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Spike train statistics and Gibbs distributions

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Abstract

This paper is based on a lecture given in the LACONEU summer school, Valparaiso, January 2012. We introduce Gibbs distribution in a general setting, including non stationary dynamics, and present then three examples of such Gibbs distributions, in the context of neural networks spike train statistics: (i) Maximum entropy model with spatio-temporal constraints; (ii) Generalized Linear Models; (iii) Conductance based Integrate and Fire model with chemical synapses and gap junctions.

Keywords Neural networks dynamics; spike train statistics; Gibbs distributions.

1 Introduction

Neurons communicate among them by generating action potentials or “spikes” which are pulses of electrical activity. When submitted to external stimuli, sensory neurons produce sequences of spikes or “spike trains” constituting a collective response and a dynamical way to encode information about those stimuli. However, neural responses are typically not exactly reproducible, even for repeated presentation of a fixed stimulus. Therefore, characterizing the relationship between sensory stimuli and neural spike responses can be framed as a problem of determining the most adequate probability distribution relating a stimulus to its neural response. There exist several attempts to infer this probability from data and / or general principles, based on Poisson or more general point processes [1, 16, 51], Bayesian approaches [29, 23], maximum entropy [17, 55] (for a review see [43]). In this paper we present several situations where the notion of Gibbs distributions is appropriate to address this problem.

The concept of Gibbs distribution comes from statistical physics. We use it here in a more general sense than the one usually taught in standard physics courses, although it is part of mathematical statistical physics [22]. We argue here that Gibbs distributions might be canonical models for spike train statistics analysis. This statement is based on three prominent examples.

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1. The so-called Maximum Entropy Principle allows one to propose spike train statistics models considering restrictions based on empirical observations. Although this approach has been initially devoted to show the role of weak instantaneous pairwise correlations in the retina [47], it has been recently applied to investigate the role of more complex events such as instantaneous triplets [19] or spatio-temporal events [55]. Probability distributions arising from the Maximum Entropy Principle are Gibbs distributions.

2. Other approaches such as the Linear-Non Linear (LN) or Generalized Linear Models (GLM) propose an ad hoc form for the conditional probability that a neuron fires given the past network activity and given the stimulus. Those models have been proven quite efficient for retina spike trains analysis [41]. They are not limited by the constraint of stationarity, but they are based on a questionable assumption of conditional independence between neurons. As we show, the probability distributions coming out from those models are also Gibbs distributions.

3. Recent investigations on neural networks models (conductance based integrate-and-fire (IF) with chemical and electric synapses) show that statistics of spike trains generated by these models are Gibbs distributions reducing to 1 when dynamics is stationary, and reducing to 2 in specific cases [7, 8, 14]. In the general case, the spike trains produced by these models have Gibbs distributions which neither match 1 nor 2.

The paper is organized as follows. After some definitions regarding spike train statistics and a presentation of Gibbs distributions we develop these three examples, with a short discussion of their advantages and drawbacks in spike trains analysis. Then, we discuss some relations between these models, mainly based on the Hammersley-Clifford theorem [24, 2, 33, 39]. This paper is a summary of several papers written by the authors and other collaborators [7, 8, 34, 9, 14]. As such it does not contain original material (except the presentation).

2 Definitions

2.1 Spike trains

We consider a network of $N$ neurons. We assume that there is a minimal time scale $\delta > 0$ corresponding to the minimal resolution of the spike time, constrained by biophysics and by measurements methods (typically $\delta \sim 1 \text{ ms}$) [11 10]. Without loss of generality (change of time units) we set $\delta = 1$, so that spikes are recorded at integer times. One then associates to each neuron $k$ and each integer time $n$ a variable $\omega_k(n) = 1$ if neuron $k$ fires at time $n$ and $\omega_k(n) = 0$ otherwise. A spiking pattern is a vector $\omega(n) \overset{\text{def}}{=} [\omega_k(n)]_{k=1}^{N}$ which tells us which neurons are firing at time $n$. We note $\mathcal{A} = \{0, 1\}^N$ the set of spiking patterns. A spike block is a finite ordered list of spiking patterns, written:

$$\omega_{n_1}^{n_2} = \{\omega(n)\}_{n_1 \leq n \leq n_2}.$$
where spike times have been prescribed between the times \( n_1 \) to \( n_2 \) (i.e., \( n_2 - n_1 + 1 \) time steps). The range of a block is \( n_2 - n_1 + 1 \) time steps from \( n_1 \) to \( n_2 \). The set of such blocks is \( \mathcal{A}_{n_2-n_1+1} \). Thus, there are \( 2^Nn \) possible blocks with \( N \) neurons and range \( n \). We call a raster plot a bi-infinite sequence \( \omega \) of spiking patterns. Obviously experimental rasters are finite, but the consideration of infinite sequences is more convenient mathematically. The set of such blocks is \( \mathcal{A}^Z \).

2.2 Transition probabilities

The probability that a neuron emits a spike at some time \( n \) depends on the history of the neural network. However, it is impossible to know explicitly its form in the general case since it depends on the past evolution of all variables determining the neural network state. A possible simplification is to consider that this probability depends only on the spikes emitted in the past by the network. In this way, we are seeking a family of transition probabilities of the form \( \mathbb{P}_n[\omega(n) \mid \omega_{n-D}^{n-1}] \), the probability that the firing pattern \( \omega(n) \) occurs at time \( n \) given a past spiking sequence \( \omega_{n-D}^{n-1} \). Here, \( D \) is the memory depth of the probability, i.e., how far in the past does the transition probability depend on the past spike sequence. We use the convention that \( \mathbb{P}_n[\omega(n) \mid \omega_{n-D}^{n-1}] = \mathbb{P}_n[\omega(n)] \) if \( D = 0 \) (memory-less case).

The index \( n \) of \( \mathbb{P}_n[ \cdot \mid \cdot ] \) indicates that transition probabilities depend explicitly on the time \( n \). We say that those transition probabilities are time-translation invariant or stationary if for all \( n \), \( \mathbb{P}_n[\omega(n) \mid \omega_{n-D}^{n-1}] = \mathbb{P}_D[\omega(D) \mid \omega_{D-1}^{0}] \) whenever \( \omega_{n-D}^{n-1} = \omega_{D-1}^{0} \) (i.e., the probability does not depend explicitly on time). In this case we drop the index \( n \).

Transition probabilities depend on the neural network characteristics such as neurons conductances, synaptic responses or external currents. They give information about the dynamics that takes place in the observed neural network. Especially, they have a causal structure where the probability of an event depends on the past. This reflects underlying biophysical mechanisms in the neural network, which are also causal.

2.3 Gibbs distribution

We define here Gibbs distributions (or Gibbs measures) in a more general setting that the one usually taught in statistical physics courses, where Gibbs distributions are considered in the realm of stationary process and maximum entropy principle. Here, we do not assume stationarity and the definition encompasses the maximum entropy distributions. The Gibbs distributions considered here are called chains with complete connections in the realm of stochastic processes [38, 39] and g-measures in ergodic theory [27]. They are also studied in mathematical statistical physics [22].

2.3.1 Continuity with respect to a raster

For \( n \in \mathbb{Z} \), we note \( \mathcal{A}_{n-1}^{\infty} \) the set of sequences \( \omega_{n-1}^{\infty} \). Assume that we are given a set of transitions probabilities, like in the previous section, possibly depending
on an infinite past\footnote{In this case, one has to assume that (i) for every $\omega(n) \in A$, $P_n[\omega(n) | \omega_{-\infty}^{-1}]$ is measurable with respect to $F_{\leq n-1}$, the sigma-algebra on $A_{-\infty}^{n-1}$; (ii) for every $\omega_{-\infty}^{-1} \in A_{-\infty}^{-1}$, $\sum_{\omega(n) \in A} P_n[\omega(n) | \omega_{-\infty}^{-1}] = 1$.} i.e. of the form $P_n[\omega(n) | \omega_{-\infty}^{n-1}]$. We give in section 3.3 an example of neural network model where such transition probabilities with an infinite memory do occur.

Even if transition probabilities involve an infinite memory $\omega_{-\infty}^{n-1}$, it is reasonable to consider situations where the effects of past spikes decreases exponentially with their distance in the past. This corresponds to the mathematical notion of \textit{continuity with respect to a raster}. We note, for $n \in \mathbb{Z}$, $m \geq 0$, and $r$ integer:

$$\omega^{mn} = \omega' \text{ if } \omega(r) = \omega'(r), \forall r \in \{n-m, \ldots, n\}.$$Consider a function $f$ depending both on discrete time $n$ and on the raster part of $\omega$ anterior to $n$. We write $f(n, \omega)$ instead of $f(n, \omega_{-\infty}^{n-1})$. The function $f$ is \textit{continuous with respect to the raster} $\omega$ if its $m$-variation:

$$\text{var}_m[f(n, \omega)] := \sup \left\{ \left| f(n, \omega) - f(n, \omega') \right| : \omega^{mn} = \omega' \right\}$$

 tends to 0 as $m \to +\infty$. This precisely means that the effect, on the value of $f$ at time $n$, as this change is more distant in the past.

\subsection*{2.3.2 Gibbs distribution}

\begin{definition}
A Gibbs distribution is a probability measure $\mu : \Omega \to [0, 1]$ such that:

(i) for all $n \in \mathbb{Z}$ and all $\mathcal{F}_{\leq n}$-measurable functions $f$:

$$\int f(\omega_{-\infty}^{n-1}) \mu(d\omega) = \int \sum_{\omega(n) \in A} f(\omega_{-\infty}^{n-1}) P_n[\omega(n) | \omega_{-\infty}^{n-1}] \mu(d\omega).$$

(ii) $\forall n \in \mathbb{Z}, \forall \omega_{-\infty}^{n-1} \in A_{-\infty}^{n-1}, P_n[\omega(n) | \omega_{-\infty}^{n-1}] > 0$.

(iii) For each $n \in \mathbb{Z}$, $P_n[\omega(n) | \omega_{-\infty}^{n-1}]$ is continuous with respect to $\omega$.

The condition (i) is a natural extension of the condition defining the invariant probability of an homogeneous Markov chain (see eq. (2) next section). In its most general sense (i) does not require stationarity and affords the consideration of an infinite memory. It defines so-called \textit{compatibility conditions}. They state that the average of a function $f(n, \omega)$ with respect to $\mu$, at time $n$ (left hand side), is equal to the average computed from transition probabilities (right hand side). This equality must hold for any time $n$.

There exist several theorems guaranteeing the existence and uniqueness of a Gibbs distribution \cite{22, 13}: this holds if the variation of transition probability decays sufficiently fast with time (typically exponentially) as $m \to +\infty$. \footnote{In this case, one has to assume that (i) for every $\omega(n) \in A$, $P_n[\omega(n) | \omega_{-\infty}^{n-1}]$ is measurable with respect to $F_{\leq n-1}$, the sigma-algebra on $A_{-\infty}^{n-1}$; (ii) for every $\omega_{-\infty}^{-1} \in A_{-\infty}^{-1}$, $\sum_{\omega(n) \in A} P_n[\omega(n) | \omega_{-\infty}^{-1}] = 1$.}
2.4 Markov chains

Straightforward examples of Gibbs distributions defined in this way are provided by Markov chains with positive transition probabilities. Recall that a Markov chain of length $D$ is defined by a set of transition probabilities $P_n[\omega(n) | \omega_{n-D}^{n-1}]$ where the memory depth $D > 0$ is finite. These transition probabilities are obviously continuous with respect to $\omega$. If we assume moreover that they are strictly positive $\forall n \in \mathbb{Z}, \forall \omega_{n-D}^{n-1} \in \mathcal{A}_{n-D}^{n-1}$ then they match (ii) in the definition above. Finally, in this case, (i) is equivalent to the following property. For any time $n_1, n_2, n_2 - n_1 \geq D$:

$$\mu[\omega_{n_2}^{n_1}] = \prod_{l=n_1+D}^{n_2} P_l[\omega(l) | \omega_{l-D}^{l-1}] \mu[\omega_{n_1+D-1}^{n_1}] . \tag{2}$$

For any times $n_1, n_2$ as above, the Gibbs-probability $\mu[\omega_{n_2}^{n_1}]$ is given by the product of the Gibbs probability of the “initial block” $\mu[\omega_{n_1+D-1}^{n_1}]$ and the products of transition probabilities from the initial time $n_1 + D$ to the last time $n_2$.

Here we have considered transition probabilities depending explicitly on time $n$. When they are time-translation invariant (homogeneous Markov chain) the definition (2.1) is the definition of the unique invariant distribution of the Markov chain (it is unique because we have assumed positive transition probabilities).

Let us now state [2] in a different form. Define:

$$\phi_n(n, \omega) \overset{\text{def}}{=} \log P_n[\omega(n) | \omega_{n-D}^{n-1}], \tag{3}$$

called a normalized Gibbs potential. Then, [2] can be stated using:

$$\mu[\omega_{n_2}^{n_1} | \omega_{n_1+D-1}^{n_1}] = \exp \sum_{l=n_1+D}^{n_2} \phi_l(l, \omega) . \tag{4}$$

This form reminds the Gibbs distribution on spin lattices in statistical physics where one looks for lattice translation-invariant probability distributions given specific boundary conditions. Given a potential of range $D$ the probability of a spin block depends on the states of spins in a neighborhood of size $D$ of that block. Thus, the conditional probability of this block given a fixed neighborhood is the exponential of the energy characterizing physical interactions within the block as well as with the boundaries. Here, spins are replaced by spiking patterns; space is replaced with time which is mono-dimensional and oriented: there is no dependence in the future. Boundary conditions are replaced by the dependence in the past.

The definition [3] of the normalized Gibbs potential extends to the case $D \to +\infty$.

\footnote{One also says that $\mu$ is compatible with the set of transition probabilities.}
3 Gibbs distributions and models of spike train statistics

In this section we review several examples of models/concepts used to analyze spike train statistics. All of them enter in the realm of Gibbs distributions defined above.

3.1 Maximum entropy models

The definition (2.1) affords time-dependent transition probabilities. On the opposite, in this section we assume that they do not depend explicitly on \( n \), or, equivalently, that they are time-translation invariant. This corresponds to the physical concept of stationarity. We denote \( \mathcal{M}_{\text{inv}} \) the set of time-translation invariant probability measures on \( \Omega \).

Assume that spike trains statistics is distributed according to an hidden probability \( \mu \). How to approach \( \mu \) from data? Maximum entropy provides a method that allows to approach \( \mu \). It selects among all the probability distributions consistent with empirical data constraints, the most random i.e. the one with the highest entropy. But, why should we choose the maximum entropy distribution? The answer is that since entropy is a measure of information, then one should choose the probability that includes the least amount of information we have about the system and no more. The result probability is a Gibbs distribution.

3.1.1 Entropy

We define the entropy rate (or Kolmogorov-Sinai entropy) of a probability \( \mu \in \mathcal{M}_{\text{inv}} \) the set of time-translation invariant probability measures as:

\[
  h[\mu] = -\limsup_{n \to \infty} \frac{1}{n+1} \sum_{\omega_0^n} \mu[\omega_0^n] \log \mu[\omega_0^n],
\]

where the sum holds over all possible blocks \( \omega_0^n \). Note, that in the case of a Markov chain \( h[\mu] \) also reads [15]:

\[
  h[\mu] = -\sum_{\omega_0^D} \mu[\omega_0^D] \log \mu[\omega_0^D],
\]

Finally, when \( D = 0 \), \( h[\mu] \) reduces to the usual definition:

\[
  h(\mu) = -\sum_{\omega(0)} \mu[\omega(0)] \log \mu[\omega(0)].
\]

We used here the notation \( h(\mu) \) instead of \( S \) or \( s \), used in statistical physics. This is the conventional notation in ergodic theory for the (Kolmogorov-Sinai) entropy where the dependence on the measure \( \mu \) is made explicit.

3.1.2 Observables

We call observable a function:
\[ O : \Omega \rightarrow \{0, 1\}, \]
\[ \omega \mapsto \prod_{u=1}^{r} \omega_{k_u}(n_u) \]  
(8)
i.e. a product of binary spike events where \( k_u \) is a neuron index and \( n_u \) a time index, with \( u = 1, \ldots, r \), for some integer \( r > 0 \). Typical choices of observables are \( \omega_{k_1}(n_1) \) which is 1 if neuron \( k_1 \) fires at time \( n_1 \) and is 0 otherwise; \( \omega_{k_1}(n_1) \omega_{k_2}(n_2) \) which is 1 if neuron \( k_1 \) fires at time \( n_1 \) and neuron \( k_2 \) fires at time \( n_2 \) and is 0 otherwise. Another example is \( \omega_{k_1}(n_1) (1 - \omega_{k_2}(n_2)) \) which is 1 if neuron \( k_1 \) fires at time \( n_1 \) and neuron \( k_2 \) is silent at time \( n_2 \). This example emphasizes that observables are able to consider events where some neurons are silent.

We say that an observable \( O \) has range \( R \) if it depends on \( R \) consecutive spike patterns, e.g. \( O(\omega) = O(\omega_{0}^{R-1}) \). We consider here that observables do not depend explicitly on time (time-translation invariance of observables). As a consequence, for any time \( n \), \( O(\omega_{0}^{R-1}) = O(\omega_{n}^{n+R-1}) \) whenever \( \omega_{0}^{R-1} = \omega_{n}^{n+R-1} \).

### 3.1.3 Potential

A function of the form:

\[ H_\beta : \Omega \rightarrow \mathbb{R}, \]
\[ \omega \mapsto \sum_{k=1}^{N} \beta_k O_k. \]  
(9)
is called a potential, where the coefficients \( \beta_k \) are finite real numbers. The range of the potential is the maximum of the range of the observables \( O_k \).

### 3.1.4 Variational principle

Fix a potential \( H_\beta \) as in (9). Assume that it has finite range \( D \). In this case, a Gibbs distribution \( \mu \) obeys the following variational principle:

\[ P[H_\beta] = \sup_{\nu \in \mathcal{M}_{inv}} (h[\nu] + \nu[H_\beta]) = h[\mu] + \mu[H_\beta], \]  
(10)where \( P[H_\beta] \) is called the topological pressure and \( \nu[H_\beta] = \sum_{k=1}^{N} \beta_k \nu[O_k] \) is the average value of \( H_\beta \) with respect to the probability \( \nu \). We use the notation \( \nu(f) \) for the average of a function \( f \) instead of \( <f> \) used in statistical physics or \( E_\nu(f) \) used in probability theory. Note that Observables and Gibbs potentials are random functions that acts on the set of raster plots \( \Omega \).

Looking at the second equality, the variational principle selects, among all possible probabilities \( \nu \), a unique one, the Gibbs distribution, realizing the supremum. A variant of this principle holds when the average value of observables \( O_k \) is constrained to a value \( C_k \), fixed e.g. by experimental observations.

\[ ^3 \text{Thus, we do not consider here hard core potentials with forbidden configurations.} \]

\[ ^4 \text{The variational principle still holds if the range is infinite and its variation \( 1 \) decays sufficiently fast with \( m \), typically exponentially} \]

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In this case $\nu [\mathcal{H}_\beta]$ becomes $\sum_{k=1}^N \beta_k C_k$ if the average value of all observables $O_k$ is constrained. In this case the variational principle (10) reduces to maximizing the entropy on the set of measures $\nu \in \mathcal{M}_{inv}$ such that $\nu [O_k] = C_k$. Then, one is led to a classical Lagrange multipliers problem where the $\beta_k$s are the Lagrange multipliers. This is the classical approach introduced by Jaynes [25]. In this setting (10) signifies: “maximizing the entropy given the information that we have of the system” i.e. the observed average value of the observables $O_k$ is $C_k$.

### 3.1.5 Topological pressure

The topological pressure is the formal analogue of free energy density. It has the following properties:

- $\mathcal{P} [\mathcal{H}_\beta]$ is a log generating function of cumulants. We have:

\[
\frac{\partial \mathcal{P} [\mathcal{H}_\beta]}{\partial \beta_k} = \mu [O_k].
\]  
  (11)

and

\[
\frac{\partial^2 \mathcal{P} [\mathcal{H}_\beta]}{\partial \beta_k \partial \beta_l} = \frac{\partial \mu [O_k]}{\partial \beta_l} = \sum_{n=0}^{+\infty} C_{O_k O_l} (n),
\]  
  (12)

where

\[
C_{O_k O_l} (n) = \mu [O_k O_l \circ \sigma^n] - \mu [O_k] \mu [O_l],
\]

is the correlation function between the two observables $O_k$ and $O_l$ at time $n$ and $\sigma$ is the time shift operator. Note that correlation functions decay exponentially fast whenever $\mathcal{H}_\beta$ has finite range. So that $\sum_{n=0}^{+\infty} C_{O_k O_l} (n) < +\infty$.

Eq. (12) characterizes the variation in the average value of $O_k$ when varying $\beta_l$ (linear response). The corresponding matrix is a susceptibility matrix. It controls the Gaussian fluctuations of observables around their mean (central limit theorem) [46, 38, 12].

- $\mathcal{P} (\mathcal{H}_\beta)$ is a convex function of $\beta$.

- Define:

\[
Z_n = \sum_{\omega^0_n} e^{\mathcal{H}_\beta (\omega^0_n)}.
\]  
  (13)

The topological pressure obeys:

\[
\mathcal{P} (\mathcal{H}_\beta) = \lim_{n \to +\infty} \frac{1}{n} \log Z_n,
\]

and is analogous to a thermodynamic potential density (free energy, free enthalpy, pressure).

**Remark 1** For $D > 0$ one cannot write the Gibbs distribution in the form:

\[
\mu [\omega^0_n] = \frac{1}{Z_n} e^{\mathcal{H}_\beta (\omega^0_n)}.
\]  
  (14)
It only obeys: \( \exists A, B > 0 \) such that, for any block \( \omega_0^n \)
\[
A \leq e^{-(n-D+1)P(H_\beta)} e^{H_\beta(\omega_0^n)} \leq B.
\]
This is actually the definition of Gibbs distributions in ergodic theory \[12\].

3.1.6 Markov chain

The choice of the potential \[9\], i.e. the choice of a set of observables, fixes the restrictions for the statistical model. A normalization procedure allows to find a normalized potential \( \phi \) equivalent to \( H_\beta \) from which the transition probabilities are constructed. This defines an homogeneous Markov chain whose invariant measure is the Gibbs distribution associated with \( H_\beta \). It is constructed as follows.

**Transition matrix**

Consider two spike blocks \( w_1, w_2 \) of range \( D \geq 1 \). The transition \( w_1 \to w_2 \) is legal if \( w_1 \) has the form \( \omega(0)\omega_{D-1}^1 \) and \( w_2 \) has the form \( \omega_{D-1}^1\omega(D) \). The vectors \( \omega(0), \omega(D) \) are arbitrary but the block \( \omega_{D-1}^1 \) is common. Here is an example of a legal transition:

\[
w_1 = \begin{bmatrix} 0 & 0 & 1 \\ 0 & 1 & 1 \\ 1 & 1 & 0 \end{bmatrix}; w_2 = \begin{bmatrix} 0 & 1 & 1 \\ 1 & 1 & 0 \end{bmatrix}.
\]

Here is an example of a forbidden transition

\[
w_1 = \begin{bmatrix} 0 & 0 & 1 \\ 0 & 1 & 1 \\ 1 & 1 & 0 \end{bmatrix}; w_2 = \begin{bmatrix} 0 & 1 & 1 \\ 0 & 1 & 0 \end{bmatrix}.
\]

Any block \( \omega_0^D \) of range \( R = D + 1 \) can be viewed as a legal transition from the block \( w_1 = \omega_0^{D-1} \) to the block \( w_2 = \omega_0^D \) and in this case we write \( \omega_0^D \sim w_1 w_2 \).

The transfer matrix \( L \) is defined as:

\[
L_{w_1, w_2} = \begin{cases} e^{H_\beta(\omega_0^D)} & \text{if } w_1, w_2 \text{ is legal with } \omega_0^D \sim w_1 w_2, \\ 0, & \text{otherwise.} \end{cases}
\]

**Perron-Frobenius theorem**

From the matrix \( L \) the transition matrix of a Markov chain can be constructed. Since \( H_\beta(\omega_0^D) > -\infty, e^{H_\beta(\omega_0^D)} > 0 \) for each legal transition. As a consequence of the Perron-Frobenius theorem \[24,35\], \( L \) has a unique real positive eigenvalue \( s_\beta \), strictly larger than the modulus of the other eigenvalues (with a positive gap), and with associated right, \( R \), and left, \( L \), eigenvectors: \( LR = s_\beta R, LL = s_\beta L \).

The following holds:

- These eigenvectors have strictly positive entries \( R(\cdot) > 0, L(\cdot) > 0 \), functions of blocks of range \( D \). They can be chosen so that the scalar product \( \langle L, R \rangle = 1 \).

---

\(^{5}\)Two potentials are said “equivalent” or cohomologous if and only if they correspond to the same Gibbs distribution \[26\]
We have:
\[ \mathcal{P}(\mathcal{H}_\beta) = \log s_\beta. \]  
(16)

The following potential:
\[ \phi(\omega_0^D) = \mathcal{H}_\beta(\omega_0^D) - G_\beta(\omega_0^D) \]  
(17)

with:
\[ G_\beta(\omega_0^D) = \log R(\omega_0^{D-1}) - \log R(\omega_1^D) + \log s_\beta, \]  
(18)

is equivalent to \( \mathcal{H}_\beta \) and normalized. It defines a family of transition probabilities:
\[ \mathcal{P}[\omega(D) \mid \omega_0^{D-1}] \overset{\text{def}}{=} e^{\phi(\omega_0^D)} > 0. \]  
(19)

These transition probabilities define a Markov chain which admits a unique invariant probability:
\[ \mu(\omega_0^{D-1}) = R(\omega_0^{D-1}) L(\omega_1^{D-1}). \]  
(20)

which is the Gibbs distribution satisfying the variational principle \(10\).

It follows that the probability of blocks of depth \( n \geq D \) is:
\[ \mu[\omega_n^0] = \frac{e^{\mathcal{H}_\beta(\omega_n^0)}}{s_\beta^{n-D+1}} R(\omega_n^{D-1}) L(\omega_1^{D-1}). \]  
(21)

In the case \( D = 0 \) the Gibbs distribution reduces to \(14\). One can indeed easily show that:

\[ \exp G_\beta = s_\beta = \sum_{\omega(0)} e^{\mathcal{H}_\beta(\omega(0))} = Z_\beta. \]

Additionally, since spike patterns occurring at distinct time are independent in the \( D = 0 \) case, \( Z_n \) in \(13\) can be written as \( Z_n = Z_\beta^n \) so that \( \mathcal{P}(\mathcal{H}_\beta) = \log Z_\beta \).

In the general case of spatio-temporal constraints, the normalization requires the consideration of normalizing function \( G_\beta \) depending as well on the blocks \( \omega_0^D \). Thus, in addition to function \( \mathcal{H}_\beta \) normalization introduces a second function of spike blocks. This increases consequently the complexity of Gibbs potentials and Gibbs distributions compared to the spatial \( (D = 0) \) case where \( G_\beta \) reduces to a constant.

### 3.1.7 Examples

We give here a few examples of Maximum Entropy Gibbs distributions, found in the literature.

- **Bernoulli model.** Here only firing rates of neurons are constrained. The potential has the form:
\[ \mathcal{H}_\beta(\omega(0)) = \sum_{i=1}^{N} \beta_i \omega_i(0) \]
This is a memory-less model, where transitions probabilities are given by neuron firing rates \( \lambda_i = \frac{e^{\beta_i}}{1+e^{\beta_i}} \). The Gibbs distribution has the form:

\[
\mu[\omega^n_m] = \prod_{i=m}^{n} \prod_{k=1}^{N} \lambda_k^{\omega_k(l)} (1 - \lambda_k)^{1-\omega_k(l)},
\]

This is thus a product probability where neurons are independent.

- **Ising model.** This model was introduced by Schneidman et al [47] for retina spike train analysis. Here, firing rates and instantaneous pairwise synchronisation probabilities are constrained. The potential has the form:

\[
H_\beta(\omega(0)) = \sum_{i=1}^{N} \beta_i \omega_i(0) + \sum_{i,j=1}^{N} \beta_{ij} \omega_i(0) \omega_j(0).
\]

This is a memory-less model where the Gibbs distribution has the classical form (14).

- **Extended spatial Ising model.** A natural extension of Ising model has been proposed by Ganmor et al [19], where triplets and more general synchronous spike events are considered. The potential has the form:

\[
H_\beta(\omega(0)) = \sum_{i=1}^{N} \beta_i \omega_i(0) + \sum_{i,j=1}^{N} \beta_{ij} \omega_i(0) \omega_j(0) + \sum_{i,j,k=1}^{N} \beta_{ijk} \omega_i(0) \omega_j(0) \omega_k(0) + \ldots
\]

This is a memory-less model where the Gibbs distribution has the classical form (14).

- **Spatio temporal Ising model.** In [31] Marre et al considered a spatio-temporal extension of the Ising model where the potential has the form:

\[
H_\beta(\omega(0)) = \sum_{i=1}^{N} \beta_i \omega_i(0) + \sum_{i,j=1}^{N} \beta_{ij} \omega_i(0) \omega_j(1).
\]

Here spatio-temporal pairs with memory depth 1 are considered. Although the Gibbs distribution has not the form (14), the authors use an approximation of the exact distribution by this form, based on a detailed balance assumption. They applied this model for spike train analysis in the cat parietal cortex.

- **General Spatio temporal model.** General models of the form (9) have been considered in [55] for the analysis of retina spike trains. A C++ implementation of methods for fitting spatio-temporal models from data is available at [http://enas.gforge.inria.fr/v3/](http://enas.gforge.inria.fr/v3/).

### 3.1.8 Applications

The maximum entropy principle has been used by several authors [47, 49, 52, 56, 35, 47, 19, 20] for Multi-electrode Arrays (MEA) spike train analysis. Efficient methods have been designed to estimate the parameters of the potential, in the spatial case [17] (Broderick et al., 2007) and in the spatio-temporal case [34].
This approach, grounded on statistical physics, attempts to find a generic model for spike statistics based on a potential of the form \( \beta \), where the observables and their related \( \beta \) parameters summarize “effective interactions” between spikes. Behind this approach exists, we believe, a physicists “dream”: inferring, from data analysis, the equivalent of the equation of states existing in thermodynamics; that is, summarizing the behaviour of a big neuronal system by a few canonical variables (analogous e.g. to temperature, pressure, volume in a gas).

To our opinion, recent remarkable investigations to exhibit critical phenomena in retina spike train statistics are part of this project (Tkačič et al., 2006, 2009).

The main advantage of this approach is the possibility of constructing different statistical models based on a priori hypotheses on the most statistically significant events (single spikes, pairs, triplets, and so on). As such, it allows to consider arbitrary forms of spatio-temporal correlations. But this strength is also a weakness. Indeed, the possible forms of potentials are virtually infinite and obviously, in the setting of neuronal dynamics, one does not have the equivalent of mechanics or thermodynamics to construct the potential from general principles.

Finally, this approach only holds for stationary data, a highly questionable assumption as far as data from living systems are concerned.

### 3.2 Generalized Linear model

We now consider a second class of Gibbs distributions related to statistical models called Linear-Nonlinear (LN) model and Generalized Linear Model (GLM) \[4, 32, 50, 37, 54, 42, 40, 1, 41\]. We focus here on the GLM and follow the presentation of Ahmadian et al \[1\].

#### 3.2.1 Conditional intensities

GLMs are commonly used statistical methods for modeling the relationship between neural population activity and presented stimuli. Let \( x \equiv x(t) \) be a time-dependent stimulus. In response to \( x \) the network emits a spike train response \( r \). This response does not only depend on \( x \), but also on the network history of spiking activity. The GLM (and LN) assimilate the spike response \( r \) as an inhomogeneous point process: the probability that neuron \( k \) emits a spike between \( t \) and \( t + dt \) is given by \( \lambda_k(t \mid H_t) dt \), where \( \lambda_k(t \mid H_t) \) is called “conditional intensity” and \( H_t \) is the history of spiking activity up to time \( t \). In the GLM this function is given by:

\[
\lambda_k(t \mid H_t) = f \left( b_k + (K_k * x)(t) + \sum_j (H_{kj} * r_j)(t) \right), \tag{23}
\]

where:

- \( f \) is a non linear function (an exponential or a sigmoid);
- \( b_k \) is some constant fixing the baseline firing rate of neuron \( k \);
- \( K_k \) is a causal, time-translation invariant, linear convolution kernel that mimics a linear receptive field of neuron \( k \);
• ∗ is the convolution product;

• \( H_{kj} \) is the memory kernel that describes possible excitatory or inhibitory post spike effects of the \( j^{th} \) observed neuron on the \( k^{th} \). As such, it depends on the past spikes, hence on \( \omega \). The diagonal components \( H_{kk} \) describe the post spike feedback of the neuron to itself, and can account for refractoriness, adaptation and burstiness depending on their shape;

• \( r_j \) is the spike train of neuron \( j \): \( r_j(t) = \sum_{r \geq 1} \delta(t-t_j^{(r)}) \), where \( t_j^{(r)} \) is the time of the \( r^{th} \) spike of \( j^{th} \) neuron.

The spike response has a history dependent structure that makes Poisson models inappropriate. Point processes affords for history dependence and generalizes Poisson process. A point process can be completely characterized by its conditional intensity function.

\[
\lambda_k(t \mid H_t) = \lim_{\Delta t \to 0} \frac{P(N_{[t+\Delta t]}=1 \mid H_t)}{\Delta t},
\]

where \( N_{[t+\Delta t]} \) is the counting process that gives the number of spikes occurring in the interval \([t + \Delta t])\). Choosing \( \Delta t \) to be a sufficiently small time interval \( \sim 1\text{ms} \), the probability of firing more than one spike is negligibly small compared to the probability of firing one spike. This assumption is biophysically plausible because neurons have refractory period. Therefore:

\[
P(\text{spike in } [t + \Delta t] \mid H_t) \approx \lambda_k(t \mid H_t)\Delta t.
\]

Here \( \lambda_k(t \mid H_t) \) is defined in continuous time, and spikes are discrete events. If we discretize the time to make the spikes emitted by the point process belong to a single bin, we have:

\[
P(\omega_k(n) = 1 \mid H_{n-1}) \approx \lambda_k(n \mid H_{n-1})\Delta t := p_k(n)
\]

### 3.2.2 Conditional independence

The GLM postulates that, given the history \( H \) and stimulus \( x \), neurons are independent (conditional independence upon the past and stimulus). In the context of transition probabilities defined on section 2.2, the response at time \( n \) is a spiking pattern \( \omega(n) \) while the history is the spike activity \( H \). As a consequence of the conditional independence assumption the probability of a spike pattern follows a Bernoulli process:

\[
P_n[\omega(n) \mid \omega_{-\infty}^{n-1}] = \prod_{k=1}^{N} p_k(n)^{\omega_k(n)}(1-p_k(n))^{1-\omega_k(n)}.
\]

### 3.2.3 Gibbs distribution

Transition probabilities are strictly positive whenever \( 0 < p_k(n) < 1 \), for all \( k,n \). If \( f \) is e.g. a sigmoid this holds provided its argument \( b_{i} + (K_{i} \ast x)(t) + \sum_{j}(H_{ij} \ast r_{j})(t) \) remains bounded in absolute value. The continuity of \( \lambda \) with respect to \( \omega \) holds whenever \( f \) is continuous and the memory kernel \( H \) is continuous with respect to \( \omega \). This second condition is fulfilled in two cases:
• $H$ depends on a finite past;
• $H$ depends on an infinite past, but the memory dependence decays sufficiently fast to ensure continuity. Since $H$ mimics synaptic influence it is typically a sum of $\alpha$-profiles that mimic PSPs (Post Synaptic Potentials). $\alpha$ profiles decay exponentially fast with time, so they match this condition. We come back to this point in section 3.3.

The Gibbs potential associated with (24) is:

$$\phi_n(\omega) = \sum_{k=1}^{N} \left( \omega_k(n) \log p_k(n) + (1 - \omega_k(n))(1 - p_k(n)) \right),$$  

(25)

It is normalized by definition.

### 3.2.4 Applications

This model has been applied in a wide variety of experimental settings [5, 13, 53, 6, 36, 54, 40]. Efficient methods has been designed to estimate the parameters [1].

To us, the main advantages of the GLM are:
• The transition probability is known (postulated) from the beginning and does not require the heavy normalization (17) imposed by potentials of the form (9);
• The model parameters have a neurophysiological interpretation, and their number grows at most as a power law in the number of neurons.
• It has good decoding performances
• It holds for non stationary data.

Its main drawbacks are:
• It postulates an ad hoc form for the transition probability of the stochastic process;
• It uses a quite questionable assumption of conditional independence: neurons are assumed independent at time $n$ when the past is given. On the opposite, the maximal entropy principle does not require this assumption.
• To us, the biophysical interpretation of the parameters $H_{kj}$ is unclear. Do they correspond to “real” connectivity? “functional” connectivity?

### 3.3 Integrate and Fire neural networks

The previous examples were mainly developed for data analysis: one speculates a form for transitions probabilities, performs parameters fitting, and then uses the model to decode or to extrapolate the statistics of complex events. Here we start from a different point of view asking the following questions: Can we have a reasonable idea of what could be the spike train statistics studying a neural network model? Do Gibbs distribution arise in these models? What is the shape of the potential? We focus here on a model proposed in [7, 8, 14] where these questions have been answered.
3.3.1 Model

The integrate-and-fire model remains one of the most ubiquitous models for simulating and analyzing the dynamics of neuronal circuits. Despite its simplified nature, it captures some of the essential features of neuronal dynamics. Denote $V(t)$ the membrane potential vector with entries $V_k(t)$. The continuous-time dynamics of $V(t)$ is defined as follows. Fix a real variable $\theta > 0$ called “firing threshold”. For a fixed time $t$, we have two possibilities:

1. Either $V_k(t) < \theta$, $\forall k = 1, \ldots, N$. This corresponds to sub-threshold dynamics.

2. Or, $\exists k, V_k(t) \geq \theta$. Then, we speak of firing dynamics.

The model proposed here is an extension of the conductance based Integrate-and-Fire neuron model introduced in [45]. The model-definition follows the presentation given in [11, 8]. Neurons are considered as points, with neither spatial extension nor biophysical structure (axon, soma, dendrites). Dynamics is ruled by a set of stochastic differential equations where parameters, corresponding to chemical conductances, depend on the action potentials emitted in the past by the neurons. In this way, the dynamical system defined here is ruled both by continuous and discrete time dynamical variables.

Subthreshold dynamics

It is defined by:

$$C_k \frac{dV_k}{dt} = -g_{L,k}(V_k - E_L) - \sum_j g_{kj}(t, \omega)(V_k - E_j) + \sum_j g_{kj}(V_j - V_k) + I_k(t),$$

(26)

where:

- $C_k$ is the membrane capacity of neuron $k$;
- $I_k(t) = i^{(ext)}_k(t) + \sigma_B \xi_k(t)$ is a current when a time-dependent part $i^{(ext)}_k(t)$ (stimulus) and stochastic part $\sigma_B \xi_k(t)$ where $\xi_k(t)$ is a white noise and $\sigma_B$ controls the noise intensity;
- $g_{L,k}$ is the leak conductance and $E_L < 0$ the leak Nernst potential;
- $g_{kj}$ mimics electric conductance (gap junctions) between neurons $j$ and $k$; these are passive and symmetric conductances;
- the term

$$g_{kj}(t, \omega) = G_{kj} \sum_{r: t^{(r)}_j(\omega) < t} \alpha_{kj} \left( t - t^{(r)}_j(\omega) \right),$$

(27)

mimics the conductance of the chemical synapse $j \rightarrow k$, where:

$$\alpha_{kj}(t) = \frac{t}{\tau_{kj}} e^{-\frac{t}{\tau_{kj}}} H(t),$$

(28)

mimics a PSP, $H(t)$ is the Heaviside function (that mimics causality); $t^{(r)}_j(\omega)$ is the $r$th spike emitted by neuron $j$ in the raster $\omega$, therefore $g_{kj}(t, \omega)$ depends on the whole spike history; $E_j$ is the reversal potential of the chemical synapse $j \rightarrow k$. 

15
Firing dynamics and reset

If, at time $t$, some neuron $k$ reaches its firing threshold $\theta$, $V_k(t) = \theta$, then this neuron emits a spike. To conciliate the continuous time dynamics of membrane potentials and the discrete time dynamics of spikes we define the spike and reset as follows.

- The neuron membrane potential $V_k$ is reset to 0 at the next integer time after $t$.
- A spike is registered at time $[t] + 1$ where $[t]$ is the integer part of $t$. This allows us to represent spike trains as events on a discrete time grid. It has the drawback of artificially synchronizing spikes coming from different neurons, in the deterministic case [11, 28]. However, the presence of noise in membrane potential dynamics eliminates this synchronization effect.
- Spikes are separated by a time scale $\tau_{sep} > 0$ which is a multiple of $\delta$ (thus an integer).
- Between $[t] + 1$ and $[t] + \tau_{sep}$ the membrane potential $V_k$ is maintained to 0 (refractory period). From time $[t] + \tau_{sep}$ on, $V_k$ evolves according to (26) until the next spike.
- When the spike occurs (at time $[t] + 1$), the raster $\omega$ as well conductances $g_{kj}(t, \omega)$ are updated.

3.3.2 Main results

This model has several variants: discrete time [7]; continuous time with chemical synapses [8] and continuous time with chemical and electric synapses [14]. We list here the main results concerning spike statistics and Gibbs distributions.

1. Whatever the values of the parameters the model admits a unique Gibbs distribution in the general sense given in section 2.3.

2. When the noise is weak and without gap junctions, the normalized Gibbs potential can be explicitly computed. It takes the form:

$$\phi_n(\omega) = \sum_{k=1}^{N} \left( \omega_k(n) \log \lambda_k(n) + (1 - \omega_k(n)) \log(1 - \lambda_k(n)) \right), \quad (29)$$

where

$$\lambda_k(n) = f \left( b_k(n - 1, \omega) + \Phi_k^{(ext)}(n - 1, \omega) + \Phi_k^{(syn)}(n - 1, \omega) \right), \quad (30)$$

where:

- $f$ is a sigmoid function;
- $b_k(n - 1, \omega)$ is a function depending on the threshold value, the leak Nernst potential, and on the integrated noise, integrated from the last time where has been reset (depending on $\omega$) up to time $n - 1$;
- $\Phi_k^{(ext)}(n - 1, \omega)$ corresponds to the integrated effects of the external current $i_k^{(ext)}$ on the membrane potential;
the term $\Phi^{(\text{syn})}_k(n-1, \omega)$ corresponds to the integrated effects of chemical synapses on the membrane potential.

3. Eq. (29), expresses that in this case, neurons are conditionally independent upon the past.

4. In this conductance-based model, conductances depend on the past via (27). One can consider as well a current-based model where conductances are fixed and current depend on the past spikes. In this case, the terms $\Phi^{(\text{ext})}_k(n-1, \omega)$ and $\Phi^{(\text{syn})}_k(n-1, \omega)$ can be written as convolutions and one recovers a potential with a form analogous to (25).

5. In the general case (gap junctions), neurons are not conditionally independent. Gap junctions induce a coupling effect which does not allow any more the factorization (29) of the potential.

6. Correlations (pairwise and higher order) are mainly due to chemical synapses and gap junctions. Additional correlations can also be induced by the stimulus using e.g. a current $i^{(\text{ext})}_j$ where time fluctuations of $i^{(\text{ext})}_j$ are correlated with $i^{(\text{ext})}_j$. But these are extra-correlations that disappear when the stimulus is removed, whereas the dynamical correlations remains.

7. The potential has an infinite range (infinite memory). However, thanks to the exponential decay of the alpha profile, one can show that the potential is continuous. This allows to propose Markovian approximation of the Gibbs distribution where the exact potential is replaced by a potential with a finite range [7, 8].

### 3.3.3 Applications

What do we finally learn from the study of this model?

- We have a positive answer to the existence of Gibbs distributions in neural networks models.
- An explicit form for the potential is known in specific cases.
- The form (30) actually also fits with maximum entropy model, in the stationary case, as shown in section 3.4.
- In this model the origin of correlations is essentially due to dynamics, not to the stimulus.
- Gap junctions play here a central role in the structure of dynamical correlations and dependence of dynamics upon history (see [14] for more details).
- The analysis holds for non stationary data.

Considering potential uses of this study to fit real data the main criticism is:
• This is a model. Is it sufficient to describe real neural networks? For example, its application to retina data is controversial since it considers only spiking cells (that mimics ganglion cells), but retina has also non firing cells like most amacrine and bipolar cells.

• In the general case there is no explicit form for the Gibbs potential.

• Even when there exists an explicit form for the potential, it has quite a lot of parameters which can be difficult to fit from data.

3.4 Relations between these approaches

In this section, we establish a connection between the three examples of Gibbs distributions considered in sections 3.1, 3.2, 3.3.

Consider a family of transition probabilities satisfying the positivity condition (ii) in section 2.3, where we furthermore assume that the memory depth is finite and that transition probabilities are time-translation invariant. As stated in section 2.4, this define an homogenous Markov chain. The transition probabilities are thus functions of blocks $\omega_D^D$ (see section 2.2 and the definition of time translation invariance). These functions can then take at most $2^{N(D+1)}$ values. The same holds for the normalized potential (3). Now, one can prove that any such function can be written as:

$$
\phi(\omega) \equiv \phi(\omega_D^D) = \sum_{l=0}^{L} \phi_l \mathcal{O}_l(\omega),
$$

with $L = 2^{N(D+1) - 1}$ and $\mathcal{O}_l$ is an observable where the time index ranges from 0 to $D$. The index $l$ parametrizes an enumeration of all possible observables with $N$ neurons and $D + 1$ time steps, where $m_0$ is the constant observable $\mathcal{O}_0 = 1$, and so on.

Now, using the positivity condition and the results in section 3.1, one can show that any family of stationary transition probabilities with memory depth $D$ can be associated with a potential of the form (9). The correspondence is actually unique. This is a straightforward application of the celebrated Hammersley-Clifford theorem [24, 2, 33, 39].

An immediate consequence of this result is that, in the stationary case with finite memory, the GLM potential (25) and the Integrate and Fire (29) correspond to a Maximum Entropy model with a potential of the form (9). The parameters $\beta_k$ in (9) are then nonlinear functions of the parameters in (25) or in (29) (see [7]). As a consequence, some of these parameters are redundant: there are a priori $2^{N(D+1)}$ non vanishing parameters $\beta_k$ while there are quite less parameters in the GLM or in the Integrate and Fire (of order $N^2$). However, the GLM assumes conditional independence between neurons, while the maximum entropy approach is precisely used to take care of (pairwise and higher order) correlations between neurons. In this sense it is more general.

In the non stationary case one can no more apply the maximum entropy principle (entropy is not defined). However, in the case where statistics depends on time on a slow time scale (compared to spike characteristic time scale) one
can use a quasi-static approach where the parameters $\beta_k$ in (9) vary slowly in time [44] (Tyrcha et al., 2012). On the opposite, the GLM allows to consider non stationary data with efficient results [41].

The IF model contains both maximum entropy models and GLM. It has a maximum entropy Gibbs distribution in the stationary case, and it reduces to GLM upon several simplifications. In its more general form it allows the consideration of non stationarity and does not rely on the conditional independence assumption. Unfortunately, its generality is a weakness since an explicit form for the potential is not known yet in the general case.

4 Conclusion

In this paper we have argued that Gibbs distribution considered in a fairly general sense could constitute generic statistical models to fit spike trains data. The example of the Integrate and Fire model suggests that such distribution could be also defined for more elaborated neural networks models (FitzHugh-Nagumo or Hodgkin-Huxley). In particular, the existence and uniqueness of a Gibbs measure holds whenever there is continuity with respect to a raster, with a sufficiently fast decay of the variation [1] [13]. As shown in [7, 8, 14] this property is ensured when interactions between neurons decay exponentially fast. This is typically the case for chemical synapses where the PSP [28] decays exponentially fast with time.

The interest of proposing Gibbs distribution constructed from neural network models is multiple. The model mimics a neurophysiological structure where interactions between neurons, stimuli, and biophysical parameters are well identified. As a consequence the model-parameters can be easily interpreted. Thus, the role of each specific biophysical parameter on spike statistics can be easily analysed. Also, the potential obtained this way is already normalized, while e.g. maximum entropy principle requires a complex procedure to achieve normalization. Finally, in this context, it is possible to study the effect of a time-dependent stimulus on spike statistics.

However, this approach, to be efficient requires (i) to have an analytical form for the potential; (ii) to be able to fit the many parameters of a non linear problem. This is yet far from being achieved.

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References

[1] Yashar Ahmadian, Jonathan W. Pillow, and Liam Paninski. Efficient Markov Chain Monte Carlo Methods for Decoding Neural Spike Trains. Neural Computation, 23(1):46–96, January 2011.

[2] J. Besag. Spatial interaction and the statistical analysis of lattice systems. Journal of the Royal Statistical Society. Series B (Methodological), 36(2):192–236, 1974.
[3] R. Bowen. Equilibrium states and the ergodic theory of Anosov diffeomorphisms, volume 470 of Lect. Notes in Math. Springer-Verlag, New York, 1975.

[4] D. R. Brillinger. Maximum likelihood analysis of spike trains of interacting nerve cells. Biol Cybern, 59(3):189–200, 1988.

[5] D. R. Brillinger. Nerve Cell Spike Train Data Analysis - a Progression of Technique. J Amer Statist Assn, 87(418):260–271, 1992.

[6] E. N. Brown, R. Barbieri, U. T. Eden, and L. M. Frank. Likelihood methods for neural spike train data analysis. Computational Neuroscience: A Comprehensive Approach, 2003.

[7] Bruno Cessac. A discrete time neural network model with spiking neurons ii. dynamics with noise. J. Math. Biol., 62:863–900, 2011.

[8] Bruno Cessac. Statistics of spike trains in conductance-based neural networks: Rigorous results. Journal of Mathematical Neuroscience, 1(8), 2011.

[9] Bruno Cessac and Adrian Palacios. Spike train statistics from empirical facts to theory: the case of the retina, volume in “Current Mathematical Problems in Computational Biology and Biomedicine”. Springer, 2012.

[10] Bruno Cessac, Héléne Paugam-Moisy, and Thierry Viéville. Overview of facts and issues about neural coding by spikes. J. Physiol. Paris, 104(1-2):5–18, February 2010.

[11] Bruno Cessac and T. Viéville. On dynamics of integrate-and-fire neural networks with adaptive conductances. Frontiers in neuroscience, 2(2), July 2008.

[12] J.R. Chazottes and G. Keller. Mathematics of Complexity and Dynamical Systems, pages 1422–1437. Springer, 2011.

[13] E. J. Chichilnisky. A simple white noise analysis of neuronal light responses. Network: Comput. Neural Syst., 12:199–213, 2001.

[14] Rodrigo Cofré and Bruno Cessac. Dynamics and spike trains statistics in conductance-based integrate-and-fire neural networks with chemical and electric synapses. Chaos, Solitons and Fractals, 2013. In press.

[15] I. P. Cornfeld, S. V. Fomin, and Ya. G. Sinai. Ergodic Theory. Springer, Berlin, Heidelberg, New York, 1982.

[16] Vere-Jones D. Daley, D. J. An introduction to the theory of point processes (2nd ed.). New York: Springer., 2003.

[17] M. Dudík, S. Phillips, and R. Schapire. Performance guarantees for regularized maximum entropy density estimation. In Proceedings of the 17th Annual Conference on Computational Learning Theory, 2004.

[18] Roberto Fernandez and Grégory Maillard. Chains with complete connections : General theory, uniqueness, loss of memory and mixing properties. J. Stat. Phys., 118(3-4):555–588, 2005.
[19] Elad Ganmor, Ronen Segev, and Elad Schneidman. The architecture of functional interaction networks in the retina. The journal of neuroscience, 31(8):3044–3054, 2011.

[20] Elad Ganmor, Ronen Segev, and Elad Schneidman. Sparse low-order interaction network underlies a highly correlated and learnable neural population code. PNAS, 108(23):9679–9684, 2011.

[21] F. R. Gantmacher. The theory of matrices. AMS Chelsea Publishing, Providence, RI, 1998.

[22] Hans-Otto Georgii. Gibbs measures and phase transitions. De Gruyter Studies in Mathematics:9. Berlin; New York, 1988.

[23] Sebastian Gerwinn, Jakob Macke, and Matthias Bethge. Bayesian population decoding of spiking neurons. Front Comput Neurosci, page 3:21, 2009.

[24] J. M. Hammersley and P. Clifford. Markov fields on finite graphs and lattices. unpublished, 1971.

[25] E.T. Jaynes. Information theory and statistical mechanics. Phys. Rev., 106:620, 1957.

[26] O. Jenkinson, R. D. Mauldin, and M. Urbanski. Zero temperature limits of gibbs-equilibrium states for countable alphabet subshifts of finite type. J. Stat. Phys, 119:765–776, 2005.

[27] M. Keane. Strongly mixing g-measures. Invent. Math., 16:309–324, 1972.

[28] C. Kirst and M. Timme. How precise is the timing of action potentials? Front. Neurosci., 3(1):2–3, 2009.

[29] Shinsuke Koyama, Uri T. Eden, Emery N. Brown, and Robert E. Kass. Bayesian decoding of neural spike trains. Annals of the Institute of Statistical Mathematics, 62:pp 37–59, 2010.

[30] G. Maillard. Introduction to chains with complete connections. Ecole Federale Polytechnique de Lausanne, winter 2007.

[31] O. Marre, S. El Boustani, Y. Frégnac, and A. Destexhe. Prediction of spatiotemporal patterns of neural activity from pairwise correlations. Phys. rev. Let., 102:138101, 2009.

[32] P. McCullagh and J. A. Nelder. Generalized linear models (Second edition). London: Chapman & Hall, 1989.

[33] John Moussouris. Gibbs and markov random systems with constraints. J. Stat. Phys., 10(1):11–33, 1974.

[34] H. Nasser, O. Marre, and B. Cessac. Spatio-temporal spike trains analysis for large scale networks using maximum entropy principle and monte-carlo method. Journal Of Statistical Mechanics, 2013. in press.
[35] Ifije E. Ohiorhenuan, Ferenc Mechler, Keith P. Purpura, Anita M. Schmid, Qin Hu, and Jonathan D. Victor. Sparse coding and high-order correlations in fine-scale cortical networks. Nature, 466(7):617–621, 2010.

[36] L. Paninski, M. Fellows, S. Shoham, N. Hatsopoulos, and J. Donoghue. Superlinear population encoding of dynamic hand trajectory in primary motor cortex. J. Neurosci., 24:8551–8561, 2004.

[37] Liam Paninski. Maximum likelihood estimation of cascade point-process neural encoding models. Network: Comput. Neural Syst., 15(04):243–262, November 2004.

[38] W. Parry and M. Pollicott. Zeta functions and the periodic orbit structure of hyperbolic dynamics, volume 187–188. Asterisque, 1990.

[39] P. Clifford. Disorder in Physical Systems: A Volume in Honour of John M. Hammersley, pages 19–32. Oxford University Press, 1990.

[40] J.W. Pillow, J. Shlens, L. Paninski, A. Sher, A.M. Litke, E.J. Chichilnisky, and E.P. Simoncelli. Spatio-temporal correlations and visual signaling in a complete neuronal population. Nature, 454(7206):995–999, Aug 2008.

[41] Jonathan W. Pillow, Yashar Ahmadian, and Liam Paninski. Model-based decoding, information estimation, and change-point detection techniques for multineuron spike trains. Neural Comput., 23(1):1–45, 2011.

[42] J.W. Pillow, L. Paninski, V.J. Uzzell, E.P. Simoncelli, and E.J. Chichilnisky. Prediction and decoding of retinal ganglion cell responses with a probabilistic spiking model. J. Neurosci, 25:11003–11013, 2005.

[43] F. Rieke, D. Warland, Rob de Ruyter van Steveninck, and William Bialek. Spikes, Exploring the Neural Code. The M.I.T. Press, 1996.

[44] Yasser Roudi and John Hertz. Mean field theory for non-equilibrium network reconstruction. Phys. Rev. Lett., 106(048702), 2011.

[45] M. Rudolph and A. Destexhe. Analytical integrate and fire neuron models with conductance-based dynamics for event driven simulation strategies. Neural Computation, 18:2146–2210, 2006.

[46] D. Ruelle. Thermodynamic formalism. Addison-Wesley,Reading, Massachusetts, 1978.

[47] E. Schneidman, M.J. Berry, R. Segev, and W. Bialek. Weak pairwise correlations imply strongly correlated network states in a neural population. Nature, 440(7087):1007–1012, 2006.

[48] E. Seneta. Non-negative Matrices and Markov Chains. Springer, 2006.

[49] J. Shlens, G.D. Field, J.L. Gauthier, M.I. Grivich, D. Petrusca, A. Sher, A.M. Litke, and E.J. Chichilnisky. The structure of multi-neuron firing patterns in primate retina. Journal of Neuroscience, 26(32):8254, 2006.

[50] E. P. Simoncelli, J. P. Paninski, J. Pillow, and O. Schwartz. Characterization of Neural Responses with Stochastic Stimuli. The cognitive neurosciences, 2004.

22
[51] Miller M. I. Snyder, D. L. Random point processes in time and space. New York: Springer., 1991.

[52] Aonan Tang, David Jackson, Jon Hobbs, Wei Chen, Jodi L. Smith, Hema Patel, Anita Prieto, Dumitru Petrusca, Matthew I. Grivich, Alexander Sher, Pawel Hottowy, Wladyslaw Dabrowski, Alan M. Litke, and John M. Beggs. A maximum entropy model applied to spatial and temporal correlations from cortical networks In Vitro. The Journal of Neuroscience, 28(2):505–518, January 2008.

[53] F. E. Theunissen, S. V. David, N. C. Singh, A. Hsu, W. E. Vinje, and J. L. Gallant. Estimating spatio-temporal receptive fields of auditory and visual neurons from their responses to natural stimuli. Network, 12(3):289–316, January 2001.

[54] Wilson Truccolo, Uri T. Eden, Matthew R. Fellows, John P. Donoghue, and Emery N. Brown. A point process framework for relating neural spiking activity to spiking history, neural ensemble and extrinsic covariate effects. J Neurophysiol, 93:1074–1089, 2005.

[55] Juan Carlos Vasquez, Olivier Marre, Adrian G Palacios, Michael J Berry, and Bruno Cessac. Gibbs distribution analysis of temporal correlation structure on multicell spike trains from retina ganglion cells. J. Physiol. Paris, 106:120–127, 2012.

[56] Shan Yu, Debin Huang, Wolf Singer, and Danko Nikolic. A small world of neuronal synchrony. Cereb. Cortex, 2008.