ \frac{dC}{dt} = J_r + J_l - J_p : \text{mmolar} \]

# IP3R de-inactivation probability
\[ \frac{dh}{dt} = \frac{(h_{\text{inf}} - h_{\text{clipped}})}{\tau_h} \times (1 + \text{noise} \times \xi \times \tau_h^{0.5}) : 1 \]

\[ h_{\text{clipped}} = \text{clip}(h,0,1) : 1 \]

\[ J_r = \left( \Omega_C \times m_{\text{inf}}^3 \times h_{\text{clipped}}^3 \right) \times \left( C_T - (1 + \rho_A) \times C \right) : \text{mmolar/second} \]

\[ J_l = \Omega_L \times \left( C_T - (1 + \rho_A) \times C \right) : \text{mmolar/second} \]

\[ J_p = \frac{O_P \times C^2}{C^2 + K_P^2} : \text{mmolar/second} \]

\[ m_{\text{inf}} = \frac{I}{I + d_1} \times \frac{C}{C + d_5} : 1 \]

\[ h_{\text{inf}} = \frac{Q_2}{Q_2 + C} : 1 \]

\[ \tau_h = \frac{1}{O_2 \times (Q_2 + C)} : \text{second} \]

\[ Q_2 = d_2 \times \frac{I + d_1}{I + d_3} : \text{mmolar} \]

# Neurotransmitter concentration in the extracellular space
\[ Y_S : \text{mmolar} \]

# Noise flag
\[ \text{noise} : 1 \text{ (constant)} \]

```python
# Milstein integration method for the multiplicative noise
astrocytes = NeuronGroup(2, astro_eqs, method='milstein')
```

In the above Brian 2 code, the multi-line astro_eqs string defines our astrocyte model. The white noise term $\xi$ in equation 13 is implemented by the special symbol $\xi$ (in units of $s^{-1/2}$) which is a predefined symbol in Brian 2. Also note that as a gating variable, $h$ has to be bound between 0 and 1, which is not guaranteed by the (approximated) nature of equation 13. We therefore restrict $h$ to this interval by replacing it by $h_{\text{clipped}}=\text{clip}(h,0,1)$ on the right hand side of the ODE for $h$ and in the formula of $J_r$ (Shuai and Jung, 2002). The noise term in equation 13 is multiplicative ($\xi(t)$ multiplies the gating variable $h$), we therefore have to use a numerical integration method that can handle multiplicative stochastic differential equations (under the Stratonovich interpretation, cf. Kloeden and Platen, 1992). We do this by specifying method='milstein' as an argument to the NeuronGroup initializer, leading to the use of the Milstein method for integration (Kloeden and Platen, 1992; Mil’shtein, 1975; also see details on example 2_gchi_astrocyte.py in Appendix B).

The above model also defines three astrocyte-specific variables that are not defined by equations: $I_{\text{bias}}$, $Y_S$, and $\text{noise}$. $I_{\text{bias}}$ and $\text{noise}$ are constant over time, but $Y_S$, the concentration of synaptically-released neurotransmitter in the extracellular space around astrocytic receptors, i.e. $Y_S$ in equation 15 depends on synaptic activity that changes and therefore does not have the (constant) flag. We will define how it gets linked to the synaptic activity further below.

In this example, we want to compare two types of astrocytes, a deterministic and a stochastic one. To distinguish them, we have introduced the above-mentioned $\text{noise}$ constant which scales the strength of the noise term in equation 13. We can therefore switch the noise term on or off, and we initialize it so that the first astrocyte is deterministic and the second is stochastic:

\[ \text{astrocytes.noise} = [0, 1] \]

To complete our model, we also need to specify how to calculate $Y_S$ in equation 15, as it is needed for the integration of the $\Gamma_A$ state variable. For now, we are only interested in the activity of the astrocyte and how it is stimulated by synaptic neurotransmitter. Therefore, we do not take into account short-term synaptic plasticity, and rather consider a trivial synaptic model stating that $Y_S$ increases by the same amount at every action potential and then decays...
Figure 2. Modeling of synaptically-activated astrocytes. A Illustration of the design principles to implement an astrocyte stimulated by synaptic inputs in Brian 2. The coupling between synapses and the astrocyte is achieved by an additional Synapses object, labeled here as ecs_syn_to_astro, which feeds into the astrocyte the sums of all individual synaptic inputs. B Simulated dynamics of the state variables of two synaptically-activated astrocytes, one deterministic (red traces) and one with noise on the gating variable $h$ according to equation 13 (blue traces). The two astrocytes shared the same synaptic input which was repeatedly triggered by action potentials arriving at a rate of 0.5 Hz. Model parameters as documented in tables in Appendix C with the exception of $\rho_c = 0.001$; $K_P = 0.1 \mu$M; $O_\beta = 5 \mu$M s$^{-1}$; $O_\delta = 0.2 \mu$M s$^{-1}$; $\Omega_{5P} = 0.1$ s$^{-1}$; $K_D = 0.5 \mu$M; $F_{ex} = 0.09 \mu$M; $K_\delta = 0.3 \mu$M; and $I_{bias} = 0$. 
exponentially (Dayan and Abbott, 2001):

\[ \text{synapses_eqs} = 'dY_S /dt = -\Omega_c * Y_S : \text{mmolar (clock-driven)}' \]
\[ \text{synapses_action} = 'Y_S += \rho_c * Y_T' \]
\[ \text{synapses} = \text{Synapses(source_neurons, target_neurons, model=synapses_eqs, on_pre=synapses_action, method='linear')} \]
\[ \text{synapses.connect()} \]

Finally we have to specify how the synapses stimulate the astrocytes. We do this by introducing a further \text{Synapses} object that connects our synapses (which thus represent the “presynaptic” source in our object) to the two astrocytes (hence regarded as “postsynaptic” targets) in an all-to-all fashion, which can be concisely expressed by a call to \text{connect()} without any arguments. Each astrocyte \( i \) senses the sum of \( Y_S \) across all \( S \) synapses that impinge on it. This can be mathematically expressed as \( Y_{S,post}^i = \sum_{j \in S} Y_{S,j} \), and the implementation in \text{Brian 2} closely follows this formulation, using the flag \text{(summed)} to denote the summing operation:

\[ \text{ecs_syn_to_astro} = \text{Synapses(synapses, astrocytes, \text{'Y_S_post = Y_S_pre : mmolar (summed)')}} \]
\[ \text{ecs_syn_to_astro.connect()} \]

The above definition will update \( Y_S \) for each astrocyte at every time step so that the integration of the \( \Gamma_A \) has access to the current value at all times.

Figure 2A summarizes the design of the astrocyte model previously described, exemplifying its implementation by \text{Brian 2} classes originally introduced in Figure 1A. A sample simulation of this model is presented in Figure 2B, which shows the dynamics of the state variables \( \Gamma_A, I, C \) and \( h \) for the deterministic (\text{red traces}) vs. stochastic astrocyte (\text{blue traces}) in response to synaptic stimulation by periodic action potentials at a rate of 0.5 Hz. It may be appreciated how noisy dynamics of the gating variable \( h \) could dramatically alter \( C \) and \( I \) dynamics compared to the deterministic scenario. This could also impact the activated fraction of astrocytic metabotropic receptors (\( \Gamma_A \)) by the \( \text{Ca}^{2+} \)-dependent Hill nonlinearity in the right hand side of equation 15, although in this example, the effect may be deemed moderate for the specific choice of values for the model’s parameters (Table C.2).

2.5 Gliotransmitter release and modulation of synaptic release

Astrocytes are not only stimulated by synapses but they can also modulate them by releasing neurotransmitters (also termed “gliotransmitters” for their glial origin; Bezzi and Volterra, 2001) in a \( \text{Ca}^{2+} \)-dependent fashion. This process generally requires astrocytic intracellular \( \text{Ca}^{2+} \) concentration to increase beyond a certain threshold, resulting in the release of a quantum of gliotransmitter into the periastrocytic space (De Pittà et al., 2013). In turn, released gliotransmitter molecules diffuse in the extracellular space, ultimately reaching extrasynaptic receptors found on synaptic elements belonging either to the very synapses that stimulate the astrocyte in the so-called “closed-loop” scenario of gliotransmission, or to other synapses in the case of “open-loop” gliotransmission (Araque et al., 2014). Among these targeted receptors, presynaptically-located ones, once bound by gliotransmitters, can ultimately regulate synaptic transmission through modulations of synaptic release probability (Engelman and MacDermott, 2004; Pinheiro and Mulle, 2008). In the simplest approximation, as elucidated in Chapter 12, this modulation can be modeled by treating the parameter \( U_0 \) in the previously-introduced Tsodyks-Markram synapse model (equation 4) no longer as a constant, but rather as linearly dependent on the fraction \( \Gamma_S \) of activated presynaptic receptors (De Pittà et al., 2011):

\[ \frac{d\Gamma_S}{dt} = O_G G_A (1 - \Gamma_S) - \Omega_G \Gamma_S \] (16)
\[ U_0 = (1 - \Gamma_S)U^*_0 + \alpha \Gamma_S, \quad (17) \]

where \( G_A \) denotes the gliotransmitter concentration in the extracellular space, and \( \alpha \) dictates whether the effect of gliotransmitters on the synapse is to decrease \((0 \leq \alpha < U^*_0)\) or increase neurotransmitter release \((U^*_0 < \alpha \leq 1)\) \cite{DePittà2011}. In Brian 2 syntax, this leads to the following synaptic equations:

```
synapses_eqs = '''
# Neurotransmitter
 dY_S/dt = -\Omega_c * Y_S : mmolar (clock-driven)
# Fraction of activated presynaptic receptors
 d\Gamma_S/dt = O_G * G_A * (1 - \Gamma_S) -
             \Omega_G * \Gamma_S : 1 (clock-driven)
# Usage of releasable neurotransmitter per single action potential:
 du_S/dt = -\Omega_f * u_S : 1 (clock-driven)
# Fraction of synaptic neurotransmitter resources available:
 dx_S/dt = \Omega_d *(1 - x_S) : 1 (clock-driven)
# released synaptic neurotransmitter resources:
r_S : 1
# gliotransmitter concentration in the extracellular space:
\[ G_A : mmolar \]
'''
```

Because the value of \( U_0 \) is only needed at the arrival of an action potential, there is no need to include equation (17) in the above code. Rather, we update \( U_0 \) at the beginning of the statements executed by synapses upon action potential arrival, i.e.

```
synapses_action = '''
 U_0 = (1 - \Gamma_S) * U_0__star + \alpha * \Gamma_S
 u_S += U_0 * (1 - u_S)
 r_S = u_S * x_S
 x_S -= r_S
 Y_S += \rho_c * Y_T * r_S
'''
```

For the sake of simplicity we retained two inefficiencies in the above code which should otherwise be avoided in larger, computationally-demanding simulations. First, we used the (clock-driven) specification (instead of (event-driven)), even though synaptic state variables need only be updated on action potential arrival (Section 2.3). This allows us to directly retrieve and plot state variables at each time step without the need to manually interpolate between their values at the timing of action potentials. For the same reason, we also defined \( r_S \) (equation 7) as an additional state variable in the synapse model in `synapses_eqs` rather than using it as an auxiliary temporary variable in the statements of `synapses_action` as we previously did (Section 2.3). This choice allows us to easily record \( r_S \) by a monitor, avoiding the need to recompute it a posteriori based on the values of the other state variables.

Finally, we need to model gliotransmitter release from the astrocyte. For this, we use a similar description to that of synaptic neurotransmitter release (Chapter 13). We thus introduce a new variable \( x_A \) which represents the fraction of gliotransmitter resources available for release from the astrocyte and thereby controls the value of \( G_A \). These two state variables decay as \cite{DePittà2011}

\[
\frac{dx_A}{dt} = \Omega_A(1 - x_A) \quad (18)
\]

13
\[
\frac{dG_A}{dt} = -\Omega_e G_A, \tag{19}
\]
while, when gliotransmitter is released, they are updated according to
\[
G_A \leftarrow G_A + \rho_e G_T U_A x_A \tag{20}
\]
\[
x_A \leftarrow x_A (1 - U_A) \tag{21}
\]
As before, we can implement the above by textual equations and statements in Brian 2, i.e.

```python
astro_eqs = '''
# [...]
# Fraction of gliotransmitter resources available:
dx_A /dt = Omega_A * (1 - x_A) : 1
# gliotransmitter concentration in the extracellular space:
dG_A /dt = -Omega_e * G_A : mmolar
'''
glio_release = '''
G_A += rho_e * G_T * U_A * x_A
x_A -= U_A * x_A
'''
```

We also need to include in the astrocytic model a mechanism to time gliotransmitter release. We do this by imaging that when the astrocyte’s Ca\(^{2+}\) concentration increases beyond a threshold \(C_\theta\), the astrocyte “fires” a gliotransmitter release event upon which the statements of `glio_release` are executed. In this fashion, we can define in the astrocyte’s `NeuronGroup` a threshold crossing for Ca\(^{2+}\) concentration (by the `threshold` keyword), upon which gliotransmitter is released, and specify by the `reset` keyword what to do following firing of a gliotransmitter release event by the astrocyte. Moreover, to avoid the `threshold` condition repeatedly triggering the release in all the time steps where the Ca\(^{2+}\) concentration is above the threshold, we use the same condition for the `refractory` keyword, thereby stating that, as long as the Ca\(^{2+}\) concentration is above threshold, no new event should be triggered. That is,

```python
astrocyte = NeuronGroup(2, astro_eqs,
threshold='C>C_Theta',
refractory='C>C_Theta',
reset=glio_release,
method='rk4')
```

Finally, to complete our model, we have to define how synapses are modulated by the astrocytes’ gliotransmitter release. We do this by imaging that when the astrocyte’s Ca\(^{2+}\) concentration increases beyond a threshold \(C_\theta\), the astrocyte “fires” a gliotransmitter release event upon which the statements of `glio_release` are executed. In this fashion, we can define in the astrocyte’s `NeuronGroup` a threshold crossing for Ca\(^{2+}\) concentration (by the `threshold` keyword), upon which gliotransmitter is released, and specify by the `reset` keyword what to do following firing of a gliotransmitter release event by the astrocyte. Moreover, to avoid the `threshold` condition repeatedly triggering the release in all the time steps where the Ca\(^{2+}\) concentration is above the threshold, we use the same condition for the `refractory` keyword, thereby stating that, as long as the Ca\(^{2+}\) concentration is above threshold, no new event should be triggered. That is,

```python
ecs_astro_to_syn = Synapses(astrocyte, synapses,
'G_A_post = G_A_pre : mmolar (summed)')
```

Figure 3B illustrates how gliotransmitter release from the astrocyte could change synaptic neurotransmitter release in our model. The top panels show Ca\(^{2+}\) traces (\(C\)) from two astrocytes that are differently stimulated by exogenous IP\(_3\) production (\(J_{ex}\) in equation 14) so that their intracellular Ca\(^{2+}\) concentration crosses the threshold for gliotransmitter release (gray dashed line) at different times. This results in one astrocyte releasing gliotransmitter in the extracellular space (\(G_A\)) only once (green traces, “weak gliotransmission”), while the other releases gliotransmitter twice (purple traces, “stronger gliotransmission”). The modulation of
A

source_neurons -> spike event -> target_neurons

synapses

ecs_astro_to_syn

G_A (summed)

gliotransmitter in the extracellular space

astrocyte

with gliotransmitter release

B

C (µM)

G_A (µM)

time (s)

0
50
100
1500

0.0
0.4
0.8
1.2
C (/µM)

−16 −14 −12 −10 −8 −6 −4 −2 0

YS (µM)

0.00
0.25
0.50
0.75
1.00

0.0
100
200
300
400
500
600

time (ms)

0
500
1000
1500

0.00
0.25
0.50
0.75
1.00

0.00
0.25
0.50
0.75
1.00

X_M

Y_M (µM)

no gliotransmission
weak gliotransmission
stronger gliotransmission

black

green

purple

15
Figure 3. (Previous page.) Modeling of modulation of synaptic release by gliotransmission. A The model uses a Synapses object `ecs_syn_to_astro` to allow synapses’ modulation by perisynaptic gliotransmitters. B (top panels) Extracellular gliotransmitter concentration ($G_A$) resulting from intracellular Ca$^{2+}$ dynamics ($C$) in two differently activated astrocytes. (Bottom panels) Dynamics of synaptic state variables ($u_S$, $x_S$) and extracellular neurotransmitter concentration ($Y_S$) in response to a sequence of action potentials (top) for a synapse without (black) and with (green and purple) gliotransmitter-mediated modulation. The two astrocytes were initialized at $t = -17.1$ s at $x_A = 1$, $I = 0.4 \mu$M, and $h = 0.9$ and were respectively stimulated by $I_{bias} = 0.8 \mu$M (weak gliotransmission) and $I_{bias} = 1.25 \mu$M (stronger gliotransmission). Other model parameters as in the Tables in Appendix C.

Synaptic release ensuing from these two different timings of gliotransmitter release, is illustrated in the lower panels, where neurotransmitter release from a synapse stimulated by a sample train of action potentials, is monitored first in the absence of gliotransmitter release from the astrocyte, and then in the presence of weak vs. stronger gliotransmission. Without gliotransmission, the extracellular neurotransmitter concentration ($Y_S$) progressively decreases with the incoming action potentials, compatibly with the onset of strong short-term synaptic depression (black traces). In the presence of gliotransmission instead, while the amount of released neurotransmitter per action potential is generally lower than in the “naive” synapse (since we assumed $\alpha < 0$ in this example), this amount tends to increase at every action potential with respect to preceding ones, and this increase is larger for stronger gliotransmission. This is consistent with the onset of short-term synaptic facilitation, and agrees with the experimental observation and the theoretical argument that gliotransmission could change the synapse’s short-term plasticity [Araque et al. 2014 De Pittà et al. 2016 see also Chapters 8 and 13].

2.6 Closed-loop gliotransmission

In the examples discussed so far, we only separately considered one-way interactions between synapses and astrocytes, modeling either modulation of astrocytic activity by synaptic neurotransmitters (Section 2.4) or modulations of synaptic release by astrocytic gliotransmitters (Section 2.5). In practice however, these two pathways may coexist, with gliotransmission feeding back in a closed-loop fashion on the very synapses that stimulate the astrocyte and trigger its gliotransmitter release. This section focuses on such a closed-loop scenario for gliotransmission.

Closed-loop gliotransmission can easily be implemented in Brian 2 by combining the model of a synaptically-activated astrocyte (Figure 2A) with that of open-loop gliotransmission (Figure 3A), resulting in the model scheme shown in Figure 4A. However, it may be noted that in doing so we end up using two independent Synapses objects (ellipses) to separately describe the extracellular space for synapse-to-astrocyte signaling ($ecs_{syn\_to\_astro}$) and the extracellular space for astrocyte-to-synapse gliotransmission ($ecs_{astro\_to\_syn}$). In reality, both neurotransmitter and gliotransmitter release could occur in the same extracellular space, and thus only a single Synapses object might be considered at the benefit of computational efficiency. Nonetheless, the choice of using two independent objects allows us to take into account the more general scenario of astrocytes that are either sensitive or not to the activity of the same synapses they modulate [Martin et al. 2015 Navarrete and Araque 2016]. This is therefore an appropriate choice when dealing with many synapses interacting with astrocytes as in the case of the neuron–glia network discussed at the end of this chapter.

To elucidate some of the possible functional implications of closed-loop gliotransmission we set out to characterize the average synaptic release for $N_{synapses}=500$ identical synapses for different input stimuli and compare it to the open-loop scenario of gliotransmission as well as to
Figure 4. Closed-loop gliotransmission. A In the most general design, modeling of closed-loop gliotransmission in Brian 2 separates between the extracellular space of synapse-to-astrocyte signaling (ecs_syn_to_astro) and the extracellular space of astrocyte-to-synapse gliotransmission (ecs_astro_to_syn). B Average extracellular concentration of synaptically-released neurotransmitter ($\langle Y_S \rangle$) for step increases of the mean rate of Poisson-generated incoming action potentials (top panel, $\nu_{in} = 0.011$ Hz, $0.11$ Hz, $1.1$ Hz, $11$ Hz for 5-s time intervals; traces averaged over 500 identical synapses.) C Corresponding average release of synaptic neurotransmitter resources as function of the rate of incoming action potentials (data points and error bars: mean ± standard deviation for 100 trials.) Parameters as in the tables in Appendix C except for $O_\beta = 3.2 \mu$M s$^{-1}$; $I_{bias} = 1 \mu$M (open-loop gliotransmission); $I_{bias} = 0 \mu$M (closed-loop gliotransmission).
the “naive” scenario without gliotransmission. *Brian 2* is optimized to deal with large objects (see Appendix [A]), so rather than simulating one synapse at a time, for 500 times in the three different scenarios (i.e. 1500 simulations in total), we simulate all synapses in all scenarios in one single run. This is achieved by considering $500 \times 3$ synapses and an astrocyte group of $500 \times 2$ elements. The first 500 synapses are modulated by gliotransmitters from the first 500 astrocytes in a closed-loop fashion, the second group of 500 synapses is modulated by open-loop gliotransmission mediated by the other 500 astrocytes; and finally, the remaining synapses do not consider any gliotransmission. In contrast to previous synaptic connection patterns, here we can directly calculate a target index for each connection, instead of evaluating a logical condition for every possible connection pair. *Brian 2* has a built-in syntax for such descriptions, which offers a much more efficient way of establishing connections. In this syntax, we provide an expression to calculate the target index $j$ based on the source index $i$ and potentially other pre- or postsynaptic properties. To exclude certain potential connections, this expression can be combined with an optional if part stating the condition for a connection to exist. Remember that in our example here, the source index $i$ and the target index $j$ each refer either to synapses or astrocytes, depending on the direction of the connection (“synapses to astrocytes” or “astrocytes to synapses”). This leads to the following *Brian 2* code:

```
ecs_syn_to_astro = Synapses(synapses, astrocyte,
    'Y_S_post = Y_S_pre : mmolar (summed)')
# Connect the first N_synapses synapses—astrocyte pairs
ecs_syn_to_astro.connect(j='i if i < N_synapses')
ecs_astro_to_syn = Synapses(astrocyte, synapses,
    'G_A_post = G_A_pre : mmolar (summed)')
# Connect the first N_synapses astrocytes—synapses pairs (closed-loop)
ecs_astro_to_syn.connect(j='i if i < N_synapses')
# Connect the second N_synapses astrocyte—synapses pairs (open-loop)
ecs_astro_to_syn.connect(j='i if i >= N_synapses and i < 2*N_synapses')
```

Figure 4B shows a reproduction of Figure 13.2 by our *Brian 2* implementation of closed-loop gliotransmission for the time evolution of the average neurotransmitter concentration in the synaptic cleft ($\langle Y_S \rangle$) in response to step increases in the rate of incoming action potentials ($\nu_{in}$, top panel). Gliotransmission dramatically changes synaptic transmission (colored vs. black traces), with the effect of closed-loop gliotransmission (purple trace) being somewhat intermediate between the scenarios of no gliotransmission (black trace) and open-loop gliotransmission (green trace).

This is further elucidated in Figure 4C where the mean neurotransmitter concentration in the extracellular space in the three scenarios is shown for different mean rates of randomly incoming action potentials. The low-pass filter characteristics of synapses without gliotransmission (top panel) turns into a bell-shaped, band-pass filter characteristics caused by (release-decreasing) open-loop gliotransmission (middle panel) (Chapter 12). In the presence of closed-loop gliotransmission however, the average concentration of synaptically-released neurotransmitter is in between those expected in the other two scenarios for low input rate values, and tends to approach the shape of the curve in the open-loop scenario for increasing rates. For high input rates however, the release-decreasing effect of gliotransmission is such that the synapse is ultimately silenced and cannot sustain further gliotransmitter release. Synaptic transmission then becomes independent of gliotransmission again as if it were in the naive scenario without gliotransmission, which accounts for the jump at $\nu_{in} > 10$ Hz.
2.7 Networks of astrocytes

Astrocytes are known to arrange in networks of different shape and connectivity depending on the brain region under consideration (Giaume et al., 2010), and to be capable of propagating Ca\(^{2+}\) signals through such networks in the form of intercellular (regenerating) waves. The mechanisms underlying such propagation can be multiple and varied (Scemes and Giaume, 2006). Here, we only focus on the well characterized mechanism of intracellular IP\(_3\) diffusion through gap junctions channels (GJCs) between neighboring astrocytes (Chapter 7).

From a modeling perspective, IP\(_3\) diffusion from one astrocyte \(j\) to a neighboring one \(i\) can be thought as a flux of IP\(_3\) (\(J_{ij}\)) which is some nonlinear (rectifying) function of the IP\(_3\) concentration gradient between cells \(i\) and \(j\), i.e. \(\Delta_{ij}I = I_i - I_j\), such as, for example (Lallouette et al., 2014, see also Chapter 7)

\[
J_{ij} = \frac{-F}{2} \left(1 + \tanh \left(\frac{|\Delta_{ij}I| - I_\theta}{\omega_I}\right)\right) \text{sgn}(\Delta_{ij})
\]  

Incidentally, we note that the above formula is reminiscent of the expression of the exogenous IP\(_3\) flux (\(J_\text{ex}\)) in equation 14, insofar as the latter may be regarded as a special case of intercellular IP\(_3\) influx to any astrocyte from a much larger external IP\(_3\) source (i.e. \(I_{\text{bias}}\) in our notation) (Goldberg et al., 2010). Because \(J_{ij}\) is a function of IP\(_3\) concentrations in connected astrocytes (i.e. \(I_i, I_j\)) by \(\Delta_{ij}I\), it is astrocyte-dependent and not constant. Therefore, once we add \(J_{ij}\) to our astrocyte equations as in Brian 2 (denoted in the code below by \(J_{\text{coupling}}\)), we must define it as an astrocytic variable (that is without the (constant) flag), i.e.

\[
\text{astro_eqs} = 'dI/dt = J_{\delta} - J_{3K} - J_{5P} + J_{\text{ex}} + J_{\text{coupling}} : \text{mmolar} \\
# [\ldots] \\
# \text{Diffusion between astrocytes} \\
J_{\text{coupling}} : \text{mmolar/second} \\
# [\ldots] \\
'''
\]

Connections between astrocytes by GJCs may conveniently be implemented by a Synapses object in Brian 2, once we regard \(J_{ij}\) as the IP\(_3\) flow from “presynaptic” astrocyte \(j\) to “postsynaptic” astrocyte \(i\) (Figure 5B). The effective total \(J_{\text{coupling}}\) to cell \(i\) by intercellular IP\(_3\) diffusion is the sum of all IP\(_3\) fluxes incoming to cell \(i\) from the \(A_i\) astrocytes connected to this latter by GJCs, i.e. \(J_{\text{coupling}}^i = \sum_{j \in A_i} J_{ij}\). In Brian 2 code, this reads

\[
\text{astro_to_astro_eqs} = 'dI/dt = I_{\text{post}} - I_{\text{pre}} : \text{mmolar} \\
# [\ldots] \\
\text{J_{coupling}}_{\text{post}} = -F/2 * (1 + \tanh((\text{abs}(\text{delta}_{I}) - I_\theta)/\omega_I)) * \\
\text{sign}(\text{delta}_{I}) : \text{mmolar/second (summed)} \\
', \\
\text{astro_to_astro} = \text{Synapses(astrocytes, astrocytes,} \\
\text{ model=astro_to_astro_eqs)'}
\]

The above code bears the caveat of defining GJCs as unidirectional when they may not be so. This caveat can be easily overcome, specifying both a connection from astrocyte \(i\) to astrocyte \(j\) and a connection from \(j\) to \(i\), whenever we want to model bidirectional IP\(_3\) diffusion between neighboring astrocytes. For example, to connect astrocytes in a ring, where every astrocyte is connected to its neighbors (Figure 5A), we can make use of the connect method of the astro_to_astro object, and specify the following condition in terms of Brian 2 predefined pre- and post-synaptic indexes, \(i\) and \(j\) respectively, and the total number of elements in the
Figure 5. Astrocytes connected in a network. A Sample astrocyte network in a ring configuration with only one cell (in red) being exogenously stimulated. Connections between cells are bidirectional, and represent GJC-mediated coupling between neighboring astrocytes. B General Brian 2 modeling principle of astrocytic networks: GJC-mediated connections can be modeled by a Synapses object (ellipse). C Intercellular Ca\(^{2+}\) wave generation and propagation in a ring of 50 identical astrocytes mediated by stimulation of cell 25 (red trace). Parameters as in Table C.2 with \(F_{ex} = 0.09 \mu M s^{-1}\); and \(I_{bias} = 1 \mu M\).
pre-synaptic group $N_{pre}$

```python
astro_to_astro.connect('j == (i + 1) % N_pre or 'j == (i + N_pre - 1) % N_pre')
```

where the `%` operator implements the modulo (remainder) operation.

Figure 5C shows a snapshot of Ca$^{2+}$ dynamics of 50 astrocytes connected in a ring, where only the 25th cell is exogenously stimulated (red trace). The fact that all cells, for some $t > 0$, display Ca$^{2+}$ fluctuations, is a direct consequence of inclusion of intercellular IP$_3$ diffusion in our model. Such diffusion allows excess IP$_3$ from the stimulated cell to be redistributed by GJCs in the ring to other cells where it ultimately triggers CICR. It may also be appreciated how, in this example, bidirectional GJC communication allows for emergence of intercellular Ca$^{2+}$ waves that propagate both from and to the stimulated cell, as evidenced by wave fronts respectively oriented like ‘\’ or like ‘/’.

2.8 Coupled neuron and astrocyte networks

The examples discussed so far provide together all the ingredients to model complex networks of interacting neurons and astrocytes (Figure 6A). However, to realistically implement such networks we also need to specify the connections among neurons, synapses and astrocytes in the physical (Euclidean) space. In the following we show how to include space in such networks, limiting our focus here to planar networks for simplicity, although the outlined procedure can easily be extended to higher dimensions.

We start by adding two cell-specific parameters, $x$ and $y$, to each neuron which store the cell’s 2D spatial coordinates and initialize them so that neurons are arranged on a grid of $N_{rows}$ rows and $N_{cols}$ columns:

```python
neurons = NeuronGroup(N_e + N_i , model=neuron_eqs ,
threshold = 'v> V_th ', reset = 'v= V_r ',
refractory = 'tau_r ', method = 'euler ')
```

Furthermore, we also add a synapse-specific constant $astrocyte_index$ to the synapse’s equations, whose value will correspond to the index of the astrocyte that ensheathes a synapse:

```python
synapses_eqs = '''
# which astrocyte covers this synapse ?
astrocyte_index : integer (constant)
''',
```

---

3 Note that the expression has been split into two strings for better readability. Python automatically merges adjacent strings.
We finally need to define the effective connections between the different cells of the network. Overall there are four different types of connections: (i) connections between neurons which defines the actual synapses; (ii) connections from synapses to astrocytes, as pathways to trigger astrocyte activation; (iii) connections from astrocytes to synapses as routes for gliotransmission and thereby modulation of synaptic release; and ultimately, (iv) connections between astrocytes by GJCs. Here, for simplicity, we assume random connectivity between all neurons, independently of their spatial coordinates (as in Figure 1C). Furthermore, we make the assumption that only excitatory synapses can activate astrocytes and be modulated by them, restricting in this way our focus on the experimentally well-characterized pathway of closed-loop glutamatergic gliotransmission (Panatier et al., 2011; Perea and Araque, 2007). In particular, we specify which astrocyte is responsible for which excitatory synapse on the basis of the spatial position of postsynaptic neurons with respect to $N_a$ astrocytes ($N_a$) which, like neurons, are arranged on a regularly-spaced grid of $N_{rows}$ rows ($N_{rows}$) and $N_{cols}$ columns ($N_{cols}$), i.e.

$$N_{rows} = \text{int} (\sqrt{N_a})$$
$$N_{cols} = \frac{N_a}{N_{rows}}$$
$$\text{grid\_dist} = \frac{\text{size}}{N_{rows}}$$
$$\text{exc\_syn\_astro\_index} = (\text{int}(x_{post}/\text{grid\_dist}) + \text{int}(y_{post}/\text{grid\_dist}))$$

We then define the network of astrocytes:

```python
astro_eqs = '''
# The astrocyte position in space
x : meter (constant)
y : meter (constant)
'''
# Arrange astrocytes in a grid
astrocytes.x = '((i / N_{rows}) * \text{grid\_dist} - N_{rows}/2.0*\text{grid\_dist})$$
astrocytes.y = '((i % N_{rows}) * \text{grid\_dist} - N_{cols}/2.0*\text{grid\_dist})$$
```

Next, we connect the astrocytes with those synapses that they are supposed to ensheathe according to astrocyte_index, i.e.

```python
ecs_astro_to_syn = Synapses(astrocytes, exc_syn,
    'G_A_post = G_A_pre : mmolar (summed)')
ecs_astro_to_syn.connect('i == astrocyte_index_post')
ecs_syn_to_astro = Synapses(exc_syn, astrocytes,
    'Y_S_post = Y_S_pre/N_{incoming} : mmolar (summed)')
ecs_syn_to_astro.connect('astrocyte_index_pre == j')
```

Finally, we specify the connectivity of the astrocyte network. In this example, we introduce recurrent connections between astrocytes by GJCs, connecting each astrocyte to all other astrocytes found at the boundary of its anatomical domain, in line with the experimental observation that neighboring astrocytes are more likely to be connected than astrocytes that are far apart (Giaume et al., 2010; Pannasch and Rouach, 2013). Given that the diameter of astrocyte is between 50–130 µm (Chao et al., 2002), we consider an intermediate value of 75 µm, whereby:

```python
astro_to_astro.connect('i != j and '
    'sqrt((x_pre-x_post)**2 +
        (y_pre-y_post)**2) < 75*um')
```
Figure 6. Recurrent neuron-glial network. A Neuron-glial network model design in Brian 2. B Simulations of neuron-glia network for a rectangular-pulse increase of external current ($I_{ex}$, top panel). The raster plot (middle panel) shows the firing activity of 25% out of all excitatory (red) and inhibitory neurons (blue) of the network, and gliotransmitter release (green) from an equal fraction of astrocytes. The network-averaged firing rate is shown at the bottom. Neural activity dramatically changes from asynchronous low-firing activity to synchronous high-firing activity following gliotransmitter release from astrocytes during the period of high stimulation (2 ≤ $t$ < 4 s). External current: $I_{ex} = 100$ pA for $t < 2$ s or $t ≥ 4$ s; $I_{ex} = 120$ pA for 2 ≤ $t$ < 4 s. Neural and synaptic parameters as those in Figure 1 (see also Table C.1). Astrocyte parameters as in Table C.2 except for $O_β = 0.5$ µM s$^{-1}$; $O_δ = 1.2$ µM s$^{-1}$; and $I_{bias} = 0$. 
We present a simulation of our neuron-glia network in Figure 6B, where we show the raster plot of the firing activity of 25% of the excitatory (red) and inhibitory neurons (blue) of the network along with gliotransmitter release events from an equal fraction of astrocytes (green), in response to a transient increase of external stimulation (rectangular pulse in the top panel). Up to the onset of stimulation (i.e. \( t < 2\) s) there is no gliotransmitter release from astrocytes, therefore the network behaves as it would be expected for a neuronal network without the astrocyte component. It may be noted in fact how the raster plot of our network, and the ensuing dynamics of the total firing rate (bottom panel), show low-frequency population activity, similar to those reported in Figure 1C for our neuronal-only network model introduced in Sections 2.2 and 2.3. For \( 2 \leq t < 4\) s, the increase of external stimulation correlates with an increase in the firing rate of the whole network, as reflected by a denser raster plot during this period. In particular for \( t > 3.5\) s, the larger neuronal firing triggers gliotransmitter release from astrocytes and thus astrocytic modulation of excitatory synaptic transmission. Because this modulation is slow-decaying (Chapter 8), it outlasts the transient increase of external stimulation and changes neural firing once the external stimulation returns to its original value (at \( t = 4\) s). We can indeed clearly see how, for \( t > 4\) s, excitatory neurons are more synchronized in firing than for \( t < 2\) s, as a consequence of gliotransmission from astrocytes. This is just one example of the many possible ways astrocytes could actively shape neural activity, which has also been suggested to participate in the genesis of cortical UP and DOWN states (Fellin et al. 2012).

Conclusions

Computational approaches to model glial physiology are hampered by the lack of definitive experimental evidence and a missing comprehensive modeling framework that could tackle the many different scales of glial signaling. “Standard” glia models have yet to be identified, and neural simulator packages therefore do not ship such models as part of their pre-built model libraries. While in theory these libraries could be extended by individual researchers to add their preferred glia model, in practice this path is only open to experienced programmers.

In this chapter, we have shown how Brian 2’s simple syntax and versatility can offer a solution to these problems, providing an ideal tool to model glial physiology, and specifically the influence of astrocytes on neural activity. Brian 2’s syntax allows the researcher to flexibly describe models by using conventional mathematical notation instead of low-level programming code (Goodman and Brette 2008; Goodman et al. 2009; Stimberg et al. 2014). Moreover, Brian 2’s core data structure NeuronGroup, which describes a neuron by a set of ODEs, parameters, and actions that are triggered by conditions, provides a versatile framework that can be borrowed to also describe non-neuronal cell types such as astrocytes. Similarly, the Synapses data structure that, in purely neural simulations, represents chemical and electrical synapses that connect neurons, can also be used to model the interactions between astrocytes and synapses, as well as GJCs between astrocytes. Importantly, this flexibility does not come at the cost of computational efficiency: without any user interaction, Brian 2 employs a code generation approach that generates highly efficient code based on the user-provided high-level description (Goodman 2010). We hope that these arguments motivate newcomers as well as experienced researchers to experiment with Brian 2 in the future and use it to model glial physiology in their research, thereby contributing to the growth of this exciting emerging field of computational research.
Appendix A  Technical remarks on $Brian$ 2

$Brian$ 2 scripts are executed by default in the so-called “runtime mode”. This mode runs the simulation loop over the time steps in Python and executes chunks of target language code that have been generated from the model description provided by the user. The choice of target language depends on the user’s system; $Brian$ 2 will prefer to use the C++ programming language but, if the user does not have a working C++ compiler, will fall back to a pure Python-based simulation. A Python-based simulation will usually be significantly slower but can give comparable performance for big networks due to the use of vectorized computation ([Brette and Goodman] 2011). The advantage of the runtime mode is that the user has full control to combine the automatically generated simulation code with arbitrary hand-written Python code. This code could dynamically change aspects of the model during the run, or interact with it in other ways. For example it could read out the model’s state and hand it over to some code for visualization or terminate the simulation based on some criterion. This mode however involves a significant overhead per simulated time step, since the program flow constantly switches between Python and the individually-generated code chunks. For small-to-medium size networks for which computations during a single time step do not take long, this overhead can critically dominate the total runtime and lead to long simulation times.

To avoid this problem and allow more efficient simulations, $Brian$ 2 also offers an alternative mode called “standalone mode”. In this mode, the complete simulation code, including the main simulation loop, are written as a set of C++ files to disk which can then be compiled and executed as a single program. The resulting files are independent of the Python platform, so that the simulation could also be run on systems where Python may not be available (for example, in robotics). Moreover, if the user code complies to some specific conventions and does not run custom Python code during a simulation, then switching from runtime to standalone mode only requires the addition of a `set_device('cpp_standalone')` line to the simulation script; $Brian$ 2 then takes care of the whole process transparently. For further details, the reader is invited to see comments in individual examples files (Appendix B) and/or refer to the online $Brian$ 2 documentation.

Appendix B  Example files

The code for all the simulations presented in this chapter has been organized in multiple standalone example files as detailed in the following. Unless stated otherwise, all simulations start from zero initial conditions, except for $h(0) = 0.9$ and $x_S(0) = x_A(0) = 1$.

example_1_COBA.py

This file implements the simulation of the neuron-only network model of Figure 1. The simulation runs for 1 s with an integration time step of 0.1 ms. Out of all neurons, we distinguish between excitatory (exc_neurons) and inhibitory ones (inh_neurons), which give rise to excitatory synapses (exc_syn) and inhibitory synapses (inh_syn), connecting from the respective population to the full population. Because the dynamics of synaptic variables are updated only at incoming action potentials (i.e. (event-driven)), we can monitor the value of these variables only at the arrival time of action potentials but not in between. However, we can reconstruct the whole synaptic dynamics by recording synaptic variables immediately after each action potential (i.e. at $t^+_i$ with $i > 0$), which is achieved by specifying the keyword argument `when='after_synapses'` in the synaptic StateMonitor. For $t > t_i$, the solutions of the synapse’s
equations 4 and 5 then read (Tsodyks, 2005):

\[ u_S(t) = u_S(t_i^+) \exp \left( -\Omega_f(t - t_i) \right) \]

\[ x_S(t) = 1 + \left( x_S(t_i^+) - 1 \right) \exp \left( -\Omega_d(t - t_i) \right) \]

whereas synaptic release by the \( i \)th action potential at time \( t_i \) is given by

\[ r_S(t_i) = u_S(t_i^+) x_S(t_i^-) = \frac{u_S(t_i^+) x_S(t_i^+)}{1 - x_S(t_i^-)} \]

**example_2_gchi_astrocyte.py**

This code implements the synaptically-stimulated astrocyte model and related simulations of Figure 2. The astrocyte’s temporal dynamics in response to synaptic activity was simulated for 30s using the derivative-free Milstein integration method with a time step of 1 ms. In the deterministic limit of \( \xi(t) \to 0 \) in equation 13, the Milstein method reduces to the classical (forward) Euler method which is suitable, at sufficiently small time steps, to numerically solve dynamics of the deterministic astrocyte model, too. Synapses are stimulated by a train of periodic action potentials at rate \( f_0 = 0.5 \text{ Hz} \) (\( f_0 \), rate of generation of action potentials by presynaptic neurons) generated by

```python
source_neurons = NeuronGroup(1, 'dx/dt = f_0 : 1', threshold='x > 1', reset='x=0', method='euler')
```

**example_3_io_synapse.py**

This file implements the open-loop model of gliotransmission and the simulations shown in Figure 3. The code considers three synaptic connection between one presynaptic \( \text{source_neurons} \) and one postynaptic \( \text{target_neurons} \), built by passing \( n=3 \) as an argument to the \( \text{synapses.connect} \) method. We further consider two astrocytes stimulated by different \( I\_\text{bias} \) values, and connect them to synapses 2 and 3 respectively, leaving synapse 1 as it is (i.e. without gliotransmission). This is done by:

```python
ces_astro_to_syn.connect(j=i+1)
```

**example_4_synrel.py**

This code runs the closed-loop model of gliotransmission for simulations in Figure 4B. The code considers \( N_{\text{synapses}} \) neurons (\( \text{source_neurons} \)), each firing action potentials drawn from an independent, inhomogeneous Poisson process with a stepped rate specified in a \( \text{TimedArray} \), i.e.

\[ \text{rate.in} = \text{TimedArray([0.011, 0.11, 1.1, 11]*Hz, dt=5*second)} \]

\[ \text{source_neurons} = \text{PoissonGroup}(N_{\text{synapses}}, \text{rates='\text{rate.in(t)'}')} \]

\[ \text{target_neurons} = \text{NeuronGroup}(N_{\text{synapses}}, '') \]

The \( \text{target_neurons} \) are used to build \( N_{\text{synapses}} \) multi-synaptic connections from the \( \text{source_neurons} \), with each connection constituted of three synapses. Out of these three synapses, the first one is connected with its own astrocyte and is, in turn, modulated by gliotransmitters released from this latter (closed-loop scenario); the second one is modulated by gliotransmitters released from another astrocyte (open-loop scenario); the third one is left as it is (scenario without gliotransmission). Since this is repeated for all \( N_{\text{synapses}} \), and overall we have \( N_{\text{astro}} \) different scenarios of gliotransmission (open-loop vs. closed-loop), we consider \( N_{\text{astro}} \times N_{\text{synapses}} \) astrocytes in total, and connect them accordingly with \( N_{\text{synapses}} \times (N_{\text{astro}}+1) \) synapses as elucidated in Section 2.6.
example_4_rsmean.py

The file provides the code to build the synaptic transfer characteristics in Figure 4C in terms of average synaptically-released neurotransmitter resources for different input rates of (presynaptically) incoming action potentials.

eexample_5_astro_ring.py

This code implements the astrocyte ring model in Figure 5. The simulation runs for 4000 s with a time step of 50 ms. Calcium concentrations shown in Figure 5C were normalized by their maximum.

eexample_6_COBA_with_astro.py

This file runs the simulation of the recurrent neuron-glial network in Figure 6. To stimulate the network by a time-varying external current we multiply $I_{\text{ex}}$ in neuron_eqs on page 4 by $\text{stimulus} = \text{TimedArray}(\{1.0, 1.2, 1.0, 1.0\}, \text{dt} = 2*\text{second})$. Neurons are placed on a square lattice of size $3.75 \times 3.75 \text{mm}$ at $50 \mu\text{m}$ distance from each other. For $t = 0$ we set $C = I = 0.01 \mu\text{M}$.

Appendix C  Model parameters used in the simulations

The following tables report constants that correspond to the model parameters used in the simulations presented in this chapter. Simulation-specific parameters are marked by ‘†’ and are reported in respective figure captions instead.

C.1 Neurons and synapses

| Symbol | Name in code | Value | Units | Description |
|--------|--------------|-------|-------|-------------|
| $C_m$  | C_m         | 198 pF |       | Membrane capacitance |
| $E_l$  | E_l         | -60 mV |       | Leak reversal potential |
| $g_l$  | g_l         | 9.99 nS|       | Leak conductance |
| $I_{\text{ex}}$ | I_ex | † pA |       | External current |
| $V_r$  | V_r         | -60 mV |       | Reset potential |
| $V_\theta$ | V_th | -50 mV |       | Firing threshold |
| $\tau_r$ | tau_r | 5 ms |       | Refractory period |

Synapse parameters

| Symbol | Name in code | Value | Units | Description |
|--------|--------------|-------|-------|-------------|
| $\Omega_d$ | Omega_d | 2 s$^{-1}$ |       | Synaptic depression rate |
| $\Omega_f$ | Omega_f | 3.33 s$^{-1}$ |       | Synaptic facilitation rate |
| $U_0$ (U_0) | U_0___star (U_0) | 0.6 – |       | Resting synaptic release probability |
| $Y_T$ | Y_T | 500 mM |       | Total vesicular neurotransmitter concentration |
| $\rho_c$ | rho_c | 0.005 – |       | Synaptic vesicle-to-extracellular space volume ratio |
| $\Omega_c$ | Omega_c | 40 s$^{-1}$ |       | Neurotransmitter clearance rate |
| $w_e$ | w_e | 50 pS |       | Excitatory synaptic conductance |
| $w_i$ | w_i | 1 nS |       | Inhibitory synaptic conductance |
| $\tau_e$ | tau_e | 5 ms |       | Excitatory synaptic time constant |
| $\tau_i$ | tau_i | 10 ms |       | Inhibitory synaptic time constant |
| $E_e$ | E_e | 0 mV |       | Excitatory synaptic reversal potential |
| $E_i$ | E_i | -80 mV |       | Inhibitory synaptic reversal potential |

Presynaptic receptors

| Symbol | Name in code | Value | Units | Description |
|--------|--------------|-------|-------|-------------|
| $O_G$  | O_G         | 1.5 $\mu\text{M}^{-1}\text{s}^{-1}$ |       | Agonist binding (activating) rate |
| $\Omega_G$ | Omega_G | 0.5 min$^{-1}$ |       | Agonist release (deactivating) rate |
C.2 Astrocytes

| Symbol | Name in code | Value | Units | Description |
|--------|--------------|-------|-------|-------------|
| C_T   | C_T          | 2     | µm    | Total cell free Ca^{2+} content |
| ρ_A   | rho_A        | 0.18  |       | ER-to-cytoplasm volume ratio |
| d_1   | d_1          | 0.13  | µm    | IP_3 dissociation constant |
| d_2   | d_2          | 1.05  | µm    | Ca^{2+} inactivation dissociation constant |
| d_3   | d_3          | 0.9434| µm    | IP_3 dissociation constant |
| d_5   | d_5          | 0.08  | µm    | Ca^{2+} activation dissociation constant |
| O_2   | O_2          | 0.2   | µm s^{-1} | IP_3 binding rate for Ca^{2+} inhibition |
| Ω_C   | Omega_C      | 6     | s^{-1} | Maximal rate of Ca^{2+} release by IP_3Rs |
| Ω_L   | Omega_L      | 0.1   | s^{-1} | Maximal rate of Ca^{2+} leak from the ER |
| O_ρ   | O_ρ          | 0.9   | µm s^{-1} | Maximal Ca^{2+} uptake rate by SERCAs |
| K_P   | K_P          | 0.05  | µm    | Ca^{2+} affinity of SERCAs |

IP_3 signaling

| Symbol | Name in code | Value | Units | Description |
|--------|--------------|-------|-------|-------------|
| O_β   | O_beta       | †     | µm s^{-1} | Maximal rate of IP_3 production by PLCβ |
| O_δ   | O_delta      | 0.6   | µm s^{-1} | Maximal rate of IP_3 production by PLCδ |
| κ_δ   | kappa_delta  | 1.5   | µm    | Inhibition constant of PLCδ by IP_3 |
| K_δ   | K_delta      | 0.1   | µm    | Ca^{2+} affinity of PLCδ |
| O_3K  | O_3K         | 4.5   | µm s^{-1} | Maximal rate of IP_3 degradation by IP_3-3K |
| K_3K  | K_3K         | 1     | µm    | IP_3 affinity of IP_3-3K |
| K_D   | K_D          | 0.7   | µm    | Ca^{2+} affinity of IP_3-3K |
| Ω_5P  | Omega_5P     | 0.05  | s^{-1} | Maximal rate of IP_3 degradation by IP 5P |

Metabotropic receptor kinetics

| Symbol | Name in code | Value | Units | Description |
|--------|--------------|-------|-------|-------------|
| O_N   | O_N          | 0.3   | µM † s^{-1} | Agonist binding rate |
| Ω_N   | Omega_N      | 0.5   | s^{-1} | Maximal inactivation rate |
| K_KC  | K_KC         | 0.5   | µm    | Ca^{2+} affinity of PKC |
| ζ     | zeta         | 10    |       | Maximal reduction of receptor affinity by PKC |

IP_3 stimulation & diffusion

| Symbol | Name in code | Value | Units | Description |
|--------|--------------|-------|-------|-------------|
| F_ex  | F_ex         | 2     | µm s^{-1} | Maximal exogenous IP_3 flow |
| I_bias| I_bias       | †     | µm    | External IP_3 drive |
| F     | F            | 0.09  | µm s^{-1} | GJIC IP_3 permeability |
| I_θ   | I_Theta      | 0.3   | µm    | Threshold gradient for IP_3 diffusion |
| ω_1   | omega_1      | 0.05  | µm    | Scaling factor of diffusion |

Gliotransmission

| Symbol | Name in code | Value | Units | Description |
|--------|--------------|-------|-------|-------------|
| C_θ   | C_Theta      | 0.5   | µm    | Ca^{2+} threshold for exocytosis |
| G_T   | G_T          | 200   | mM    | Total vesicular gliotransmitter concentration |
| Ω_A   | Omega_A      | 0.6   | s^{-1} | Gliotransmitter recycling rate |
| U_A   | U_A          | 0.6   |       | Gliotransmitter release probability |
| ρ_e   | rho_e        | 0.0065|       | Astrocytic vesicle-to-extracellular volume ratio |
| Ω_e   | Omega_e      | 60    | s^{-1} | Gliotransmitter clearance rate |
| α     | alpha        | 0     |       | Gliotransmission nature |
References

Abbott, L. F. (2008). Theoretical neuroscience rising. *Neuron*, 60(3):489–495.

Aleksin, S. G., Zheng, K., Rusakov, D. A., and Savtchenko, L. P. (2017). ARACHNE: A neural-neuroglial network builder with remotely controlled parallel computing. *PLoS Comput. Biol.*, 13(3):e1005467.

Araque, A., Carmignoto, G., Haydon, P. G., Oliet, S. H. R., Robitaille, R., and Volterra, A. (2014). Gliotransmitters travel in time and space. *Neuron*, 81(4):728–739.

Beenhakker, M. P. and Huguenard, J. R. (2010). Astrocytes as gatekeepers of GABA<sub>B</sub> receptor function. *The Journal of Neuroscience*, 30(45):15262–15276.

Bezzi, P. and Volterra, A. (2001). A neuron-glia signalling network in the active brain. *Curr. Opinion Neurobiol.*, 11:387–394.

Brette, R. (2012). On the design of script languages for neural simulation. *Network*, 23(4):150–156.

Brette, R. and Goodman, D. F. M. (2011). Vectorized algorithms for spiking neural network simulation. *Neural Comput*, 23(6):1503–1535.

Brette, R., Rudolph, M., Carnevale, T., Hines, M., Beeman, D., Bower, J. M., Diesmann, M., Morrison, A., Goodman, P. H., Harris, J. F. C., Zirpe, M., Natschläger, T., Pecevski, D., Ermentrout, B., Djurfeldt, M., Lansner, A., Rochel, O., Vieville, T., Muller, E., Davison, A. P., El Boustani, S., and Destexhe, A. (2007). Simulation of networks of spiking neurons: a review of tools and strategies. *J Comput Neurosci*, 23(3):349–398.

Brunel, N. (2000). Dynamics of sparsely connected networks of excitatory and inhibitory spiking neurons. *J Comput Neurosci*, 8(3):183–208.

Cannon, R. C., Gewaltig, M.-O., Gleeson, P., Bhalla, U. S., Cornelis, H., Hines, M. L., Howell, F. W., Muller, E., Stiles, J. R., Wils, S., and Schutter, E. D. (2007). Interoperability of Neuroscience Modeling Software: Current Status and Future Directions. *Neuroinform*, 5(2):127–138.

Carnevale, N. T. and Hines, M. L. (2006). *The NEURON Book*. Cambridge University Press.

Chao, T. I., Rickmann, M., and Wolff, J. R. (2002). The synapse-astrocyte boundary: an anatomical basis for an integrative role of glia in synaptic transmission. In Volterra, A., Magistretti, P. J., and Haydon, P. G., editors, *The Tripartite Synapse: Glia in Synaptic Transmission*, chapter 1, pages 3–23. Oxford University Press, New York.

Dayan, P. and Abbott, L. F. (2001). *Theoretical neuroscience*, volume 806. Cambridge, MA: MIT Press.

De Pittà, M., Brunel, N., and Volterra, A. (2016). Astrocytes: orchestrating synaptic plasticity? *Neuroscience*, 323:43–61.

De Pittà, M., Goldberg, M., Volman, V., Berry, H., and Ben-Jacob, E. (2009a). Glutamate-dependent intracellular calcium and IP<sub>3</sub> oscillating and pulsating dynamics in astrocytes. *J. Biol. Phys.*, 35:383–411.
De Pittà, M., Volman, V., Berry, H., and Ben-Jacob, E. (2011). A tale of two stories: astrocyte regulation of synaptic depression and facilitation. PLoS Comput. Biol., 7(12):e1002293.

De Pittà, M., Volman, V., Berry, H., Parpura, V., Liaudet, N., Volterra, A., and Ben-Jacob, E. (2013). Computational quest for understanding the role of astrocyte signaling in synaptic transmission and plasticity. Front. Comp. Neurosci., 6:98.

De Pittà, M., Volman, V., Levine, H., and Ben-Jacob, E. (2009b). Multimodal encoding in a simplified model of intracellular calcium signaling. Cogn. Proc., 10(S1):55–70.

De Pittà, M., Volman, V., Levine, H., Pioggia, G., De Rossi, D., and Ben-Jacob, E. (2008). Coexistence of amplitude and frequency modulations in intracellular calcium dynamics. Phys. Rev. E, 77(3):030903(R).

Engelman, H. S. and MacDermott, A. B. (2004). Presynaptic ionotropic receptors and control of transmitter release. Nature Rev., 5:135–145.

Falcke, M. (2004). Reading the patterns in living cells: the physics of Ca$^{2+}$ signaling. Adv. Phys., 53(3):255–440.

Fellin, T., Ellenbogen, J. M., De Pittà, M., Ben-Jacob, E., and Halassa, M. (2012). Astrocyte regulation of sleep circuits: experimental and modeling perspectives. Front. Comp. Neurosci., 6(65).

Fuhrmann, G., Segev, I., Markram, H., and Tsodyks, M. (2002). Coding of temporal information by activity-dependent synapses. J. Neurophysiol., 87:140–148.

Gewaltig, M.-O. and Diesmann, M. (2007). NEST (NEural Simulation Tool). Scholarpedia, 2(4):1430.

Giaume, C., Koulakoff, A., Roux, L., Holeman, D., and Rouach, N. (2010). Astroglial networks: a step further in neuroglial and gliovascular interactions. Nat. Rev. Neurosci., 11(2):87–99.

Goldberg, M., De Pittà, M., Volman, V., Berry, H., and Ben-Jacob, E. (2010). Nonlinear Gap junctions enable long-distance propagation of pulsating calcium waves in astrocyte networks. PLoS Comput. Biol., 6(8):e1000909.

Goodman, D. and Brette, R. (2008). Brian: a simulator for spiking neural networks in Python. Front. Neuroinform., 2:5.

Goodman, D. F. M. (2010). Code generation: A strategy for neural network simulators. Neuroinform, 8(3):183–196.

Goodman, D. F. M., Brette, R., Goodman, D. F. M., and Brette, R. (2009). The Brian simulator. Front. Neurosci., 3.

Hunter, J. (2007). Matplotlib: A 2D Graphics Environment. Computing in Science Engineering, 9(3):90–95.

Kloeden, P. E. and Platen, E. (1992). Numerical Solution of Stochastic Differential Equations.

Lallouette, J., De Pittà, M., Ben-Jacob, E., and Berry, H. (2014). Sparse short-distance connections enhance calcium wave propagation in a 3D model of astrocyte networks. Front. Comp. Neurosci., 8:45.
Li, Y. and Rinzel, J. (1994). Equations for InsP$_3$ receptor-mediated $[\text{Ca}^{2+}]_i$ oscillations derived from a detailed kinetic model: A Hodgkin-Huxley like formalism. *J. Theor. Biol.*, 166:461–473.

Martín, R., Bajo-Grañeras, R., Moratalla, R., Perea, G., and Araque, A. (2015). Circuit-specific signaling in astrocyte-neuron networks in basal ganglia pathways. *Science*, 349(6249):730–734.

McDougal, R. A., Morse, T. M., Carnevale, T., Marenco, L., Wang, R., Migliore, M., Miller, P. L., Shepherd, G. M., and Hines, M. L. (2017). Twenty years of ModelDB and beyond: building essential modeling tools for the future of neuroscience. *J Comput Neurosci*, 42(1):1–10.

Mil’stejn, G. (1975). Approximate integration of stochastic differential equations. *Theory of Probability & Its Applications*, 19(3):557–562.

Nadkarni, S., Jung, P., and Levine, H. (2008). Astrocytes optimize the synaptic transmission of information. *PLoS Comput. Biol.*, 4(5):e1000088.

Naeem, M., McDaid, L., Harkim, J., Wade, J. J., and Marsland, J. (2015). On the role of astroglial syncytia in self-repairing spiking neural networks. *IEEE Trans. Neural Networks and Learning Systems*, page In press.

Navarrete, M. and Araque, A. (2010). Endocannabinoids potentiate synaptic transmission through stimulation of astrocytes. *Neuron*, 68(1):113–126.

Nimmerjahn, A. (2009). Astrocytes going live: advances and challenges. *J. Physiol.*, 587:1639–1647.

Panatier, A., Vallée, J., Haber, M., Murai, K., Lacaille, J., and Robitaille, R. (2011). Astrocytes are endogenous regulators of basal transmission at central synapses. *Cell*, 146:785–798.

Pannasch, U. and Rouach, N. (2013). Emerging role for astroglial networks in information processing: from synapse to behavior. *Trends in Neurosciences*, 36(7):405–417.

Perea, G. and Araque, A. (2007). Astrocytes potentiate transmitter release at single hippocampal synapses. *Science*, 317:1083–1086.

Pinheiro, P. S. and Mulle, C. (2008). Presynaptic glutamate receptors: physiological functions and mechanisms of action. *Nat. Rev. Neurosci.*, 9:423–436.

Poskanzer, K. E. and Yuste, R. (2016). Astrocytes regulate cortical state switching in vivo. *Proceedings of the National Academy of Sciences*, 113(19):E2675–E2684.

Savtchenko, L. P. and Rusakov, D. A. (2014). Regulation of rhythm genesis by volume-limited, astroglia-like signals in neural networks. *Phil. Trans. Royal Soc. B: Biological Sciences*, 369(1654):20130614.

Scemes, E. and Giaume, C. (2006). Astrocyte calcium waves: What they are and what they do. *Glia*, 54:716–725.

Shuai, J.-W. and Jung, P. (2002). Stochastic properties of $\text{Ca}^{2+}$ release of inositol 1,4,5-trisphosphate receptor clusters. *Biophysical Journal*, 83(1):87–97.

Stiles, J. R., Bartol, T. M., et al. (2001). Monte Carlo methods for simulating realistic synaptic microphysiology using MCell. In Schutter, E. D., editor, *Computational neuroscience: realistic modeling for experimentalists*, pages 87–127. CRC Press, Boca Raton, FL.
Stimberg, M., Goodman, D. F. M., Benichoux, V., and Brette, R. (2014). Equation-oriented specification of neural models for simulations. *Frontiers in Neuroinformatics*, 8.

Tsodyks, M. (2005). Activity-dependent transmission in neocortical synapses. In Chow, C. C., Gutkin, B. D., H., C. M., and Dalibard, J., editors, *Methods and Models in Neurophysics*, pages 245–265. Elsevier.

Tsodyks, M., Pawelzik, K., and Markram, H. (1998). Neural networks with dynamic synapses. *Neural Computation*, 10(4):821–835.

van der Walt, S., Colbert, S., and Varoquaux, G. (2011). The NumPy array: A structure for efficient numerical computation. *Computing in Science Engineering*, 13(2):22–30.

Volman, V., Ben-Jacob, E., and Levine, H. (2007). The astrocyte as a gatekeeper of synaptic information transfer. *Neural computation*, 19(2):303–326.

Wade, J. J., McDaid, L. J., Harkin, J., Crunelli, V., and Kelso, J. A. S. (2011). Bidirectional coupling between astrocytes and neurons mediates learning and dynamic coordination in the brain: a multiple modeling approach. *PLoS One*, 6(12):e29445.

Wallach, G., Lallouette, J., Herzog, N., De Pittà, M., Ben-Jacob, E., Berry, H., and Hanein, Y. (2014). Glutamate-mediated astrocytic filtering of neuronal activity. *PLoS Comput. Biol.*, 10(12):e1003964.

Zorec, R., Araque, A., Carmignoto, G., Haydon, P., Verkhratsky, A., and Parpura, V. (2012). Astroglial excitability and gliotransmission: An appraisal of Ca^{2+} as a signaling route. *ASN Neuro*, 4(2):e00080.