

### UNIVERSITY MASTER'S DEGREE

### PHYSICAL TECHNOLOGY: RESEARCH AND APPLICATIONS

### MASTER'S THESIS

Exploring the relationship between Hawkes processes and self-organized criticality in living systems

#### Antonio Rivas Blanco

Tutor(s): Jorge Hidalgo Aguilera and Serena di Santo

Line of research: Modelling of complex systems and their interdisciplinary applications

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Tutor 1: Jorge Hidalgo Aguilera

Tutor 2: Serena di Santo

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Tutor 1: Jorge Hidalgo Aguilera

Tutor 2: Serena di Santo

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Nombre y apellidos: Antonio Rivas Blanco

DNI: 49832223D

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## Abstract

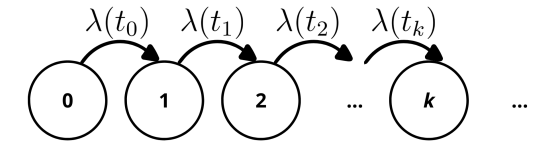
### Chapter 1

### Introduction

#### 1.1 Point processes

Within the large framework of complex systems, stochastic processes lend us a hand to decypher properties of living systems, bridging randomness with structured behaviour. This processes are used to model the dynamics of systems which evolve randomly in time. This is why they are ideal for describing natural phenomena such as the spread of diseases [1], social networks [2] or ecological systems [3]. Mathematically, a stochastic process is a collection of random variables [4], generally ordered in time  $\{X_t\}_{t\in T}$ , where t is the time and  $X_t$  is the system state at time t. T is the time index set, which can be discrete or continuous, in this work we will focus on the discrete case because we are interested in the study of point (Hawkes) processes for modeling neurons.

Point processes are a type of stochastic process that describe the occurrence of events in time or space. We will be interested in time point processes because we are going to model the spiking activity of neurons. For our purposes, they will be characterized by two parameters, the time of occurrence of the events  $t_k$  and the intensity or rate of occurrence of these events  $\lambda$ . This rate tell us how likely is that an event occurs at time t given the history of the process (probability density function, PDF) as pictured in Figure 1.1.

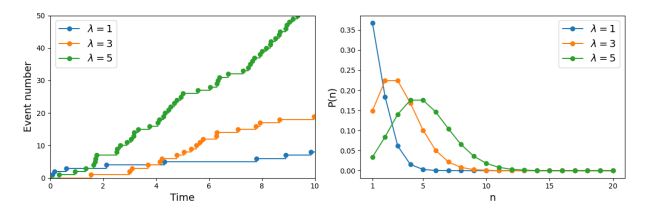


**Figure 1.1.** Representation of a point process. The intensity function  $\lambda(t)$  is a time-dependent function.

In general, the rate is a function of the history of the process, which makes the process non-Markovian, but in our case, it will be a Markovian process, which means that the rate depends only on the last event that occurred as we will see. An example of a Markovian point process is the Poisson process, which is a simple and one of the most studied point processes because they are present in many everyday situations such as the arrival of customers at a store, occurrence of defects on a Production line. They are also present in some physics phenomena, for instance, the decay of radioactive particles or the arrival of photons at a detector. These processes are characterized by a rate of occurrence of events  $\lambda$ . The dynamics of these processes are described by the Poisson distribution which is the probability distribution of a random variable N such that the probability that N = n is:

$$P(N=n) = \frac{\lambda^n}{n!} e^{-\lambda}.$$
 (1.1)

Furthermore, the mean value and the variance of the distribution are also equal to  $\lambda$ . Poisson processes can be homogeneous or inhomogeneous, depending on whether the rate is constant or time-dependent. In Figure 1.2 we can see an example of a homogeneous Poisson process.



**Figure 1.2.** Left: event number in time for different rates. Right: Probability of having a certain number of events for different rates.

### 1.2 Hawkes processes

On the other hand if we consider a non-homogeneous Poisson process, the rate is a function of time,  $\lambda(t)$ , which is the case of the Hawkes process. The rate can be written in several ways [5, 6, 7, 8]. We will use the the expression from [5]:

$$\lambda(t|t_1,\dots,t_k) = \mu + n\sum_{i=1}^k \phi(t-t_i),$$
 (1.2)

where  $\mu$  is the background rate of a homogeneous Poisson process, n is a parameter that controls the strength the self-excitation, and  $\phi(t)$  is the kernel function that describes the influence of the past events on the rate of occurrence of the events. The kernel function is a non-negative and monotonically non-increasing function that integrates to 1. Typical choices for the kernel function are the exponential or the power-law functions. In this work we will focus on the exponential kernel. From Eq 1.2 we can see that the rate depends on the history of the process, making it non-Markovian in general, but

with an exponential kernel, the process becomes Markovian. The kernel function can be written as:  $\phi(t) = \sum_{t_i < t} \alpha e^{-\beta(t-t_i)}$  so the rate becomes:

$$\lambda(t) = \mu + \sum_{t_i < t} \alpha e^{-\beta(t-t_i)}$$

$$= \mu + \sum_{\substack{t_i < t_k \\ t_{k: last event}}} \alpha e^{-\beta(t-t_k+t_k-t_i)}$$

$$= \mu + e^{-\beta(t-t_k)} \sum_{\substack{t_i < t_k \\ \lambda(t_k)}} \alpha e^{-\beta(t_k-t_i)}$$

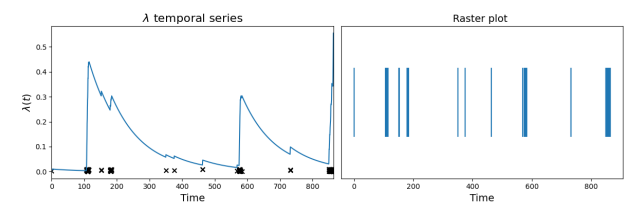
$$= \mu + e^{-\beta(t-t_k)} \left(\lambda(t_k) - \mu + n\right).$$

$$(1.3)$$

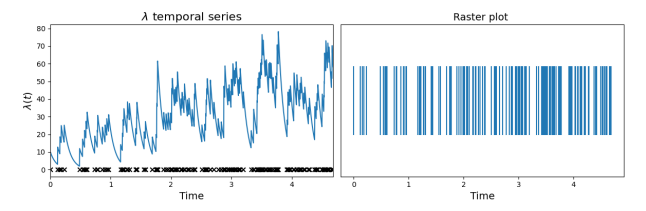
Where we have used the following expression for the rate of the Hawkes process at time  $t_k$ :

$$\lambda(t_k) = \mu + \sum_{t_i < t_k} \alpha e^{-\beta(t_k - t_i)} \Rightarrow \sum_{t_i < t_k} \alpha e^{-\beta(t_k - t_i)} = \lambda(t_k) - \mu + \alpha \tag{1.4}$$

Despite being a Markovian process, it is still an inhomogeneous Poisson process because the rate is not constant. In addition, it is a self-exciting process, which means that the occurrence of an event increases the probability of the occurrence of another event. This is why it is used to model the spiking activity of neurons, where the occurrence of a spike increases the probability of the occurrence of another spike. This self-excitation will enable the appearance of bursts of activity that we will measure. The parameters chosen for the kernel function will be  $\alpha=\beta=1$  and we will vary the background rate  $\mu$  from values much smaller than 1 to values greater than 1. In Figures 1.3 and 1.4 we can see typical diagrams of Hawkes processes with these parameters.

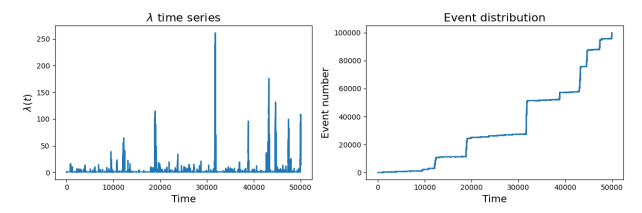


**Figure 1.3.** On the left, a temporal series of K = 150 events of a Hawkes process with  $\mu = 0.01$ , on the right, a raster plot of the same process.

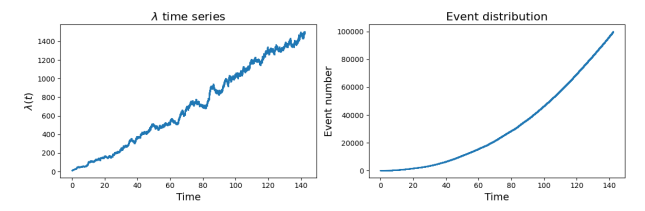


**Figure 1.4.** On the left, a temporal series of K = 150 events of a Hawkes process with  $\mu = 0.01$ , on the right, a raster plot of the same process.

As shown in Figure 1.3, when the background rate is smaller than 1, events are less likely to occur, but when they do, they tend to form avalanches of activity thanks to the self-excitation. On the other hand, when the background rate is greater than 1, events occur more frequently, forming avalanches of activity more frequently and longer, as shown in Figure 1.4. If we ignore the time of occurrence of the events and we focus only on the structure of  $\lambda$  and therefore of the events, we can see that the process with  $\mu = 0.01$  has a bursty structure, while the process with  $\mu = 10$  has a more regular structure. This phenomenon is exposed in Figures 1.5 and 1.6.



**Figure 1.5.** First, a temporal series of  $K = 10^5$  events of a Hawkes process with  $\mu = 0.01$ , on the right, the event distribution.



**Figure 1.6.** First, a temporal series of  $K=10^5$  events of a Hawkes process with  $\mu=10$ , on the right, the event distribution.

In most cases, the motivation of study of point processes is counting the events, but in our case we also are interested in the time of occurrence of the events which will let us define bursts or avalanches of activity that we will use to describe the dynamics of the system. Additionally,

Unless otherwise stated, n=1 HABLAR DE CRITICIDAD A PARTIR DE AQUÍ, EJEMPLOS DE CRITICAL BRANCHING PROCESSES, ETC.

### Chapter 2

## **Objectives**

Ordenar los objetivos una vez escrito el trabajo para que coincidan con como se presenta.

The main objectives of this Master's thesis are:

- To understand what Hawkes processes are, where we can find them, how to generate them computationally and relate them with neuroscience.
- To understand the importance of time binning and reproduce the results of the original paper [5] and compare them with the results obtained in this work.
- ¿Criticality?
- To study the behaviour of a self-exciting process with n=2 and compare it with the case n=1.
- To study the behaviour of an inhibitory and excitatory neuron coupled.

### Chapter 3

## Methodology

In the following sections, the methodology for data generation, managment and analysis will be presented. To address these issues, we will use Python [9, 10] due to its versatility and the wide range of libraries available. The two used will be NumPy [11] and Matplotlib [12] for the visualization.

#### 3.1 Time series factory

The first step is the generation of time series, there are two ways to do this: the slow one and the fast one. The first one is discretizing the time and calculating the rate at each time step according with Eq 1.2, then accept or reject the event if  $p < \lambda \cdot dt$  for a random number  $p \in \mathcal{U}[0,1]$ . This method works for small time series, but for large ones is not efficient because the summation of the kernel function has to be done at each time step. The pseudo-code for this method is presented in Algorithm 1.

#### Algorithm 1 Slow method to generate Hawkes processes.

```
 \begin{aligned} & \textbf{Require: } t_m ax, \ n_{intervals}, \ \lambda(t_0) = \mu, \ p \\ & dt \leftarrow \frac{t_{max}}{n_{intervals}} \\ & \textbf{for } i = 0 \ \text{to } n_{intervals} \ \textbf{do} \\ & \lambda(t_k) \leftarrow \mu + n \sum_{t_i < t_k} \phi(t_k - t_i) \\ & \textbf{if } \lambda(t_k) \cdot dt > p \ \textbf{then} \\ & t_{event} \leftarrow t_k \\ & \textbf{end if} \\ & \textbf{end for} \end{aligned}
```

The fast method takes advantage of Monte Carlo methods [13] to generate the time series. The idea of this procedure consists in computing the inter-event time instead of the time of the event. To get to the algorithm, we start from the following expression:

$$PDF(\text{inter-event time} = \Delta t) = \lambda(t + \Delta t)e^{-\int_t^{t+\Delta t} \lambda(t')dt'}$$
 (3.1)

To demonstrate this, we have to take a look at the Figure 3.1 and recall that  $\lambda$  is a probability per unit of time.

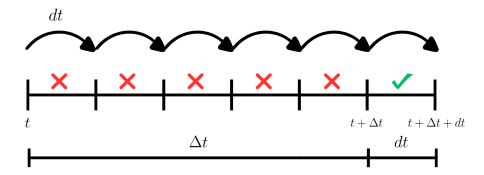


Figure 3.1. Diagram to calculate the cumulative probability of the inter-event time.

The probability per unit of time of having an event in the interval  $[t+\Delta t, t+\Delta t+dt]$  is the probability of no events in the interval  $[t, t+\Delta t]$  times the probability of happening in the interval  $[t+\Delta t, t+\Delta t+dt]$ . Putting words into mathematics, we have that the probability of not having an event in the interval  $[t, t+\Delta t]$  is:

$$P(\text{event} \in [t + \Delta t + dt]) = (1 - \lambda(0) \cdot dt) (1 - \lambda(dt) \cdot dt) (1 - \lambda(2dt) \cdot dt) \dots$$

$$= \prod_{k=0} \underbrace{(1 - \lambda(kdt) \cdot dt)}_{e^{\ln(1 - \lambda(kdt)dt)}} = e^{\sum_{k=0} \ln(1 - \lambda(kdt)dt)} = \dots \quad \text{Using } \ln(1 - \varepsilon) \approx -\varepsilon$$

$$= e^{-\sum_{k=0} \lambda(kdt)dt} \underbrace{=}_{dt \to 0} e^{-\int_{t}^{t+\Delta t} \lambda(t')dt'}.$$
(3.2)

Knowing that the probability of having an event in the interval  $[t + \Delta t, t + \Delta t + dt]$  is  $\lambda(t + \Delta t)dt$ , we have:

$$P(\text{event} \in [t + \Delta t, t + \Delta t + dt]) \mathcal{U} = \lambda(t + \Delta t) dt \cdot e^{-\int_{t}^{t + \Delta t} \lambda(t') dt'} PDF(\text{inter-event time} = \Delta t) \mathcal{U}. \tag{3.3}$$

Having that we can calculate the inter-event time following the next steps.

PDF (inter-event time 
$$= \Delta, t$$
)  $= \lambda(t + \Delta t) \underbrace{e^{\int_t^{t+\Delta t} \lambda(t'dt')}}_{\text{No events during } (t, t+\Delta t)}$ 

In order to generate  $\Delta t$ , we will use the inverse transform method [14], therefore we have to calculate the cumulative probability of the inter-event time:

$$\operatorname{accum}(\Delta t) = \int_{0}^{\Delta t} \operatorname{PDF}(\Delta t') d\Delta t' = u \in \mathcal{U}[0, 1]$$

$$\int_{0}^{\Delta t} \underbrace{\lambda(t + \Delta t') e^{\int_{t}^{t + \Delta t'} \lambda(t') dt'}}_{-\frac{d}{d\Delta t'} \left[e^{-\int_{t}^{t + \Delta t} \lambda t' dt'}\right]} d\Delta t' = u \quad \text{Using Barrow rule}$$

$$-e^{-\int_{t}^{t + \Delta t} \lambda(t') dt'} \Big|_{0}^{\Delta t} = 1 - e^{-\int_{t}^{t + \Delta t} \lambda(t') dt'} = u \quad \text{Taking logarithms}$$

$$\int_{t}^{t + \Delta t} \lambda(t') dt' = -\ln(1 - u) = \ln(\bar{u})$$

To compute the inter-event time, we have to generate  $\bar{u} \sim$  and solve the equation. Having in mind this relation and using Eq 1.4 we have: EN EL LA ECUACIÓN REFERENCIADA Y LA SEGUNDA EXPONENCIAL SE PUEDE PONER  $\lambda(t_k^-)$  en lugar de  $\lambda(t_k)$ ?.

$$u = 1 - e^{-\mu(t - t_k)} e^{-(\lambda(t_k) + \alpha - \mu) \cdot \int_{t_k}^{t} e^{-\beta(t') - t_k} dt'}$$

$$u = 1 - \underbrace{e^{-\mu(t - t_k)}}_{P(t_{k+1}^{(1)} > t)} \underbrace{e^{-\left[(\lambda(t_k) + \alpha - \mu)\beta^{-1}\left(1 - e^{-\beta(t - t_k)}\right)\right]}}_{P(t_{k+1}^{(2)} > t)}$$
(3.5)

Then we apply the composition method [7]. If we take  $t_{k+1} = \min(t_{k+1}^{(1)}, t_{k+1}^{(2)})$ ; then  $t_{k+1} \sim P(t_{k+1} > t)$ , hence:

$$\operatorname{Prob}(t_{k+1} = \min\left(t_{k+1}^{(1)}, t_{k+1}^{(2)}\right) \le t) = 1 - \operatorname{Prob}\left(\min\left(t_{k+1}^{(1)}, t_{k+1}^{(2)}\right) > t\right)$$

$$= 1 - \operatorname{Prob}\left(t_{k+1}^{(1)} > t\right) \cdot \operatorname{Prob}\left(t_{k+1}^{(2)} > t\right)$$
(3.6)

where we have used that the probability that the smaller is greater than t is that each separately is greater than t because both have to be greater than t. As we can see the expressions in Eqs 3.5 and 3.6 are the same, so we can use the composition method to generate the inter-event time. Then, the algorithm to generate the inter-event time is:

1. Generate  $t_{k+1}^{(1)} \sim P\left(t_{k+1}^{(1)} > t\right) = e^{-\mu(t-t_k)}$  using

$$P\left(t_{k+1}^{(1)} \le t\right) = 1 - \underbrace{e^{-\mu(t-t_k)}}_{\bar{u_1} \in \mathcal{U}[0,1] \Rightarrow = u_1 - \bar{u_1}} = u_1 \in \mathcal{U}[0,1]$$

This is done by generating  $u_1 \in \mathcal{U}[0,1]$  and solving the equation.

$$u_{1} = 1 - e^{-\mu \left(t_{k+1}^{(1)} - t_{k}\right)}$$

$$\ln(u_{1}) = -\mu \left(t_{k+1}^{(1)} - t_{k}\right) \Rightarrow t_{k+1}^{(1)} = t_{k} - \frac{\ln(u_{1})}{\mu}$$
(3.7)

2. Generate  $t_{k+1}^{(2)} \sim P\left(t_{k+1}^{(2)} > t\right) = e^{-\left((\lambda(t_k) + \alpha - \mu)\beta^{-1}\left(1 - e^{-\beta\left(t_{k+1}^{(2)} - t_k\right)}\right)\right)}$  in a similar way as before:

$$u_{2} = 1 - e^{-\left((\lambda(t_{k}) + \alpha - \mu)\beta^{-1} \left(1 - e^{-\beta \left(t_{k+1}^{(2)-t_{k}}\right)}\right)\right)}$$

$$-\ln(u_{2}) = \left((\lambda(t_{k}) + \alpha - \mu)\beta^{-1} \left(1 - e^{-\beta \left(t_{k+1}^{(2)-t_{k}}\right)}\right)\right)$$

$$1 + \frac{\beta \ln u_{2}}{\lambda(t_{k}) + \alpha - \mu} = e^{-\beta \left(t_{k+1}^{(2)-t_{k}}\right)}$$

$$t_{k+1}^{(2)} = t_{k} - \beta^{-1} \ln \left(1 + \frac{\beta \ln u_{2}}{\lambda(t_{k}) + \alpha - \mu}\right)$$
This number must be positive
$$(3.8)$$

3. Choose  $t_{k+1} = \min\left(t_{k+1}^{(1)}, t_{k+1}^{(2)}\right)$ 

4. Calculate the rate at  $t_{k+1}$  using Eq 1.3 and go back to step 1.

With this method, we can generate time series efficiently. The pseudo-code for this method is presented in Algorithm 2.

#### **Algorithm 2** Algorithm to generate K Hawkes events

```
Require: \alpha, \beta, \lambda(t_0) = \mu, K

for k = 0 to K do

u_1, u_2 \leftarrow \mathcal{U}[0, 1]

t_{k+1}^{(1)} \leftarrow \frac{\ln(u_1)}{\mu}

t_{k+1}^{(2)} \leftarrow \beta^{-1} \ln \left( 1 + \frac{\beta \ln u_2}{\lambda(t_k) + \alpha - \mu} \right)

t_{k+1} \leftarrow \min \left( t_{k+1}^{(1)}, t_{k+1}^{(2)} \right)

\lambda(t_{k+1}) \leftarrow \mu + e^{-\beta(t_{k+1} - t_k)} \left( \lambda(t_k) - \mu + n \right)

end for
```

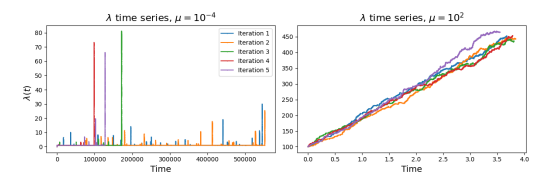
#### 3.2 When physics and cooking merge

Once we have a method to generate time series, we can start analyzing them. We should stablish a control parameter to distinguish between different regimes of the process. In this case, we will define resolution parameter  $\Delta > 0$  as a time interval which will help us to identify clusters of activities. Our time series will have K events that occur in times  $\{t_1, \ldots, t_K\}$ . Considering this, a cluster os events starts at time  $t_i$  and ends at time  $t_j$  if  $t_j - t_i \leq \Delta$ . The number of events in the cluster (cluster size) is the number of events in the interval  $[t_i, t_j]$  and it's duration is  $t_j - t_i$ . The extreme cases are when  $\Delta$  is smaller than the minimum inter-event time, in this case, each event is a cluster of size 1 and duration 0. On the other hand when  $\Delta$  is greater than the largest inter-event time, all the events are in the same cluster of size K and duration  $t_K - t_1$ . Between these two extremes, we will have different regimes of the process. To identify these regimes, we need a phase diagram, in our case, it will be a percolation diagram where we will plot the percolation strength  $P_{\infty}$  as a function of the resolution parameter  $\Delta$ . The percolation strength is defined as the fraction of events that are in the largest cluster over the total number of events. Three different set of parameters will be used to generate the time series in order to compare them. The parameters  $\alpha$  and  $\beta$  will be fixed to 1 in all cases, the other parameters are shown in Table 3.1.

Table 3.1. Configuration of the parameters for the simulations

Configuration	$\mu$	n
First	1	0
Second	$10^{-4}$	1
Third	$10^{2}$	1

The percolation diagram will be generated by generating 1000 time series for each configuration and calculating the percolation strength for each one because in general they are not stationary processes as we can observer in Figure 3.2.



**Figure 3.2.** Five a temporal series of  $K = 10^5$  events of Hawkes processes with  $\mu = 10^{-4}$  on the left side and  $\mu = 10^2$  on the right one.

Beginning with the first configuration, we have an homogeneous Poisson process which we know that its interevent time will be distributed randomly with a probability of having an inter-event time  $x_i$  given by  $P(x_i) = \mu e^{-\mu x_i}$ . Consequently, two consecutive events will be a part of a cluster fixing the resolution parameter to  $\Delta$  with a probability of

$$P(x_i \le \Delta) = 1 - e^{-\mu\Delta} \qquad \forall i. \tag{3.9}$$

This represents the probability in a homogeneous 1D percolation model [15], where we can identify a non percolant phase and a percolant phase separated by the critical point  $\Delta^*$ . We can calculate this parameter if we know the maximum inter-event time of the time series. Let us assume that our time series has K events, therefore, it will percolate if the condition we have just stablished is satisfied. We can calculate this threshold as the average of the maximum inter-event time in K samples from the inter-event time distribution solving the following equation:

$$K \int_{\Delta^*}^{\infty} P(x)dx = 1$$

$$K \int_{\Delta^*}^{\infty} \mu e^{-\mu x} dx = 1$$

$$-K \left[ e^{-\mu x} \right]_{\Delta^*}^{\infty} = K \left[ e^{-\mu \Delta^*} - e^{-\mu \omega} \right]_{0}^{0} = 1$$

$$K e^{-\mu \Delta^*} = 1$$

$$\Delta^*(K) = -\frac{\ln(K)}{\mu}$$

$$(3.10)$$

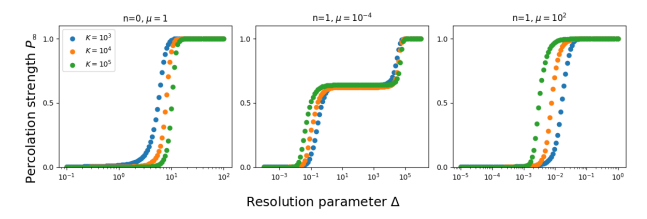
### Chapter 4

### Results

This section provides the main results of the investigation. First, the results reproduced from the original paper [5] are presented. Then, the results of the analysis with n=2 are shown. Finally, we have studied the behaviour of an inhibitory and excitatory neuron coupled.

### 4.1 Results from the original paper

The first result is the percolation phase diagram is shown in Figure 4.1. It displays the percolation strength  $P_{\infty}$  versus the resolution parameter  $\Delta$ .



**Figure 4.1.** Percolation phase diagrams for different event number K taking average values of R = 1000 realizations.

The first plot configuration is a Markovian (n=0) Poisson process with rate  $\mu$ . This is the simplest case, where the inter-event time  $x=t_i-t_{i-1}$  follows an exponential distribution  $P(x_i)=\mu e^{\mu x_i}$ . The other two plots are Hawkes processes for  $\mu\ll 1$  and  $\mu\gg 1$  that are also Markovian as we have chosen an exponential kernel (REFERENCIAR AQUÍ A LA PARTE EN LA QUE SE EXPLICA EN METODOLOGÍA.) . In one hand, a double transition is observed when  $\mu=10^{-4}$ , in the other hand, a single transition occurs when  $\mu=10^2$ .

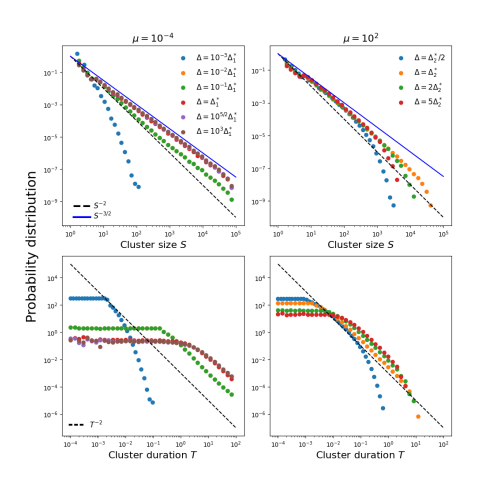
Once we have the phase diagram, we can study avalanche statistics. Given a resolution parameter  $\Delta$ , we can spot clusters or avalanches of activity. A cluster starts when a neuron fires and ends if

the neuron does not fire for a time greater than  $\Delta$ . We define the size of a cluster as the number of spikes it contains and the duration as the time between the first and last spike. We have studied the avalanches for  $K=10^5$  events and R=1000 realizations to obtain the average values since the process is highly not stationary. We will study the size and duration of the avalanches for the three different regions of the phase diagram for  $\mu=10^{-4}$  and the two regions of the phase diagram for  $\mu=10^2$ . These regions are separated by two thresholds, a pseudocritical threshold  $\Delta_1^*$  and the threshold of the second transition at  $\Delta_2^*$ . We can compute these with the following formulas [5]:

$$\Delta_1^* \simeq \frac{\log(K)}{\langle \lambda \rangle} = \frac{\log(K)}{\mu + \sqrt{2\mu K}} \tag{4.1}$$

$$\Delta_2^* = \frac{\log(K)}{\mu} \tag{4.2}$$

Once we have the thresholds, we can study the avalanches for the different regions of the diagram. The results are shown in Figure 4.2.



**Figure 4.2.** Avalanche statistics for a self-exciting Hawkes process with n = 1 for  $K = 10^5$  events averaged over R = 1000 realizations.

For  $\mu = 10^{-4}$ , the results show a power-law distribution for the size and duration of the avalanches. In the case of duration, the exponent is  $\tau = 2$  and for the size, we can notice a transition of the exponent from  $\alpha = 2$  to  $\alpha = 3/2$  as we increase the resolution parameter  $\Delta$ . The first exponent corresponds to the universality class of 1D percolation, whereas the second is compatible with the universality class of mean-field branching process. However, if  $\Delta \ll \Delta_1^*$ , the behaviour is subcritical for the size and duration of the avalanches.

For  $\mu=10^2$ , the result shows another power-law distribution for both size and duration of the avalanches unless  $\Delta \ll \Delta_2^*$ , where the behaviour is subcritical. In this case, the exponents are  $\alpha=\tau=2$  corresponding to the universality class of 1D percolation.

HABLAR AQUÍ DE LA INFLUENCIA DEL MENOR NÚMERO DE EVENTOS, SE SIGUE PRODUCIENDO LA TRANSICIÓN, PERO PARA OTROS VALORES DE DELTA

#### 4.2 Results for n=2

In the article, the authors have studied a process which is critical itself because the parameter n is fixed to n = 1. We have studied the case n = 2 to see if the process is still critical. In the Figure 4.3 two time series for n = 1 and n = 2 are shown.

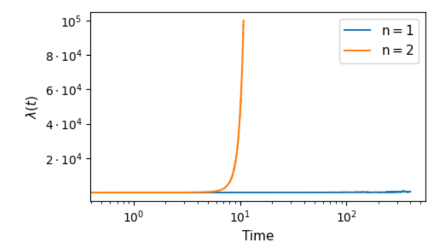
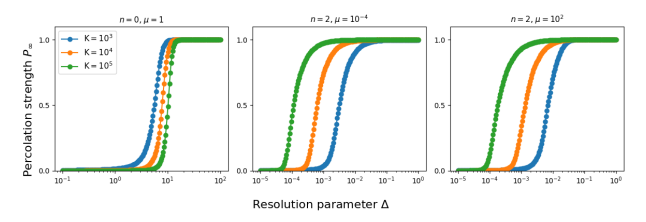


Figure 4.3. Time series for n = 1 and n = 2.

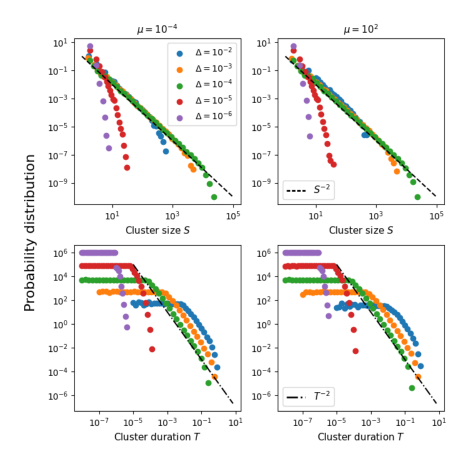
Similarly to the previous section, first we obtain the phase diagram in order to observe the transitions. In this case, Eqs 4.1-4.2 are not valid. Therefore, we will obtain this parameter graphically from the phase diagrams shown in Figure 4.4.



**Figure 4.4.** Percolation phase diagrams for a Hawkes process with n=2.

As we can see, now we have a single transition for  $\mu=10^{-4}$  and  $\mu=10^2$  corresponding to 1D

percolation, consequently, the exponents for the size and duration should be  $\alpha = \tau = 2$ . In a similar way, we have studied the avalanches for  $K = 10^5$  events and R = 1000 realizations to obtain the average values. The statistics of the avalanches are shown in Figure 4.5.



**Figure 4.5.** Avalanche statistics for a self-exciting Hawkes process with n=2 for  $K=10^5$  events averaged over R=1000 realizations.

4.3 Inhibitory and excitatory neurons coupled

## Chapter 5

## Conclusions

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### Anexo

# REVISAR LOS CÓDIGOS PARA QUE ESTÉN ACTUALIZADOS

Script 5.1. Script with the main functions.

```
1 import numpy as np
  import matplotlib.pyplot as plt
  def algorithm (rate, mu, n):
       Algorithm that computes interevent times and Hawkes intensity for a self-exciting
6
      process
      \#Output: rate x_k, x_k
8
9
      # 1st step
      u1 = np.random.uniform()
11
       if mu == 0:
12
           F1 = np.inf
       else:
          F1 = -np.\log(u1) / mu
16
      # 2nd step
17
      u2 = np.random.uniform()
18
       if (rate - mu) == 0:
19
           G2 = 0
20
21
           G2 = 1 + np.log(u2) / (rate - mu)
22
23
      # 3rd step
       if G2 \ll 0:
26
          F2 = np.inf
       else:
28
          F2 = -np \cdot log(G2)
29
30
      # 4th step
31
      xk = \min(F1, F2)
32
33
      # 5th step
      rate_tk = (rate - mu) * np.exp(-xk) + n + mu
       return rate_tk, xk
37
  def generate_series(K, n, mu):
38
39
       Generates temporal series for K Hawkes processes
40
41
```

```
42
       ##Inputs:
       K: Number of events
43
       n: Strength of the Hawkes process
44
       mu: Background intensity
45
46
       ##Output:
47
       times: time series the events
48
       rate: time series for the intensity
49
50
       times\_between\_events = [0]
51
       rate = [mu]
        for _ in range(K):
53
            rate_tk, xk = algorithm(rate[-1], mu, n)
54
            rate.append(rate_tk)
56
            times_between_events.append(xk)
       times = np.cumsum(times_between_events)
57
58
       return times, rate
59
   def identify_clusters(times, delta):
60
61
        Identifies clusters in a temporal series given a resolution parameter delta
62
63
       ## Inputs:
64
       times: temporal series
65
       delta: resolution parameter
66
67
68
       ## Output:
       clusters: list of clusters
69
70
       clusters = []
71
       current_cluster = []
72
        for i in range (len(times) - 1):
73
            if times[i + 1] - times[i] \le delta:
74
                if not current_cluster:
75
                     current_cluster.append(times[i])
76
                current\_cluster.append(times[i + 1])
78
            else:
79
                if current_cluster:
                     clusters.append(current_cluster)
80
                     current_cluster = []
81
82
       return clusters
83
   def generate_series_perc(K, n, mu):
84
85
        Generates temporal series for K Hawkes processes
86
87
       ##Inputs:
       K: Number of events
89
       n: Strength of the Hawkes process
90
       mu: Background intensity
91
92
       ##Output:
93
       times_between_events: time series the interevent times
94
       times: time series the events
95
       rate: time series for the intensity
96
97
       times\_between\_events = [0]
       rate = [mu]
99
        for \underline{\quad} in range(K):
100
            rate_tk, xk = algorithm(rate[-1], mu, n)
101
            rate.append(rate_tk)
            times_between_events.append(xk)
103
```

```
times = np.cumsum(times_between_events)
104
105
        return times_between_events, times, rate
106
   def calculate_percolation_strength(times_between_events, deltas):
107
108
        Calculate the percolation strength for a given set of deltas (resolution parameters)
109
110
       ## Inputs:
       {\tt times\_between\_events} \colon \; {\tt time} \; \; {\tt series} \; \; {\tt of} \; \; {\tt interevent} \; \; {\tt times}
        deltas: list of resolution parameters
113
114
       ## Output:
115
        percolation_strengths: list of percolation strengths
116
117
        percolation\_strengths = []
119
120
        for delta in deltas:
121
            cluster_sizes = []
            current_cluster_size = 1 # The first event is always a cluster
123
124
            for time in times_between_events:
                if time < delta:
126
                     current_cluster_size += 1
127
                else:
128
                     if current_cluster_size > 1: # Only consider clusters with more than one
129
       event
130
                         cluster_sizes.append(current_cluster_size)
                     current_cluster_size = 1 # The next event is always a cluster
132
            if current_cluster_size > 1: # Consider the last cluster if it ends at the last
       event
                cluster_sizes.append(current_cluster_size)
134
135
            if len(cluster_sizes) != 0: # Check if cluster_sizes is not empty to avoid
136
       errors
                max_cluster_size = max(cluster_sizes)
            else:
                max\_cluster\_size = 0
139
140
            percolation_strengths.append(max_cluster_size / len(times_between_events))
141
        return percolation_strengths
143
144
145
   """def calculate_percolation_strength(times_between_events, deltas):
146
        percolation\_strengths = []
147
148
        for delta in deltas:
149
            cluster_sizes = []
            # Initialize the size of the current cluster
            current_cluster_size = 1 # The first event is always a cluster
152
            for i in range(len(times_between_events)):
154
                if times_between_events[i] <= delta:
155
156
                     current_cluster_size += 1
                else:
                     if current_cluster_size > 1: # Only consider clusters with more than one
       event
                         cluster_sizes.append(current_cluster_size)
159
                    # Reset the size of the current cluster
160
                     current_cluster_size = 1 # The next event is always a cluster
161
```

```
162
           # Add the size of the last cluster
163
            if current_cluster_size > 1: # Only consider clusters with more than one event
164
                cluster_sizes.append(current_cluster_size)
            max_cluster_size = max(cluster_sizes)
167
168
            percolation_strengths.append(max_cluster_size / len(times_between_events))
169
       return percolation_strengths"""
170
171
   def model(n_max, mu_E, mu_I, tau, n_EE, n_IE, n_EI, n_II, dt):
172
173
       Solve the equations of the mena field model for a given number of iterations n_max
174
       Inputs:
       n_max: number of iterations
       mu_E: Poisson rate of excitatory neurons
       mu_I: Poisson rate of inhibitory neurons
179
       tau: characteristic time of the system
180
       n_EE: influence of excitatory neurons on excitatory neurons
181
       n_IE: influence of excitatory neurons on inhibitory neurons
182
       n_EI: influence of inhibitory neurons on excitatory neurons
183
       n_II: influence of inhibitory neurons on inhibitory neurons
184
       dt: time step
185
186
       Outputs:
187
       time: time series
188
189
       t_events_E: times of events of excitatory neurons
       t_events_I: times of events of inhibitory neurons
190
191
       rates_E: rates of excitatory neurons
       rates_I: rates of inhibitory neurons
       n_E = n_I = n = 0
194
       t_{events} = [0]
195
       t_events_I = [0]
196
       rates_E = [mu_E]
       rates_I = [mu_I]
198
       time = [0]
199
        while n \le n_max:
200
           # Excitation neurons
201
            l\_Enew = rates\_E[-1] + dt * (mu\_E- rates\_E[-1])/tau
202
            if np.random.uniform() < rates_E[-1]*dt:
203
                l_Enew += n_EE
204
                t_{events}: append (time [-1]+dt*np.random.uniform())
205
                n_E += 1
206
            if np.random.uniform() < rates_I[-1]*dt:
                l_Enew -= n_IE
                t_{\text{events}} E. append (time[-1] + dt*np.random.uniform())
209
                n_E += 1
210
211
           # Inhibition neurons
212
            l\_Inew \, = \, rates\_I \, [-1] \, + \, dt \, * \, (mu\_I \!\! - \, rates\_I \, [-1]) / tau
213
            if np.random.uniform() < rates_E[-1]*dt:
214
215
                l\_Inew += n\_EI
216
                t_{events}I.append(time[-1]+dt*np.random.uniform())
217
                n_I += 1
            if np.random.uniform() < rates_I[-1]*dt:
                l\_Inew = n\_II
219
                t_{events}I.append(time[-1]+dt*np.random.uniform())
220
221
                n_I += 1
            rates_E.append(l_Enew)
222
            rates_I.append(l_Inew)
223
```

```
time.append(time[-1]+dt)
224
225
            n = n\_E + n\_I
226
        return time, t_events_E, t_events_I, rates_E, rates_I
227
228
   def identify_clusters_model(times, delta):
229
230
       Identifies clusters in a temporal series given a resolution parameter delta
231
       Computes the size and duration of clusters
232
233
       ## Inputs:
234
       times: temporal series
235
       delta: resolution parameter
236
237
       ## Output:
       clusters: list of clusters
239
       clusters_sizes: list of sizes of clusters
240
       clusters_times: list of durations of clusters
241
242
       clusters = []
243
       current_cluster = []
244
        for i in range (len (times) -1):
245
            if times[i + 1] - times[i] \ll delta:
246
                if not current_cluster:
247
                    current_cluster.append(times[i])
248
                current\_cluster.append(times[i + 1])
249
            else:
250
251
                if current_cluster:
                    clusters.append(current_cluster)
252
253
                    current_cluster = []
254
       clusters_sizes = [len(cluster) for cluster in clusters]
255
       clusters\_times = [cluster[-1] - cluster[0]] for cluster[in] clusters]
256
       return clusters, clusters_sizes, clusters_times
257
258
   def bivariate_algorithm(rate1, rate2, muE, muI, nEE, nII, nEI, nIE):
259
260
       Algorithm that computes interevent times and Hawkes intensity for a bivariate Hawkes
261
       process
262
       #Inputs:
263
       rate1: Previous excitation rate
264
       rate2: Previous inhibition rate
265
       nEE: "Strength" of the autoexcitation process
266
       nII: "Strength" of the autoinhibition process
267
       nEI: "Strength" of the excitation to the inhibition
       nIE: "Strength" of the inhibition to the excitation
       muE: Background intensity of the excitation
270
       muI: Background intensity of the inhibition
271
272
273
       #Output: ratex_k, x_k, reaction (0 for excitatory events and 1 for inhibitory events)
274
275
       _{-}, xk1 = algorithm (rate1, muE, nEE)
276
       _{-}, xk2 = algorithm(rate2, muI, nII)
277
278
       xks = [xk1, xk2]
280
        if xk1 < 0:
281
            print (xk1)
282
            xk1 = 0
283
        if xk2 < 0:
284
```

```
print (xk2)
285
           xk2 = 0
286
287
       reaction = np.argmin(xks)
       if reaction = 0:
290
            rate1_tk = (rate1 - muE) * np.exp(-xk1) + nEE + muE
291
           rate2\_tk = (rate2 - muI) * np.exp(-xk1) + nEI + muI
292
       else:
293
            rate1_tk = (rate1 - muE) * np.exp(-xk2) + nIE + muE
294
           rate2\_tk = (rate2 - muI) * np.exp(-xk2) + nII + muI
295
296
       if rate1_tk <= muE:</pre>
297
           rate1\_tk = muE
298
       if rate2_tk <= muI:</pre>
           rate2\_tk = muI
301
       xk = xks[reaction]
302
303
       return rate1_tk, rate2_tk, xk, reaction
304
305
   def generate_series_bivariate(K, nEE, nII, nEI, nIE, muE, muI):
306
307
       Generates temporal series for K bivariate Hawkes processes
308
       ##Inputs:
310
       K: Number of events
311
312
       nEE: "Strength" of the autoexcitation process
       nII: "Strength" of the autoinhibition process
313
       nEI: "Strength" of the excitation to the inhibition
314
       nIE: "Strength" of the inhibition to the excitation
315
       muE: Background intensity of the excitation
316
       muI: Background intensity of the inhibition
317
318
       ##Output:
319
       times_between_events: time series the interevent times
       times: time series the events
       rate1: time series for the intensity of process 1 (Excitation)
       rate2: time series for the intensity of process 2 (Inhibition)
323
       reactions: list the event type (0 for excitation. 1 for inhibition)
324
325
       times\_between\_events = [0]
326
       rate1 = [muE]
327
       rate2 = [muI]
328
       reactions= []
329
       for _ in range(K):
           rate1_tk, rate2_tk, xk, reaction = bivariate_algorithm(rate1[-1], rate2[-1], muE,
        muI, nEE, nII, nEI, nIE)
           rate1.append(rate1_tk)
332
           rate2.append(rate2_tk)
333
            reactions.append(reaction)
334
            times_between_events.append(xk)
335
       times = np.cumsum(times_between_events)
336
337
       return times_between_events, times, rate1, rate2, reactions
```