# Synaptic Maturation Project

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#### 1 Intro

The model follows the following framework, with 3 possible states and the transitions between them:

$$\begin{array}{c} \text{Potential Synapse Pool} \xrightarrow[\overline{\text{elimination}}]{\overline{\text{constant Population}}} & \text{Immature Population} \\ \hline \\ \hline \\ \hline \end{array} \\ \begin{array}{c} \underline{\text{maturation}} \\ \hline \\ \hline \end{array} \\ \text{Mature Population} \\ \end{array}$$

For synapses in the Mature Population, we also model "synaptic size". When a synapse becomes mature, we initiate a size of 0, and then simulate the stochastic dynamics of the synaptic weight using a Kesten process

$$x_{t+1} = \epsilon_t x_t + \eta_t$$

where  $x_t$  is the synaptic size at time t and  $\epsilon_t$  and  $\eta_t$  are random variables drawn from some distribution (we use Gaussian, with the mean of  $\epsilon$  near 1 — or just below — and the mean of  $\eta$  near 0, to ensure a stationary distribution).

## 2 Model 1: Random Walks

We first model this setup using random walks (with Gillespie's algorithm) and predefined transition probabilities c, e, m, i, for creation, elimination, maturation and immaturation (dematuration) respectively. We set parameters such as potential synapse pool size, time, and Kesten process time step. When a synapse "dematures", we adopt a simplified approach by removing those synapses of size 0. This random walk model has the advantage that its stochastic nature captures the randomness inherent in biological processes, providing a realistic simulation of individual synapse behaviour. However, it is more computationally expensive and less analytically insightful (being reliant on simulations).

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Below is an example simulation of this model, with parameters total_time = 100.0 epsilon, eta = 1.0, 0.0 total_pool_size = 1000 sigma_epsilon, sigma_eta = .5, .5 c, m, e, i = 0.2,0.2,0.01,0.05 kesten_timestep = 0.01
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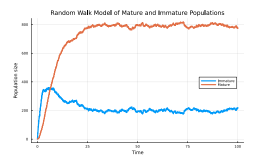


Figure 1: Single simulation of populations using Gillespie's algorithm

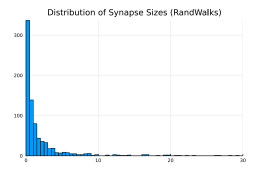


Figure 2: Final distribution of synaptic sizes

We can also keep track of the synapse sizes over time:

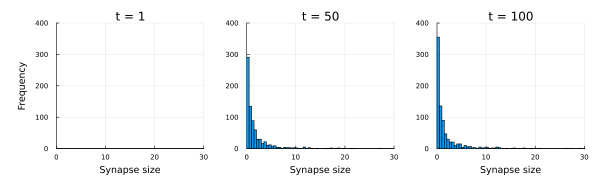


Figure 3: Evolution of the distribution of synaptic sizes over time.

# 3 Model 2: Differential Equations

#### 3.1 Setup

We then defined a set of differential equations to model the setup in another way. The equations are as follows:

$$\frac{dN_I}{dt} = cN_P - (e+m)N_I + iN_M$$
 
$$\frac{dN_M}{dt} = mN_I - iN_M$$
 
$$\frac{dN_P}{dt} = eN_I - cN_P$$

where c, e, m, i are as before, and  $N_I, N_M, N_P$  denote the number of Immature, Mature, and Potential (pool) synapses. Again, when a synapse "dematures", we remove those synapses of size 0. This differential equations model preserves the stochasticity of synapse sizes (with the Kesten process), but loses the stochastic nature of the state transitions and models average population behaviour. However, it is more computationally efficient and lends itself more easily to analysis.

Using the same parameter values as in Section 2, this gives:

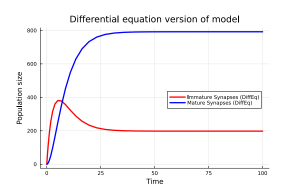


Figure 4: Populations solution using differential equations

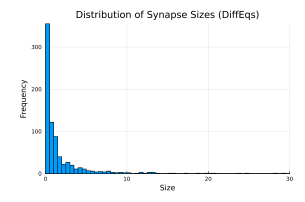


Figure 5: Final distribution of synaptic sizes

This closely matches the random walk version. To check, let us run multiple trials.

## 3.2 Multiple Trials

We run the above models over N trials (e.g. N=10 below) and this produces:

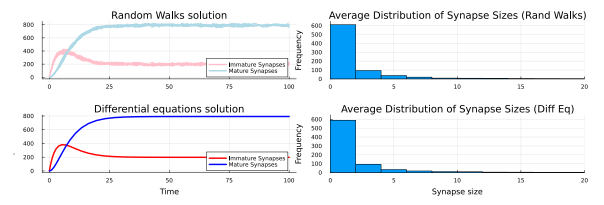


Figure 6: Multiple trials of Random Walks (top) and Differential Equations (bottom).

#### 3.3 Steady State Solutions to the Differential Equations

Taking the differential equations

$$\frac{dN_I}{dt} = cN_P - (e+m)N_I + iN_M$$

$$\frac{dN_M}{dt} = mN_I - iN_M$$

$$\frac{dN_P}{dt} = eN_I - cN_P$$

we can find the final steady state values of  $N_I, N_m, N_P$  by letting the derivatives equal 0 and solving.

This gives

$$0 = cN_P - (e+m)N_I + iN_M \Rightarrow cN_P = (e+m)N_I - iN_M$$
$$0 = mN_I - iN_M \Rightarrow mN_I = iN_M$$
$$0 = eN_I - cN_P \Rightarrow eN_I = cN_P$$

We also have that  $N_I + N_M + N_P = \text{total}$ . This gives us:

$$N_I + \frac{m}{i}N_I + \frac{e}{c}N_I = \text{total}$$

$$\frac{i}{m}N_M + N_M + \frac{ei}{cm}N_M = \text{total}$$

$$\frac{c}{e}N_P + \frac{cm}{ei}N_P + N_P = \text{total}$$

which yields the solutions:

$$N_{I} = \frac{\text{total}}{(1 + \frac{m}{i} + \frac{e}{c})}$$

$$N_{M} = \frac{\text{total}}{(1 + \frac{i}{m} + \frac{ei}{cm})}$$

$$N_{P} = \frac{\text{total}}{(1 + \frac{c}{e} + \frac{cm}{ei})}$$

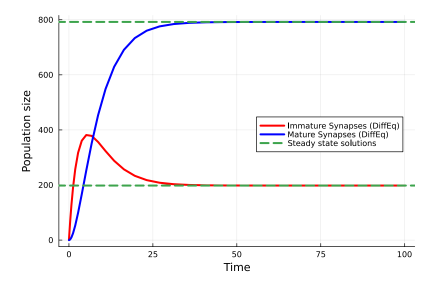


Figure 7: Differential equations model with steady state solutions

# 3.4 Phase Plane (Vector Field)

Let us plot the **phase plane** of the system. This is a graphical representation of a dynamical system with two state variables. Each point on the plane represents a specific state of the system, and trajectories show how these states evolve over time. A vector field on the phase plane consists of arrows at each point indicating the direction and speed of the system's evolution at that state.

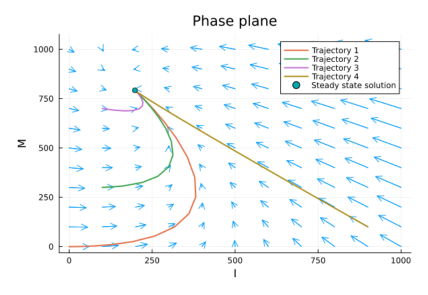


Figure 8: Phase plane of immature (x axis) vs mature (y axis) populations over time. The coloured lines show the different trajectories of populations with different initial values.

We can also plot the nullclines, i.e. the lines where the derivatives are zero. So for the first, we have that

$$cN_P - (e+m)N_I + iN_M = 0 \Rightarrow N_M = \frac{\text{total } c - cN_I - (e+m)N_I}{c - i}$$

and for the second

 $mN_I - iN_M = 0 \Rightarrow N_M = \frac{m}{i}N_I$ 

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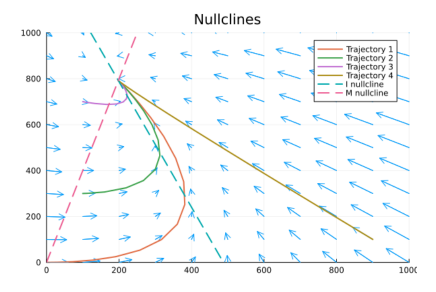


Figure 9: Here we plot the nullclines of the system. Where they intersect is the steady state solution.

# 4 Weight-Dependent Random Walk Model

Above we simplified the approach by just removing the mature synapses with size 0, but we next model the mature-to-immature transition in a weight-dependent manner. We do this by defining a probability function over synapse size. Realistically, this must capture the fact that smaller synapses are more likely to demature (and large synapses are very unlikely to demature). This can be modelled simply with an exponential function

 $p(s \text{ dematuring }) = Ae^{-\frac{s}{\lambda}}$ 

where s is the synapse size, and  $A, \lambda$  are parameters determining the exponential function. e.g.,

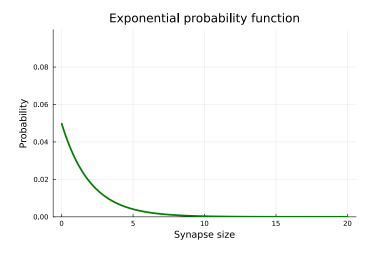


Figure 10: Exponential function for  $A=0.05, \lambda=2$ . Here we have A=0.05 as this was the value of i used in previous simulations.

Adopting this in a constant timestep stochastic state transition model, we get the following:



Figure 11: Single simulation of populations using weight dependent dematuring probabilities

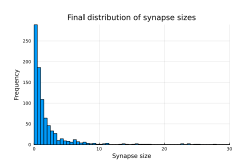


Figure 12: Final distribution of synaptic sizes.

# 5 Weight-Dependent Differential Equation Model

We similarly use this weight dependent idea in a differential equations setup, where we determine the rate of mature-to-immature by the exponential pdf. This gives:

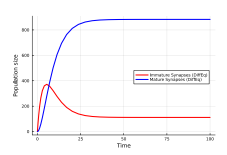


Figure 13: Differential Equation solution of populations using weight dependent dematuring probabilities

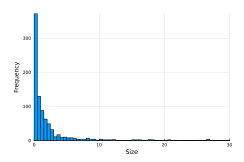
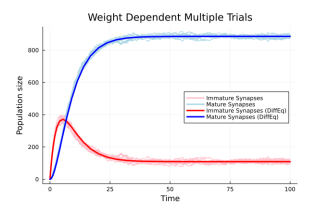


Figure 14: Final distribution of synaptic sizes.

Running this over some trials and comparing the probabilistic version with differential equations:



# 6 Going Forward

### 6.1 Calculating the average mature-to-immature fraction

The plan for the next step is to try make this a bit more analytically or computationally tractable. The idea is to work out the stationary limiting distribution  $p_1$  of the synapse sizes (from the Kesten process, subject to certain conditions), and using the exponential weight-dependent probability function  $p_2$ , calculate the "average" fraction of synapses that make the mature-to-immature transition with the integral from 0 to  $\infty$  of the product of  $p_1$  and  $p_2$ . This may be calculable analytically (I have yet to work it out, seems hard), but if not, it can be done numerically (e.g. Julia's QuadGK package was used below). This "average" value can then be implemented in a probabilistic state transition model, or in the differential equations model.

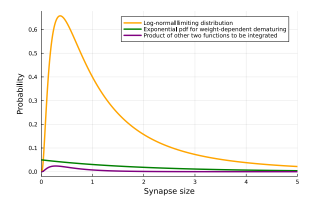


Figure 15: Limiting log-normal synapse size distribution (orange), exponential weight-dependent dematuring pdf (green), and their product (purple)

Using the distributions given above,  $\int_0^\infty p_1 p_2 \approx 0.0191$  (QuadGK numerical integration package). Implementing this value in the state transition model from Section 5 gives the below simulation:

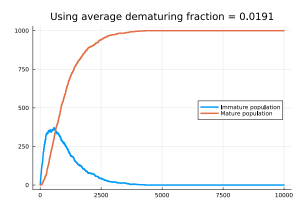


Figure 16: State transition model using constant average dematuring fraction of 0.0191

### 6.2 Incorporating developmental periods

The above models outlined above could be considered to only deal with a specific age in brain development because the parameters  $c, m, e, \lambda, \sigma_{\epsilon}, \sigma_{\eta}$  are all fixed. Ultimately we want these to vary across the developmental trajectory. We will do this either by

- Fitting the model parameters separately to the data at each developmental timepoint recorded
- Fitting some parametric function for each parameter to model its smooth change across development (i.e. fitting all the developmental timepoints in one go).