Hawkes Processes Modeling, Inference and Control: An Overview

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Abstract—Hawkes Processes are a type of point process which model self-excitement among time events. It has been used in a myriad of applications, ranging from finance and earthquakes to crime rates and social network activity analysis. Recently, a surge of different tools and algorithms have showed their way up to top-tier Machine Learning conferences. This work aims to give a broad view of the recent advances on the Hawkes Processes modeling and inference to a newcomer to the field.

I. INTRODUCTION

Point Processes are tools for modeling the arrival of time events. They have been broadly used to model both natural and social phenomena related to arrival of events in a continuous time-setting, such as the queuing of customers in a given store, the arrival of earthquake aftershocks [36], [16], the failure of machines at a factory, the request of packages over a communication network, and the death of citizens in Ancient societies [8].

Predicting, and thus being able to effectively intervene in all these phenomena is of huge commercial and/or societal value, and thus there has been an intensive investigation of the theoretical foundations of this area.

Hawkes Processes (HP) are a type of point process which models self- and mutual-excitation, i.e., when the arrival of an event makes future events more likely to happen. They are suitable for capturing epidemic, clustering, and faddish behaviour on social and natural time-varying phenomena. The excitation effect is represented by an additional function to the intensity of the process (i.e., the expected arrival rate of events): the triggering kernel, which quantifies the influence of events of a given process in the self- and mutual triggering of its associated intensity functions. Much of the Hawkes Processes’ research has been devoted to modeling the triggering kernels, handling issues of scalability to large number of concurrent processes and quantity of data, as well as speed and tractability of the inference procedure.

Regarding the learning of the triggering kernels, one of the methods involves the assumption that it can be defined by simple parametric functions, s.a. one or multiple exponentials, Gaussians, Rayleigh, Mittag-Leffler [7] functions and power-laws. Much of the work dealing with this type of approach concerned with enriching these parametric models [20], [48], [54], [53], [11], scaling them for high dimensions, i.e., multivariate processes of large dimensions [5], [22], dealing with distortions related to restrictions on the type of available data [55], and proposing adversarial losses as a complement to the simple Maximum Likelihood Estimation (MLE) [57].

Another way of learning the triggering function is by assuming that it is represented by a finite grid, in which the triggering remains constant along each of its subintervals. Regarding this piecewise constant (or non-parametric) approach, most notably developed in [32], [6], the focus has been on speeding up the inference through parallelization and online learning of the model parameters [59], [1].

A more recent approach, enabled by the rise to prominence of Deep Learning models and techniques, which accompany the increase of computational power and availability of data of the recent years, regards modeling the causal triggering effect through the use of neural network models, most notably RNNS, LSTMs and GANs. These models allow for less bias and more flexibility than the parametric models for the modeling of the triggering kernel, while taking advantage of the numerous training and modeling techniques developed by the booming connectionist community.

In addition, regarding the control of self-exciting point processes, i.e., the modification of the process parameters towards more desirable configurations, while taking into account an associated ‘control cost’ to the magnitude of these modifications, recent works either make use of Dynamic Programming (Continuous Hamilton-Jacobi-Bellman Equation-based approach) [61], [60], Kullback-Leibler Divergence penalization (a.k.a. ‘Information Bottleneck’) [47], and Reinforcement Learning-based, as well as Imitation Learning-based techniques [45], [24].

Although there have been some interesting reviews and tutorials regarding domain-specific applications of Hawkes Processes in Finance [14], [5] and Social Networks [38], a broad view of the inference and modeling approaches is still lacking. A close work to ours is the one presented in [56] which, although very insightful, lacks coverage of important advances, such as the previously mentioned control approaches, as well as the richer variants of Neural Network-based models.

In the following, we introduce the mathematical definitions involving HPs, then carefully describe the advances on each of the aforementioned approaches, then finish by a summarization, along with some considerations.

II. THEORETICAL BACKGROUND

In this and the next section, we present the mathematical definitions used throughout the remaining sections of the paper.

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In the present work, we restrain ourselves to the Marked Temporal Point Processes (MTPP), i.e., the point process in which each event is defined by a time coordinate and a mark (or label). An intuitive example of a MTPP is shown in Figure 1.

The key definitions are those of: Counting Process, Intensity Function, Triggering Kernel, Impact Matrix, (Log-)likelihood, Covariance, Bartlet Spectrum, Higher-order Moments and Branching Structure.

A. Multivariate Marked Temporal Point Processes

Realizations of univariate MTPPs, here referred to by \( S \) are one or more sequences of events \( e_i \), each composed by the time coordinate \( t_k \) and the mark \( m_k \), s.a.:

\[
S = \{(t_0, m_0), ..., (t_N, m_N)\}
\]  

Marks may represent, for example, a specific user in a social network or a specific geographic location, among others. For more complex problems, as in the check-in times prediction of [58], a composite mark may represent an user of interest and a specific location.

An easy way to generalize this notation would be to refer to multiple realizations of multivariate MTPPs as \( S = S_{i,j} \), where \( i \) would refer to the dimension of the process, while \( j \) would refer to the index of the sequence. Now, regarding only the purely temporal portion of the process, i.e., the time coordinates \( t_k \), it is also common to express them by means of a Counting Process \( N(t) \), which is simply the cumulative number of event arrivals up to time \( t \):

\[
\int_0^t dN_s,
\]  

where:

- \( dN_{t_k} = 1 \), if there is an event at \( t_k \);
- \( dN_{t_k} = 0 \), otherwise.

This is illustrated in Figure 2. Associated to each temporal point process, there is an Intensity Function, which is the expected rate of arrival of events:

\[
\lambda(t)dt = E\{dN_t = 1\},
\]  

which may depend or not on the history of past events. Such dependence results in a so-called “Conditional Intensity Function” (CIF):

\[
\lambda(t)dt = E\{dN_t = 1|\mathcal{H}\},
\]  

where \( \mathcal{H} \) is the history of all events up to time \( t \):

\[
\mathcal{H} : \{t_{i,j} \in S|t_{i,j} < t\}
\]  

This concept will be further discussed in the next section.

B. Hawkes Processes

The simplest example of a temporal point process is the Homogeneous Poisson Process (HPP), in which the intensity is a positive constant:

\[
\lambda(t) = \mu,
\]  

for \( \mu \in \mathbb{R}_+ \).

In the case of an Inhomogeneous Poisson Process (IPP), the intensity \( \lambda(t) \) is allowed to vary. Both HPP and IPP are shown in Figure 3.

Both the Homogeneous and the Inhomogeneous Poisson Processes have, in common, the fact that each consecutive event interval sampled from the Intensity Function is independent of the previous ones. When analyzing several natural phenomena, one may wish to model how events in each
dimension $i$ of the process, which may be representing a specific social network user, an earthquake shock at a given geographical region, or a percentual jump in the price of a given stock, just to cite a few examples - affect the arrival of events in all the dimensions of the process, including its own.

In particular, we are interested in the cases where the arrival of one event makes further events more likely to happen, which is reflected as an increase in the value of the intensity function after the time of said event. When this increase happens to the intensity Function of the same $i$-th dimension of the event, the effect is denominated self-excitation. When the increase happens to the intensity of other dimensions, we refer to it as mutual excitation.

Hawkes Processes (HP) model self-excitation in an analytical expression for the intensity through the insertion of an extra term, which is designed to capture the effect of all the previous events of the process in the current value of the CIF. For a univariate Hawkes, we have:

$$\lambda_{HP}(t) = \mu + \sum_{t_i < t} \phi(t - t_i),$$  \hspace{1cm} (7)

while, for the multivariate case, with dimension $D$, we are going to have both self-excitation ($\phi_{ii}(t)$) and mutual-excitation terms ($\phi_{ij}(t)$, s.t. ($i \neq j$)):

$$\lambda_{HP}^i(t) = \mu_i + \sum_{j=1}^{D} \sum_{t_j < t} \phi_{ij}(t - t_{ij}).$$  \hspace{1cm} (8)

In the case that the kernel matrix

$$\Phi(t) = \begin{bmatrix} \phi_{11}(t) & \phi_{12}(t) & \phi_{13}(t) & \ldots & \phi_{1n}(t) \\ \phi_{21}(t) & \phi_{22}(t) & \phi_{23}(t) & \ldots & \phi_{2n}(t) \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ \phi_{d1}(t) & \phi_{d2}(t) & \phi_{d3}(t) & \ldots & \phi_{dn}(t) \end{bmatrix},$$  \hspace{1cm} (9)

can be factored into

$$\alpha = \begin{bmatrix} \alpha_{11} & \alpha_{12} & \alpha_{13} & \ldots & \alpha_{1n} \\ \alpha_{21} & \alpha_{22} & \alpha_{23} & \ldots & \alpha_{2n} \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ \alpha_{d1} & \alpha_{d2} & \alpha_{d3} & \ldots & \alpha_{dn} \end{bmatrix},$$  \hspace{1cm} (10)

with $\alpha \in \mathbb{R}$, and

$$\kappa(t) = \begin{bmatrix} \kappa_{11}(t) & \kappa_{12}(t) & \kappa_{13}(t) & \ldots & \kappa_{1n}(t) \\ \kappa_{21}(t) & \kappa_{22}(t) & \kappa_{23}(t) & \ldots & \kappa_{2n}(t) \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ \kappa_{d1}(t) & \kappa_{d2}(t) & \kappa_{d3}(t) & \ldots & \kappa_{dn}(t) \end{bmatrix},$$  \hspace{1cm} (11)

with

$$\Phi(t) = \alpha \odot \kappa(t),$$  \hspace{1cm} (12)

where “$\odot$” correspond to the Hadamard (element-wise) product. $\alpha$, here denominated Impact Matrix, can implicitly capture a myriad of different patterns of self- and mutual excitation, as exemplified in Figure 4.

Fig. 4: Examples of impact matrices.

For being of practical value, realizations of HPs are constrained to having finite number of events for any sub-interval of the simulation horizon $[0, T]$. This corresponds to having the kernel function (or kernel matrix) satisfying the following stationarity condition:

$$\left\| \int_0^\infty \Phi(t) dt \right\| = \left\| \left\{ \int_0^\infty \phi_{ij}(t) dt \right\}_{1 \leq i,j \leq D} \right\| \leq 1,$$  \hspace{1cm} (13)

where $\| \cdot \|$ correspond to the spectral radius of the matrix, i.e., the largest value among its eigenvalues.

Note that, for $D = 1$:

$$\| \Phi(t) \| = \int_0^\infty \phi(t) dt.$$  \hspace{1cm} (14)

If this stationarity condition is satisfied, we have that the process will reach wide-sense stationary state, i.e., when the properties of the process, most notably its “moments”, vary only as a function of the relative distance, here referred to as “$\tau$”, of its points.

The first order moment, or statistics of the HP, is defined as:

$$\Lambda_i = E\{ \lambda_{HP}^i(t) \}$$  \hspace{1cm} (15)

$$= \lim_{T \to \infty} \frac{1}{T} \int_0^\infty \lambda_{HP}^i(t) dt$$  \hspace{1cm} (16)

$$= (\| \Phi(t) \|)^{-1} \mu_i,$$  \hspace{1cm} (17)

while the second-order statistics, or stationary covariance, is defined as:

$$\nu_{ij}^2(t' - t) dt' dt'' = E\{ dN_i^j dN_j^i \} - \Lambda_i \Lambda_j dt dt''$$  \hspace{1cm} (18)

$$- \epsilon_{ij} \Lambda_i \delta(t' - t) dt,$$  \hspace{1cm} (19)

$$= \epsilon_{ij} \Lambda_i \delta(t - t') dt$$  \hspace{1cm} (20)

where $\epsilon_{ij}$ is 1, if $i = j$, and 0, otherwise, while $\delta(t)$ refers to the Dirac delta distribution.

The Fourier Transform (or, sometimes, the Laplace Transform) of this stationary covariance is referred to as Bartlett
Spectrum. In Hawkes’ seminal paper [13], high importance is given to the fact that, when assuming some specific parametric functions for the excitation matrix $\Phi(t)$, it is possible to find simple formulas for the covariance of process in the frequency domain. One example is the univariate case for $\phi_{ij}(t)$ given by a simple exponential:

$$
\phi(t) = \alpha e^{-\beta t},
$$

(21)

we have:

$$
\mu^*(s) = \mathcal{L}\{\nu(t)\} = \frac{\alpha \mu (2\beta - \alpha)}{2(\beta - \alpha)(s + \beta - \alpha)},
$$

(22)

where $\mathcal{L}\{\cdot\}$ refers to the Laplace Transform.

Going beyond the first- and second-order statistics, it is also possible to define statistics of higher orders, see [26], [8]. Although they become less and less intuitive and tractable, as their order increases, the work in [1] makes use of 3rd-order statistics, $K^{ijk}$, in a specific application of the Generalized Method of Moments for learning the impact matrix of multivariate HPs. It is defined as:

$$
K^{ijk} dt = \int \int_{\tau,\tau'} e^{i\kappa}\left( E(dN_i^t) E(dN_j^{\tau}) E(dN_k^{\tau + \tau'}),
$$

(23)

for $1 \leq i, j, k \leq D$, and it is connected to the skewness of $N_t$.

Now, regardless of the function family chosen for modeling $\Phi(t)$ and $\mu$, its fitness will be computed by measuring its likelihood over a set of sequences similar to the set of sequences used for training the model. The likelihood function given in the logarithmic form, i.e., the loglikelihood, of a multivariate HP over a set $S$ of $M$ sequences is given by:

$$
llh_S(\mu, \theta, \mathbb{F}) = \sum_{i=1}^{D} \sum_{j \in N_i} \log \lambda_{HP}(t_j^i) - \frac{1}{\mu} \int_0^T \sum_{i=1}^{D} \lambda_{HP}(t) dt
$$

(28)

Thus, the goal of learning a HP over $S$, is the act of, given a family $\mathbb{F}$ of functions assumed for the shape of the excitation functions (e.g., exponential, power-law, recurrent neural network), finding vectors $\mu$ and $\theta$ s.t.:

$$
(\mu, \theta) = \arg \max llh_S(\mu, \theta, \mathbb{F})
$$

(30)

Another concept which is of relevance in some inference methods is that of a Branching Structure. It defines the ancestry of each event in a given sequence, i.e., specifies if the $n$-th event $t_n$ was caused by a preceding event $t_i$ $(0 \leq i < n)$ or by the baseline intensity $\mu$. This concept is illustrated in Figure 5 in which the probabilities can be defined by

$$
p_{ji} = \frac{\phi(t_i - t_j)}{\lambda(t_i)}, \text{ for } (j \geq 1)
$$

(31)

and

$$
p_{0i} = \frac{\mu}{\lambda(t_i)}.
$$

(32)

C. Simulation Algorithms

Regarding the experimental aspect of HPs, synthetic data may be generated through the following methods:

1) Ogata’s Modified Thinning Algorithm [35]: It starts by sampling the first event at time $t_0$ from the baseline intensity. Then, each posterior event $t_i$ is obtained by sampling it from a HPP with intensity fixed as the value calculated at $t_{i-1}$, and then:

- Accepting it with probability $\frac{\lambda(t_i)}{\lambda^*(t_{i-1})}$, where $\lambda^*(t_{i-1})$ is the value of the intensity at time $t_{i-1}$, while $\lambda_{t_i}$ is the value calculated through Equation 7
- Or rejecting it and proceed to resampling a posterior event candidate.

2) Perfect Simulation [33]: It derives from the fact the HP may be seen as a superposition of Poisson Processes. It proceeds by sampling events from the baseline intensity, taken as the initial level, and then sampling levels of descendant events for each of the events sampled at the previous level. From each event $t_{0,i}$ sampled from the baseline intensity, we associate an IPP with intensity defined as $\phi(t - t_{0,i})$, and then sample its descendant events. Next, we take each descendant event sampled and also associate it with its corresponding IPP, and so on, until all the levels were explored over the simulation horizon $[0, T]$.

In the next section, we focus on the HP models which assume simple parametric forms for the excitation functions, along with its variants, which we will refer to as “Parametric HPs”.

![Fig. 5: Illustrative example of the concept of Branching Structure. A given edge $t_i \to t_j$ means that $t_i$ triggered $t_j$ with the probability $p_{ji}$.](image-url)
III. PARAMETRIC HPs

In the present section, we discuss the HP models which assume simple parametric forms for the excitation functions. Figure 4 shows some examples of commonly used functions for parametric HPs. Much has been done recently in terms of incrementing these models for dealing with specific aspects of some domains, such as social networks [65], [20], audio streaming [48], medical check-ups [55], among others.

As a way of modeling the daily oscillations of the triggering effects on Twitter data, [20] proposes a time-varying excitation function for HPs. The probability \( P \) of getting a retweet over the time interval \([t, t + \delta t]\), with small \( \delta \), is modeled as:

\[
P(Retweet in [t, t + \delta t]) = \lambda(t)\delta t,
\]

in which the time-dependent rate is dependent on previous events as:

\[
\lambda(t) = p(t) \sum_{t_i < t} d_i \phi(t - t_i),
\]

with \( p(t) \) being the infectiousness rate, \( t_i \) as the time corresponding to the i-th retweet arrival, and \( d_i \) as the number of followers of the i-th retweeting individual.

Furthermore, the memory kernel \( \phi(s) \), a probability distribution for the time intervals between a tweet by the follower and its retweet by the follower, has been shown to be heavily tailed in a variety of social networks [65]. It is fitted to the empirical data by the function:

\[
\phi(s) = \begin{cases} 
0, & \text{for } s < 0 \\
c_0, & \text{for } 0 \leq s \leq s_0 \\
c_0(s/s_0)^{(1+\theta)}, & \text{for } s > s_0
\end{cases}
\]

where the parameters \( c_0, s_0 \) and \( \theta \) are known.

The model is define so that the daily cycles of human activity are naturally translated into cycles of retweet activity. The time dependence of the infectious rate is, therefore, defined as:

\[
p(t) = p_0 \left( 1 - r_0 \sin \left( \frac{2\pi}{\tau_m} (t + \phi_0) \right) \right) e^{-\frac{(t - t_0)}{\tau_m}}
\]

The parameters \( p_0, r_0, \phi_0 \) and \( \tau_m \) correspond to the intensity, the relative amplitude of the oscillation, its phase, and the characteristic time of popularity decay respectively. Those are fitted through a Least Square Error (LSE) minimization procedure over the aggregation of retweet events over time bins \( \delta t \).

Another improvement over the traditional parametric forms for HPs involve the addition of a nonlinearity on the expression for the CIF:

\[
\lambda_{HP}(t) = g \left( \mu + \sum_{t_i < t} \phi(t - t_i) \right),
\]

in which \( g(\cdot) \) correspond to a so-called link function, e.g., sigmoid:

\[
g(x) = \frac{1}{1 + e^{-x}}.
\]

The work in [48] proposes a procedure for simultaneously learning \( g(\cdot), \mu \) and \( \phi(t) = \alpha \kappa(t), \) with \( \kappa(t) \) taken as \( e^{-t} \), for assuring convergence, of the algorithm.

The procedure is formulated using a moment-matching idea over a piecewise-constant approximation for \( g(\cdot) \), which leads to the definition of the objective function as a summation:

\[
\min_{\theta} \frac{1}{n} \sum_{i=1}^{n} \left( N_i - \int_{0}^{t_i} g(w \cdot x_i) dt \right)^2,
\]

with \( w = (\mu, \alpha)^T \), and \( x_t = (1, \sum_{i \in \mathcal{N}_t} \kappa'(t - t_i))^T \).

The algorithm is run by recursively updating the estimates \( \hat{\omega} \), with the Isotron Algorithm (see [18]), and \( \hat{\gamma} \), with a projected gradient descent step.

Theoretical bounds for the approximation error of the method are also given, along with extensions of the algorithm for general point processes, monotonically decreasing nonlinearities, low-rank processes and multidimensional HPs.

Another method of modeling the triggering kernel is through a composition of Gaussian kernels of different bandwidths, as in [54]. The core idea is that the maximum nonzero frequency component of the kernel is bounded as the same value of the intensity function, since this one is simply a weighted sum of the basis functions. Therefore, for every value of the tolerance \( \varepsilon \), it is possible to find a frequency value \( \omega_0 \) s.t.:

\[
\int_{\omega_0}^{\infty} |\hat{\lambda}(\omega)| d\omega \leq \varepsilon
\]

From this \( \omega_0 \), the method then defines the triggering function \( \phi(t) \) as a composition of \( K \) Gaussian functions, with \( K \) equally spaced values of bandwidth over the interval \([0, \omega_0]\).

The estimate of the value of the intensity function, for transforming into the frequency domain, is done through a kernel density estimation with gaussian kernels of bandwidth fixed as the Silverman’s rule of thumb:

\[
h = \left( \frac{4\sigma^5}{3n} \right)^{1/5} \approx 1.06\sigma n^{-1/5},
\]

where \( \sigma \) is the standard deviation of the time intervals, and \( n \) is the number of events of a given sequence.

After the Gaussian functions are defined, it remains to estimate the coefficients of the impact matrix, now a impact tensor \( \Theta \), through a convex surrogate loss penalized by parameters related to the sparsity of the matrix, the temporal sparsity of the kernels, and the pairwise similarity:

\[
\argmin_{\theta \geq 0} -\mathcal{L}_\Theta + \alpha_S \|A\|_1 + \alpha_G \|A\|_{1,2} + \alpha_F E(A),
\]

where:

* \( \|A\|_1 \) is the 11-norm of the tensor, which is related to its temporal sparsity, which causes the excitation functions to go to zero at infinity, therefore maintaining the stability of the process;
* \( \|A\|_{1,2} \) is related to the sparsity over the K basis functions of a given node of the process, and it enforces the local independence of the process;
E(A) is a coefficient to enforce the pairwise similarity of the process, which means that, if u and u’ are similar types of events, then \( \phi_u(t) \) and \( \phi_{u'}(t) \) should also be close to each other, as well as \( \phi_u(t) \) and \( \phi_{u'}(t) \);

\( \alpha_S, \alpha_G \) and \( \alpha_P \) are coefficients to be tuned for the model.

The estimation is done through an Expectation-Maximization procedure close to those of [32] and [66], which first randomly initializes the impact tensor and the vector of baseline intensities \( \mu \), and then iterates through:

1) Estimating the probability that each event was generated by each of the compositional basis kernels, as well as the baseline intensity;
2) Averaging the probabilities over all events of all training sequences for updating the coefficients of each basis function and the baseline intensity.

The two steps are repeated until convergence of the parameter estimates.

Another interesting way of learning multivariate marked HPs towards domain-specific applications, described in [53], is by assuming that the M generated sequences \( \{t_i, c_i\}_{i=1}^{I} \) \((t_i \in [0, T], c_i \in C = \{1, \ldots, C\})\), each one corresponding to an specific agent, are a product of two factors:

1) A underlying “registered” point process, defined with parameters \( \theta \), which would describe the common factors among all agents:

\[
\lambda(t) = \sum_{j=1}^{J} \exp_{t_j}(f_{j}(t; \theta, H^C_l)),
\]

for \( J \geq 1, \) \( f_{j} \) as linear functions of time, and \( H^C_l = \{(t_i, c_i)|t_i < t, c_i \in C\} \);

2) And sample-specific Warping functions \( \{W_m\}_{m=1}^{M} \), s.t. \( W_m : [0, T] \rightarrow [0, T] \),

and associated Unwarping functions \( \{W_m^{-1}\}_{m=1}^{M} \), which would capture the Idiosyncrasies of each sample/agent, with the following restrictions:

- \( E[W_m^{-1}(t)] = t; \)
- \( W_m^{-1}(t) \) is monotonically increasing on \([0, T] \).

As an example, the job-hopping among technology companies may have a core common aspect, but there are also traces of individuality (idioms) in the behaviour of employers of Amazon, as compared to those of Microsoft.

Learning both Intensity and Warping functions jointly is equivalent to the following optimization problem:

\[
\min_{\theta, \{W_m\}} \left(-\sum_{m=0}^{M} \log L(W_m^{-1}(S_m)) + \gamma R(\{W_m^{-1}\})\right),
\]

with the loglikelihood \( L(W_m^{-1}(S_m)) \) defined as:

\[
L(\theta, W_m^{-1}(S_m)) = \prod_{i=1}^{I_m} \lambda_{c_i}(W_m^{-1}(t_{im})),
\]

and the regularization term \( R(\{W_m^{-1}\}) \) as:

\[
R(\{W_m^{-1}\}) = \int_{0}^{T} \frac{1}{M} \sum_{m=1}^{M} W_m^{-1}(s) - s ds
\]

The optimization is done through recursively estimating the Unwarping function, and then using the estimate for each sequence to optimize for the registered process parameter \( \theta \), alternating until convergence.

The procedure is made feasible through two key assumptions:

- The intensity function is of exponential shape, from which the markovian property over the effects of History simplifies the \( \theta \)-learning step;
- The assumption of the Warping function as a piecewise-linear function between each consecutive pair of events:

\[
W_m^{-1}(t) = a_m^t + b_m^t, \text{ if } t \in [t_i, t_{i+1}),
\]

which allows to represent the Warping and Unwarping functions explicitly, while also turning the problem into a typical constrained nonlinear programming problem.

Another setting in which the parametric choice of kernels is highly convenient is with respect to the scalability of the inference procedure for high dimensional networks and sequences with large number of events, which occur in several domains, such as social interactions data, which is simultaneously large (i.e., large number of posts), high-dimensional (numerous users) and structured (i.e., the users interactions are not in a random fashion but, instead, present some regularities).

One interesting inference method towards this direction is the work presented in [22], which achieves a complexity \( O(nd) \), with \( n \) as the number of nodes, and \( d \) as the dimension of the impact matrix of the process.

The referred method, entitled Scalable Low-Rank Hawkes Processes (SLRHP), takes advantage of the memoryless
property of the exponential and the underlying regularity of large networks connected to social events: The memoryless property, which means that, in HPs with exponential excitation functions, the effect of all past events over the intensity value of a given point can be computed by just the time of the last event before said point, speeds up the intensity computing portion of the inference procedure iterations, while the underlying regularity of large impact matrices associated with the social phenomena allow the dynamics of large-dimensional HPs to be captured by impact matrices of much smaller magnitudes.

The baseline rates and excitation functions of model are then defined using a low-rank approximation:

\[ \mu_i(t) = \sum_{j=1}^{r} P_{ij} \bar{\mu}_j \]  
(48)

\[ \phi_{mi}(t) = \sum_{j,l=1}^{r} P_{ij} P_{ml} \bar{\phi}_{lj}(t), \]  
(49)

where \( P \in \mathbb{R}^{d \times r} \) is a projection matrix from the original d-dimensional space to a low-dimensional space \( r \) \( (r << d) \). This projection can also be seen as a low-rank approximation of the excitation function matrix \( \Phi \), in which:

\[ \Phi = P \bar{\Phi} P^T \]  
(50)

Through having that \( r << d \), the formulated low-rank approximated inference algorithm SLRHP manages to:

1) Capture a simplified underlying regularity impose inferred intensity rates’ parameters by adopting sparsity-inducing constraints to the model parameters;

2) Lower the number of parameters for both the baseline rates and excitation kernels, with the d natural rates and \( d^2 \) triggering kernels are lowered to \( r \) and \( r^2 \), respectively. This advantage is diminished slightly by the additional cost of inferring the \( (d \times r) \)-sized projection matrix \( P \).

Another way of dealing with scalability issues of multivariate HPs, both in terms of the total number of events in the sequences and the number of nodes, is through the mean-field treatment, as described in [4]. The key step of this method is to consider that the arrival intensities of each node of the process is wide-sense stationary, which implies the stability condition for the excitation matrix, and fluctuates only slightly around its mean value.

This last assumption, entitled \textit{Mean-Field Hypothesis}, posits that, if \( \lambda^i \) corresponds to the intensity of the \( i \)-th node, and \( \Lambda^i \) corresponds to the empirical estimator of the first-order statistics of said node,

\[ \tilde{\Lambda}^i = \frac{N_{T}^i}{T}, \]  
(51)

where \( N_{T}^i \) corresponds to the total number of events arrived at node \( i \) up to the final time of the simulation horizon \([0, T]\), then we have that:

\[ r^i = \frac{|\lambda^i - \tilde{\Lambda}^i|}{\Lambda^i} << 1. \]  
(52)

From this, we can recover the parameters \( \theta^i \) from the intensity function \( \lambda^i \), and the intensity function from the first-order statistics \( \Lambda^i \), as:

\[ \log \lambda^i \simeq \log \tilde{\Lambda}^i + \frac{\lambda^i - \tilde{\Lambda}^i}{\Lambda^i} - \frac{(\lambda^i - \tilde{\Lambda}^i)^2}{2(\Lambda^i)^2} \]  
(53)

and

\[ \lambda^i = \mu^i + \int_{0-}^{t} \phi^{ji}(t) dN^i_t \]  
(54)

The method yields mean-field estimates for the parameters with error which decays proportionally to the inverse of the final time \( T \) of the sequences:

\[ \mathbb{E}(\theta^i) \approx \theta^i_{MF} \]  
(55)

\[ \mathbb{C}(\theta^{ji}, \theta^{j'i}) \sim \frac{1}{T}, \]  
(56)

where \( \theta^i_{MF} \) refers to the mean-field estimator of the parameters.

Elaborating further on some difficulties of inferring parametric kernels from real-data, an interesting method regarding truncated sequences is described in [55].

In the case of learning HP parameters from real-data, one often has to deal with sequences which are only partially observed, i.e., the time event arrivals are only available over a finite time-window.

This poses a challenge concerning the robustness of the learning algorithms, since the triggering pattern from unobserved events are not considered: The inference deals with the error induced by computing intensity values over finite time windows, i.e., by computing intensity values along simulation horizons \([0, T]\), the closed-form equation would assume that its value at 0 is simply the baseline rate \( \mu \).

From the expression for the CIF:

\[ \lambda_{HP}(t) = \mu + \sum_{t_i < t} \phi(t - t_i), \]  
(57)

for \( t \in [0, T] \), we have that, taking some value \( T' \in [0, T] \), we may split the triggering effect term in two parts:

\[ \lambda_{HP}(t) = \mu + \sum_{t_i < T'} \phi(t - t_i) + \sum_{T' \leq t_i < T} \phi(t - t_i), \]  
(58)

for \( t \in [T', T] \).

If we are observing the sequences only over the interval \([T', T]\), the second term is implicitly ignored, what may have severe degradation of the learning procedure, specially in the case that the excitation functions decay slowly.

The method proposed handles this issue through a sequence-stitching method. The trick is to sample "candidate predecessor events" and choosing the most likely one from their similarity w.r.t. the observed events. The augmented sequences would then be used for the actual HP parameters’ learning algorithm.

In practice, this means that, given \( M \) sequences \( f_m \) realized over \([T', T]\), the method would not learn from the regular MLE formula

\[ \theta^* = \arg \max_{\theta} \mathcal{L}(\{f_m\}_{m=1}^{M}, \theta), \]  
(59)
but, instead, from some expression which takes into account the expected influence of unobserved predecessor events

\[ \theta_{SDC}^* = \arg\max_{\theta} \mathbb{E}_s \mathcal{H}_T \mathcal{L}([f, \hat{m}], \theta), \]  

(60)

where \( \mathcal{H}_T \) corresponds to the distribution over all possible sequences of events happening before time \( T' \). In practice, this expectation is computed not over the real distribution, but over some finite number of (relatively few) samples, such as 5 or 10. This finite sample approximation converts Equation 60 onto

\[ \theta_{SDC}^* = \arg\max_{\theta} \sum_{f_{\text{stitch}} \in \mathcal{K}} p(f_{\text{stitch}}) \mathcal{L}(f_{\text{stitch}}, \theta), \]  

(61)

where \( p(f_{\text{stitch}}) \) is the probability of the stitched sequence obtained from the concatenation of the original observed sequence and one of the sample predecessor candidate sequence. This probability is obtained by normalizing over similarity values over the candidate sequences obtained from some similarity function \( S(\cdot, \cdot) \) of the form:

\[ S(f_k, f) = e^{-\frac{|f(f_k) - f(f)|^2}{\sigma}} \]  

(62)

where \( f(\cdot) \) is some feature of the event sequence. By fixing the excitation pattern matrix \( \kappa \) as composed of exponentials \( \kappa(t) = e^{-\beta t} \) and imposing sparsity constraint \( R(\alpha) = ||\alpha||_1 \) over the impact matrix, the equivalent problem,

\[ \theta^* = \arg\max_{\mu \geq 0, \alpha > 0} \sum_{f_{\text{stitch}} \in \mathcal{K}} p(f_{\text{stitch}}) \mathcal{L}(f_{\text{stitch}}, \theta) + \gamma R(\alpha) \]  

(63)

is shown to be solved through Expectation-Maximization updated for both \( \mu \) and \( \alpha \). The method is more suitable for slowly-decaying excitation patterns, in which the influence of the unobserved events would be more prominent. In the case of exponentials with large values for the decay factor \( \beta \), the improvement margins mostly vanish.

One interesting improvement regarding the training procedure of MLE for HP parametric functions involves the complementary use of adversarial and discriminative learning, as in [57]. Although adversarial training has gained an ever increasing relevance for neural-network based models in the last years, due to the popularization of Generative Adversarial Networks [12] and its variants, as will be discussed further in Section \( \square \) keeping the assumption of a simple parametric shape for the excitation function is a way to insert domain-specific knowledge in the inference procedure.

The key idea of this complementary training is that, while the discriminative loss, here defined as the Mean-Squared Error (MSE) between discretized versions of predicted and real sequences, would direct the parameter updates towards smoother prediction curves, the adversarial loss would tend to push the temporally evenly distributed sampled sequences towards those more realistic-looking.

For the Gradient Descent-based updates, a discretization of the point process is carried out, so as to approximate the predictions through a recursive computation of the integral of the intensity function (the compensator portion of the loglikelihood function) in a closed-form expression. The parameters of the complementary training are actually initialized by a purely MLE procedure, which was found to be more insensitive to initial points. The \( MLE + GAN \) training updates then follows. The full procedure can be summarized by the following steps:

1) Subdivide the \( M \) original sequences on \([0, T]\) into training (on \([0, T^{tn}]\)), validation (on \([T^{tn}, T^{vd}]\)) and test (on \([T^{vd}, T]\)) portions, using previously defined parameters \( T^{tn} \) and \( T^{vd} \);
2) Initialize the parameters of the model through the ‘purely’ MLE procedure;
3) Sample \( M \) sequences from the model over the interval \([0, T^{tn}]\);
4) By choosing a specific parameter shape for the excitation function and binning both the original and simulated sequences over equally-spaced intervals, define the closed-form expression for the MSE (discriminative) loss \( L_{MSE} \) over all the sequences and dimensions of the process;
5) Define the GAN (adversarial) loss of the model over the sequences as:

\[ L_{GAN} = \begin{cases} 
E_{St^n \in \mathcal{P}(S^{tn})} [F_W(Y_{\theta}(S^{tn}))] - E_{St^n \in \mathcal{P}(S^{tn})} [F_W(Y_{\theta}(S^{tn}))], \\
- E_{St^n \in \mathcal{P}(S^{tn})} [F_W(Y_{\theta}(S^{tn}))],
\end{cases} \]  

for the critic network \( FW \) and for the training model \( G \),

(64)

where \( \mathcal{P}(S^{tn}) \) corresponds to the underlying probability distribution we assumed to have generated the original sequences \( S^{tn} \). \( F_W(\cdot) \) refers to a neural network which can compute the so-called Wasserstein distance, a metric for difference among distributions which will be further explained in Section \( \square \) and \( Y_{\theta}(\cdot) \) corresponds to the parametric model for sequence generation.

6) Compute the joint loss for MLE and GAN portions as:

\[ L_{MSE+GAN} = \delta L_{MSE} + (1 - \delta) L_{GAN}, \]  

(65)

7) Compute gradients of \( L_{MSE+GAN} \) over each parameter of model \( Y_{\theta} \);
8) Update parameter estimates of training model with \( \eta \nabla_{\theta}(L_{MSE+GAN}) \), for some learning rate \( \eta \);
9) Repeat steps 3 to 8 until convergence.

The method is an interesting combination of the enriched dynamical modeling from the adversarial training strategy with the robustness over small training sets of the parametric-based MLE estimation for HPs.

In this section, we provided a comprehensive analysis of the progresses in HP modeling and inference for excitation functions assumed to be of a simple parametric shape, along with their compositions and variants. In the next section, we will discuss about advances in nonparametric HP excitation function strategies.
IV. Nonparametric HPs

Nonparametric HPs consider that some rigid and simple parametric assumption for the triggering kernel may not be enough to capture all the subtleties of the excitation effects that could not be retrieved from the data. They may be broadly divided between two main approaches:

1) Frequentist
2) Bayesian
We will briefly discuss them in this section.

A. Frequentist Nonparametric HPs

The frequentist approach to HP modeling and inference consists in assuming that the excitation function (or matrix) can be defined as a binned grid (or a set of grids), in which the values of the functions would be taken as piecewise constant inside each bin, and the width of the bin will be (hopefully) expressive enough to model the local variations of the self-excitation effect.

They were first developed in the works of [23], [3] and [6]. In the case of [23], the final values of the bins were found by solving a discretized Ordinary Differential Equation, implied by the branching structure of the discretized triggering kernel and background rate, and the second-order statistics of the model, in the frequency domain, between the triggering kernel, the frequency domain, between the triggering kernel, the background rate and the second-order statistics of the model, also obtained in a discretized way over the data.

The increased expressiveness of this type of excitation model incurs in two main drawbacks:

- The bin division grid concept is close to that of a histogram over the distance among events, what usually requires much larger datasets for leading to accurate predictions, as opposed to parametric models, which would behave better on shorter and fewer sequences but most likely underfit on large sequence sets;
- The time of the inference procedure may also be much larger, since it involves sequential binning computation procedures which can not take advantage of the markov property of parametric functions such as the exponentials.

Two improvements, to be discussed in the present subsection, deal exactly with these drawbacks through:

- Acceleration of the computations over each sequence and/or over each bin of the excitation matrix/function;
- Reduction of the times of binning computational procedures through a so-called online update of the bin values.

One Acceleration strategy is developed in [1], which replaces the task of estimating the excitation functions directly through estimating their cumulative values, i.e., their integrated values from zero up to infinity, what would be enough to quantify the causal relationships among each node. That is, instead of estimating \( \phi_{ij}(t) \), for each node, the method would estimate a matrix \( G = \{g_{ij}\} \), in which:

\[
g_{ij} = \int_0^\infty \phi_{ij}(t) dt, \forall (i, j) \in D \times D
\]

The method, entitled Nonparametric Hawkes Process Cumulant (NPHC), then proceeds to compute, from the sequences, moment estimates \( \tilde{M} \) up to the third-order. It then finds some estimate \( \tilde{G} \) of this cumulant matrix which minimizes the \( L^2 \) squared error between these estimated moments and the actual moments \( M(G) \), which are uniquely determined from \( G \):

\[
\tilde{G} = \arg\min_G \|M(G) - \tilde{M}\|^2
\]

This \( L^2 \) minimization comes from the fact that, by defining:

\[
R = (I^D - G)^{-1},
\]

one may express the first-, second- and third-order moments of the process as:

\[
\Lambda^i = \sum_{m=1}^{D} R_{im} \mu^m
\]

\[
C^{ij} = \sum_{m=1}^{D} \Lambda^m R_{im} R_{jm}
\]

\[
K^{ijk} = \sum_{m=1}^{D} (R_{im} R_{jm} C^{km} + R_{im} C^{jm} R_{km} \pm C^{im} R_{im} R_{jm} R_{km}),
\]

and thus we may find an estimator \( \tilde{R} = \arg\min_R \mathcal{L}(R) \), with \( \mathcal{L}(R) \) defined as:

\[
\mathcal{L}(R) = (1 - \kappa) \|K^e - \tilde{K}^e\|_2^2 + \kappa \|C^e - \tilde{C}\|_2^2,
\]

where \( \kappa \) is a weighting parameter, \( \|\cdot\|_2 \) is the Frobenius norm and \( K^e = \{K^{ij}\}_{1 \leq i, j \leq D} \) is a two-dimensional compression of the tensor \( K \). From this expression, by setting

\[
\kappa = \frac{\|\tilde{K}^e\|_2^2}{\|\tilde{K}^e\|_2^2 + \|\tilde{C}\|_2^2},
\]

one may arrive to

\[
\tilde{G} = I^D - \tilde{R}^{-1}
\]

The estimates of moments in the algorithm are actually computed through truncated and discretized (binned) countings along a single realization of the process and, since the real-data estimates are usually not symmetric, the estimates are averaged along positive and negative axis.

Also, for \( D = 1 \), the estimate \( \tilde{G} \) can be estimated solely from the second-order statistics. For higher-dimensional processes, it is the skewness of the third-order moment which uniquely fixes \( \tilde{G} \).

Another improvement of the method consists in updating the parameters of the discretized estimate of the excitation function through a single pass over the event sequence, i.e., an online learning procedure [59].
In the case of the referred algorithm, the triggering function is assumed to:

1) be positive,
   \[ \phi(t) \geq 0, \forall t \in \mathbb{R} \]  
(76)

2) have a decreasing tail, i.e.,
   \[ \sum_{k=m}^{\infty} (t_k - t_{k-1}) \sup_{x \in (t_{k-1}, t_k]} |f(x)| \leq \epsilon_f(t_{m-1}), \forall m > 0, \]  
(77)

3) belong to a Reproducing Kernel Hilbert Space, which here is used as a tool for embedding similarity among high-dimensional and complex distributions onto lower-dimensional ones.

The method proceeds by taking the usual expression for the loglikelihood function

\[ llh_1(\lambda) = -\sum_{d=1}^{D} \left( \int_0^t \lambda_d(\tau) d\tau - x_{d,k} \log \lambda_d(t_k) \right) \]  
(78)

and optimizing, instead, over a discretized version of it

\[ llh_1(\lambda) = \sum_{d=1}^{D} \sum_{k=1}^{M(t)} \left( \int_{t_{k-1}}^{t_k} \lambda_d(\tau) d\tau - x_{d,k} \log \lambda_i(t_k) \right) \]  
(79)

\[ = \sum_{d=1}^{D} L_{d,t}(\lambda_d), \]  
(80)

with \( t_1, \ldots, t_{N(t)} \) denoting the event arrival times over an interval \([0, t]\), and with a partitioning \( \{0, t_1, \ldots, t_{M(t)}\} \) of this interval \([0, t]\) such that

\[ t_{k+1} = \min_{\tau_i \geq t_k} \{ t_k + \delta, \tau_i \} \]  
(81)

The discretized version can then be expressed as

\[ llh_1^{(\delta)}(\lambda) = \sum_{d=1}^{D} \sum_{k=1}^{M(t)} \left( \int_{t_{k-1}}^{t_k} (t_k - t_{k-1}) \lambda_d(t_k) \right) \]  
(82)

\[ - x_{d,k} \log \lambda_i(t_k) = \sum_{d=1}^{D} L_{d,t}^{(\delta)}(\lambda_d) \]  
(83)

The optimization procedure is done at each slot of the \( M(t) \) partition, taking into account:

- A truncation over the intensity function effect, i.e.,
  \[ \phi(t) = 0, \forall t > z, \]  
(84)

so as to simplify the optimization of the integral portion of the loss. The error over this approximation is shown to be bounded by the decreasing tail assumption.

- A Tikhonov regularization over the coefficients \( \mu_d \) and \( \phi_{d,t} \), which is simply the addition of weighted \( ||\mu_d||^2 \) and \( ||\phi_{d,t}||^2 \) terms to the loss function, so as to keep their resulting values small;

- A projection step for the triggering function optimization part, so as to keep them all positive.

The recent improvements over frequentist nonparametric HP estimation were shown to focus on two main strategies:

- Speeding up inference through replacement of the excitation matrix as objective by the matrix of cumulants, which were shown to be enough to capture the mutual influence among each pair of nodes;
- And an online learning procedure, which used some assumptions over the kernels (positive, decreasing tail RKHS) so as to recover estimates of the HP parameters over a single pass on some partitioning of the event arrival timeline.

**B. Bayesian Nonparametric HPs**

Another nonparametric treatment of HPs revolves around the assumption of the triggering kernel and the background rates to be modeled by distributions (or mixtures of distributions) from the so-called “Exponential Family”, which, through their conjugacy relationships, allow for closed-form computations of the sequential “Exponential Family”, which, through their conjugacy relationships, allow for closed-form computations of the sequential updates in the model. These were mainly proposed in the works of [11], [9] and [64].

In [11], HPs have also been used for modeling clustering of documents streams, capturing the dynamics of arrival time patterns, for being used together with textual content-based clustering.

The logic is that news and other media-related information sources revolve around a given occurrence- such as a natural catastrophe, a political action, or a celebrity scandal- are related not only regarding its word content, but also their time occurrences, as journalists tend to release more and more contents about a topic of high public interest, but tend to slow down the pace of publication as this interest gradually vanishes or shifts toward other subjects.

The main idea is to unite both Bayesian Nonparametric Inference, which is a scalable clustering method which allows for new clusters to be added as the number of samples grow, with Hawkes Processes. The corresponding Bayesian Nonparametric model, the Dirichlet Process, would capture the diversity of event types, while the HPs would capture the temporal dynamics of the event streams.

A Dirichlet Process \( DP(\alpha, G_0) \) can be roughly described as a probability distribution over probability distributions. It is defined by a concentration parameter \( \alpha \), proportional to the level of discretization (‘number of bins’) of the underlying sampled distribution, and a base distribution \( G_0 \), which is the distribution to be discretized. As an example, for \( \alpha \) equal to 0, the distribution is completely concentrated at a single value, while, in the limit that \( \alpha \) goes to infinity, the sampled distribution becomes continuous.

The corresponding hybrid model, the Dirichlet-Hawkes Process (DHP) is defined by:

- \( \mu \), an intensity parameter;
- \( G_0(\theta) \), a base distribution over a given parameter space \( \theta \in \Theta \);
- A collection of excitation functions \( \phi_{\theta}(t, t') \).

After an initial time event \( t_1 \) and an excitation function parameter \( \theta_1 \) are sampled from these base parameters \( \mu \), and
$G_0(\theta)$, respectively, the DHP is then allowed to alternate between:

1) Sampling new arrival events $t_i$ from the current value of $\theta$ for the excitation function, with probability

$$\mu + \sum_{i=1}^{n-1} \phi_\theta(t_n, t_i)$$

$$\text{(85)}$$

2) Or sampling a new value for $\theta$ with probability

$$\sum_{i=1}^{n-1} \phi_\theta(t_n, t_i)$$

$$\mu + \sum_{i=1}^{n-1} \phi_\theta(t_n, t_i)$$

$$\text{(86)}$$

In this way, you are dealing with a superposition of HPs, in which the arrival events tend towards processes with higher intensities, i.e., the preferential attachment, but which also allows for diversity, since there is always a nonzero probability of sampling a new HP from the baseline intensity $\mu$.

By defining the excitation functions $\phi_\theta$ as a summation of parametric kernels

$$\phi_\theta(t_i, t_j) = \sum_{i=1}^K \alpha_i^\theta \kappa_i(t_i - t_j)$$

$$\text{(87)}$$

the model can be made even more general.

The approach in [9], besides the random histogram assumption for the triggering kernel, similar to the frequentist case, also considers the case of it being defined by a mixture of Beta distributions, and have the model being updated through a sampling procedure (Markov chain Monte Carlo).

The work in [64] proposes a Gamma distribution over the possible values of $\mu$ and the triggering kernel to be modeled as:

$$f(\cdot) = \frac{f(\cdot)^2}{2},$$

$$\text{(88)}$$

where $f(\cdot)$ is a Gaussian Process [37].

These assumptions allow for closed-form updates over the posterior distributions over the background rate and the triggering kernel, which are claimed to be more scalable and efficient than the plain binning of the events.

In the next section, we will explore the neural architectures, which were introduced for modeling and generation of HPs through a more flexible representation of the effect of past events in the intensity function.

V. NEURAL NETWORK-BASED HPs

In this section, we discuss the neural network-based formulations of HP modeling. The main idea is to capture the influence of past events on the intensity function in a nonlinear, and thus hopefully more flexible, way.

This modeling approach makes use of recurrent models, which, in their simplest formulation, encode sequences of states

$$(x_0, x_1, ..., x_N)$$

$$\text{(89)}$$

and outputs

$$(y_0, y_1, ..., y_N)$$

$$\text{(90)}$$

in a way that each state $x_{i+1}$ can be obtained by a composition of the immediately preceding state $x_i$ and a so-called hidden state $h_i$ which captures the effect of the other past states:

$$h_i = \sigma_h(W_x x_i + W_h h_{i-1} + b_h)$$

$$\text{(91)}$$

$$y_i = \sigma_y(W_y h_i + b_y),$$

$$\text{(92)}$$

in which $W_y$, $W_x$, $W_h$, $b_h$ and $b_y$ are parameters to be fitted by the optimization procedure, while $\sigma_h$ and $\sigma_y$ are nonlinearities, such as a sigmoid or a hyperbolic tangent function.

In the case of HPs, the state to be modeled would be the intensity function along a sequence of time event arrivals, and an additional assumption would be that its intensity value decays exponentially between consecutive events.

In the case of most NN-based models, the inference also counts with a mark distribution for the case of marked HPs, in which a multinomial, or some other multi-class distribution, is fitted together with the recurrent intensity model.

Arguably the first from such type of models, the Recurrent Marked Temporal Point Process [10] jointly models marked event data by using a RNN with exponentiated output for modeling the intensity.

For sequences of the type $\{t_i, y_i\}_{i=1}^N$, in which $t_i$ corresponds to the time of the i-th event arrival, while $y_i$ refers to the type of event or mark, we have a hidden cell $h_i$ described by:

$$h_i = \max \left\{ W^y y_i + W^t t_i + W^h h_{i-1} + b_h, 0 \right\},$$

$$\text{(93)}$$

and a Conditional Intensity Function defined as a function of this hidden state

$$\lambda(t) = \exp \left( v^t h_i + w^t (t - t_i) + b^t \right),$$

$$\text{(94)}$$

while a K-sized mark set can have its probability modeled by a multinomial distribution:

$$P(y_{i+1} = k|h_i) = \frac{\exp (V^y_k h_i + b^y_k)}{\sum_{k=1}^K \exp (V^y_k h_i + b^y_k)}$$

$$\text{(95)}$$

As the likelihood of the whole sequence can be defined as a product of conditional density functions for each event:

$$\text{llh} \left( \{t_i, y_i\}_{i=1}^N \right) = \prod_{i=1}^N f(t_i, y_i),$$

$$\text{(96)}$$

with

$$f(t) = \lambda(t) \exp \left( -\int_{t_n}^t \lambda(t) dt \right)$$

$$\text{(97)}$$

$$= \exp \left( v^t h_j + w^t (t - t_i) + b^t + \frac{1}{w} \exp (v^t h_j + b^t) \right)$$

$$\text{(98)}$$

$$- \frac{1}{w} \exp (v^t h_i + w^t (t - t_i) + b^t) \right),$$

$$\text{(99)}$$

where $t_n$ is the latest event occurred before time $t$. From this $f(t)$, we may estimate the time of the next event as:

$$t_{i+1} = \int_{t_i}^{\infty} t f(t) dt$$

$$\text{(100)}$$
This allows us to optimize the parameters of the RNN model over the loss equal to this likelihood function, composed by these conditional density functions.

Given a set of M training sequences \( S_j = \{t_i^j, y_i^j\}_{i=1}^{N_j} \), we want to optimize the weight parameters over a loss function defined as:

\[
L(S_j) = \sum_j \sum_i \log \left( P(y_i^j|\mathbf{h}_i) + \log f(d_{i+1}^j|\mathbf{h}_i) \right)
\]

(101)

This optimization procedure is usually done through the Backpropagation Through Time (BPTT) algorithm, which proceeds by ‘unrolling’ the RNN cells by a fixed number of steps, then calculating the cumulative loss along all these steps, together with the gradients over each of W’s, V’s and b’s, then updating these parameters with a pre-defined learning rate until convergence.

One improvement for the RNN-based modeling approach is described in [51], and consists of treating the mark sequences as derived from another RNN model, instead of the multinomial distribution. This RNN for the marks is then jointly trained with the RNN for event arrival times.

Another improvement over this NN-based modeling approach is the Neural Hawkes Process [29], which uses a variant of the basic RNN, entitled Long Short-Term Memory (LSTM) [17], applied to the intensity function modeling.

The Neural Hawkes Process models the intensity function of a multi-type event sequence by associating each k-th event type with a corresponding intensity function \( \lambda_k(t) \), s.t.:

\[
\lambda_k = f_k(w_k^T \mathbf{h}(t)),
\]

(102)

with

\[
\mathbf{h}(t) = \mathbf{o}_t \odot (2\sigma(2\mathbf{c}(t) - 1)),
\]

(103)

The variables \( \mathbf{o}_t \) and \( \mathbf{c}(t) \) are defined through the following update rules:

\[
c_{i+1} = f_{i+1} \odot c(t_i) + i_{i+1} \odot z_{i+1},
\]

(104)

\[
i_{i+1} = \sigma(W_f k_i + U_i h(t_i) + d_i),
\]

(105)

\[
f_{i+1} = \sigma(W_f k_i + U_f h(t_i) + d_f),
\]

(106)

\[
z_{i+1} = 2\sigma(W_z k_i + U_z h(t_i) + d_z) - 1,
\]

(107)

\[
\mathbf{o}_{i+1} = \sigma(W_o k_i + U_o h(t_i) + d_o),
\]

(108)

and the value of the \( \mathbf{c}(t) \) is assumed to decay exponentially among consecutive events as:

\[
c(t) = c_{i+1} + (c_{i+1} - c_{i+1}) \exp(-\delta_{i+1}(t-t_i)),
\]

(109)

for

\[
c_{i+1} = c_{i+1} \odot c(t_i) + \tilde{i}_{i+1} \odot z_{i+1}
\]

(110)

\[
\delta_{i+1} = \sigma(W_\delta k_i + U_\delta h(t_i) + d_\delta).
\]

(111)

The W’s, U’s and d’s of the model are trained so as to maximize the log-likelihood over a set of sequences. Compared with the previous RNN-based model, in the Neural Hawkes Process:

1) The baseline intensity \( \mu_k \) is not implicitly considered constant, but instead is allowed to vary;
2) The variations of the cell intensity are not necessarily monotonic, because the influences of each event type on the cell values may decay at a different rate;
3) The sigmoid functions along the composition equations allow for an enriched behaviour of the intensity values.

All this contributes to an increased expressiveness of the model. Besides, as in the regular LSTM models, the “forget” gates \( f_{i+1} \) are trained so as to control how much influence the past values of \( \mathbf{c}(t) \) will have on its present value, thus allowing the model to possess a "long-term" memory.

Another variant of the RNN-based HPs, introduced in [52], models the baseline rate and the history influence as separate RNNs each. The baseline rate is taken as a time series, with its corresponding RNN updating its state at equally spaced intervals, such as five days. The event history influence RNN updates its state at each event arrival.

Both the background rate time series \( \{y_t^i\}_{t=1}^{T} \) and the marked event sequence \( \{m_i, t_i\}_{i=1}^{N} \) are modeled by LSTM cells:

\[
(h_t^y, c_t^y) = \text{LSTM}_y(y_t, h_{t-1}^y + c_{t-1}^y),
\]

(112)

\[
(h_t^m, c_t^m) = \text{LSTM}_m(m_t, h_{t-1}^m + c_{t-1}^m),
\]

(113)

These \( h \) and \( c \) states correspond to the hidden state and the long-term dependency terms, respectively, similarly to the Neural Hawkes Process. Both terms are concatenated in a single variable \( e_t \), for jointly training both RNN models:

\[
e_t = \tanh(W_f [h_t^y, h_t^c] + b_f)
\]

(114)

\[
U_t = \text{softMax}(W_U e_t + b_U),
\]

(115)

\[
u_t = \text{softMax}(W_u [e_t, U_t] + b_u)
\]

(116)

\[
s_t = W_s e_t + b_s,
\]

(117)

with \( U \) and \( u \) denoting the main event types and subtypes, respectively, and \( s \) denoting the composed timestamp of each event. The loss over which the model is trained is defined in a cross-entropy way:

\[
\sum_{j=1}^{N} \left( -W_U^j \log(u_t^j) - W_u^j \log(u_t^j) - \log \left( f(s_t^j|h_{t-1}^j) \right) \right),
\]

(118)

with

\[
f(s_t^j|h_{t-1}^j) = \frac{1}{\sqrt{2\pi\sigma}} e^{-\frac{(s_t^j - \hat{s}_t^j)^2}{2\sigma^2}},
\]

(119)

where \( \hat{s}_t^j \) is the model predicted output for the corresponding event \( s_t^j \).

The model weights are then jointly trained, over the total loss function and under some correction for the frequency ratio of each event type, for both the background rate time series values and the event arrivals RNN, and are shown to outperform more ‘rigid’ models.

Now, regarding the generation of HP sequences, both RMTPP and Neural Hawkes Processes have in common
the fact that they intend to model the intensity function of underlying process, so that new sequences may be sampled in a way to reproduce the behaviour of the original dataset. This intensity modeling, however, has three main drawbacks:

1) It may be unnecessary, since the sequences may be simply produced by unrolling cells of corresponding RNN models;

2) The sequences from these intensity-modeling approaches are sampled using a ‘Thinning algorithm’ [35], which may incur in slowed-down simulations, in the case of repeatedly rejected event intervals.

3) These methods are trained by maximizing the loglikelihood over the training sequences, which is asymptotically equivalent to minimizing the KL-Divergence over original and model distributions. This MLE approach is not robust in the case of multimodal distributions.

The model in [49] proposes approximating a generative model for generating event sequences by using an alternative metric of difference among distributions, the Wasserstein (or Earth-Moving) distance, already discussed in Section III.

In the model, entitled Wasserstein Generative Adversarial Temporal Point Process (WGANTPP), this Wasserstein loss is shown to be equal to:

\[
L' = \left[ \frac{1}{m} \sum_{i=1}^{m} f_w(g_\theta(z^{(i)})) - \frac{1}{m} \sum_{i=1}^{m} f_w(\xi) \right] + \nu \sum_{i,j=1}^{m} \frac{|f_w(\xi_i) - f_w(g_\theta(\xi_j))|}{|\xi_i - g_\theta(\xi_j)|},
\]

where the second term, along with the constant \(\nu\) corresponds to the so-called Lipschitz constraints, related to the continuity of the models.

\(\{\xi_i\}_{i=1}^{m}\) are real data sequences, while \(\{z^{(i)}\}_{i=1}^{n}\) are sequences sampled from a Homogeneous Poisson Process with a rate \(\lambda_z\) which is simply the expected arrival rate over all training sequences. The generator network \(g_\theta\) and discriminator \(f_w\) are defined as:

\[
g_\theta(z) = \rho = \{t_1, ..., t_n\}\]

\[
t_i = \eta_d^w(B_d^z h_i + b_d^z)
\]

\[
h_i = \eta_d^h(A_g^z t_i + B_g^h h_{i-1} + b_g^h)
\]

\[
f_w(\rho) = \rho = \sum_{i=1}^{n} a_i
\]

\[
a_i = \eta_d^a(B_d^a h_i + b_d^a)
\]

\[
h_i = \eta_d^h(A_g^h t_i + B_g^h h_{i-1} + b_g^h),
\]

with the \(\eta\)'s defined as nonlinearities. The \(A\)'s, \(B\)'s and \(b\)'s here are the parameters to be tuned by a Stochastic Gradient Descent procedure.

The distance metric \(|\cdot|_1\), for the case of purely temporal point processes in \([0, T]\), can be shown to be equivalent to

\[
\sum_{i=1}^{n} |t_i - \tau_i| + (m - n) \times T - \sum_{i=n+1}^{m} \tau_i,
\]

which has an intuitive graphical interpretation.

As previously discussed, the generator is trained so as to “fool” the discriminator, while the discriminator is trained so as to distinguish generated sequences from those of real data. This adversarial training procedure is roughly equivalent to gradient updates with opposite signs over their respective parameters: positive sign for the discriminator and negative sign for the generator.

In the end of the training procedure, one hopefully gets a generator network capable of producing sequences virtually indistinguishable from the real data ones.

This method, however, consists of training a network for generating entire sequences, and so the generator model learned may not accurately generate conditional output sequences from input ones. Another model, described in [50], deals with this task by generating, from partially observed sequences, the future events of these same sequences conditioned on their history, i.e., instead of aiming to capture the underlying distribution of a set of full sequences, the model performs a “sample agnostic” in-sample prediction.

Analogously to one of the parametric models described in Section III the learning procedure of this in-sample neural-network based prediction model takes advantages of both types of divergence measures: MLE loss (or KL Divergence) and Wasserstein distance.

The former aims for a rigid and unbiased parameter matching among two given probabilistic distributions, which is sensitive to noisy samples and outliers, while the latter has biased parameter updates, but is sensitive to underlying geometrical discrepancies among sample distributions. This combined loss is a way to balance both sets of priorities. In the case of long-term predictions, in which initial prediction errors propagate and magnify themselves throughout the whole stream, this joint loss was found to strengthen the effectiveness of the inference procedure.

The proposed model borrows on the seq2seq architecture ([43]) and aims to model endings of individual sequences conditioned on their partially observed history of initial events, and inserts an adversarial component in the training to increase the accuracy of long-term predictions. A network, designated as generator, encodes a compact representation of the initial partial observation of the sequence and outputs a decoded remaining of this same sequence. That is to say that, for a full sequence:

\[
\{t_1, t_2, ..., t_{n+m}\},
\]

the seq2seq modeling approach learns a mapping

\[
g_\theta(\xi) = \rho
\]

such that

\[
\xi = \{t_1, t_2, ..., t_n\}
\]

and

\[
\rho = \{t_{n+1}, t_{n+2}, ..., t_{n+m}\}
\]

This mapping is defined through a composition of LSTM cells:

\[
h_i = \eta_d^h(A_g^h t_i + B_g^h h_{i-1} + b_g^h)
\]
\[
t_{i+1} = \eta_d^h(B_d^g h_i + b_d^g),
\]
with \(\eta_d^h\) and \(\eta_d^g\) defined as nonlinear activation functions.

The learning procedure consists of finding values for A’s, B’s and b’s which maximize the conditional probability
\[
P(\rho | \xi) = \prod_{i=1}^{n+m-1} P(t_{i+1}|h_i, t_1, ..., t_i),
\]
what is carried through by parameter gradient updates over the combined loss Wasserstein and MLE loss. The adversarial component of the training, the discriminator \(f_w(\cdot)\), is modeled as a residual convolutional network (see [15]) with a 1-Lipschitz constraint, which is related to the magnitude of the gradients of the discriminative model. The full optimization problem then becomes
\[
\min_{\delta} \max_w \sum_{l=1}^{M} f_w(\{\xi_l, \rho_l\}) - \sum_{l=1}^{M} f_w(\{\xi_l, g_\theta(\xi_l)\}) - \lambda [f_w(\hat{\xi}) - 1] - \alpha \log(P_\theta(\rho | \xi))
\]
where \(\lambda\) is the Wasserstein loss and \(\alpha\) is the MLE loss.

Another improvement for NN-based modeling, proposed in [62], involves a so-called self-attention strategy [46] to improve the accuracy of the resulting network. The i-th event tuple \((t_i, m_i)\) is embedded as a variable \(x_i\):
\[
x_i = t_p + pe(m_i, t_i),
\]
which simultaneously encodes information about the event mark through
\[
t_p = e_m W_E,
\]
with \(e_m\) as an one-hot encoding vector of the mark and \(W_E\) as an embedding matrix, and information about the time interval among consecutive events through a sinusoidal-based positional encoding vector \(pe(m_i, t_i)\), with its k-th entry defined as:
\[
pe_k(m_i, t_i) = \sin(\omega_k \times i + w_k \times t_i)
\]
From this encoded variable \(x_i\), a hidden state \(h_{u,i}\) is then defined for each category \(u\) of the marks, which captures the influence of all previous events:
\[
h_{u,i+1} = \frac{\sum_{j=1}^{i} f(x_{i+1}, x_j) g(x_j)}{\sum_{j=1}^{i} f(x_{i+1}, x_j)}
\]
Through a series of nonlinear transformations, the intensity \(\lambda_u(t)\) for the u-th mark is then computed.

A further improvement of neural HP models, described in [41], involves the graph properties of multivariate HPs, which may be embedded in a NN modeling framework through the recently proposed Graph Convolutional Networks (GCN)[19].

The method is composed of a GCN module, for capturing meaningful correlation patterns in a large sets of event sequence, followed by an usual RNN module, for modeling the temporal dynamics. In short, the time sequences are modeled as a HPs, and the adjacencies among different processes are encoded as a graph. The novel (GCN+RNN) model is meant to extract significant local patterns from the graph.

The output of the initial GCN network module, which is simply a matrix of the form \(\chi = [\mu, A]\), which includes the baseline vector \(\mu\) and the adjacency matrix \(A\), are fed into the RNN-based module, there taken as a LSTM. Then, the output of this RNN module are input to a further module, a fully connected layer, for calculating the changes \(d\chi\) to be applied to the current parameter matrix \(\chi\). Then, after each training step T, the predicted value of this parameter matrix becomes as
\[
\chi(T) = \chi(T - 1) + d\chi(T - 1)
\]

The work in [58] proposes a model for check-in time prediction which is composed by a LSTM-based module, in which the feature vector is composed so as to capture each relevant aspect of the problem: the event time coordinate \(t_i\), an additional field to indicate if the check-in occurred on a weekday or during the weekend, the euclidean distance between a given check-in location and the location (l) of the previous check-in, the location type of the check-in (e.g., Hotel, Restaurant, etc.), the number of users overlapping with a given location, and the check-ins by friends of the user. This aggregates social, geographical and temporal information in a single NN-based HP-like predictive model.

All these variants of neural-network point process models allow for more flexible (nonlinear) representation of the effect of past events in the future ones, besides putting at the inference procedure’s disposal a myriad of Deep Learning tools and techniques which have enjoyed a surge of popularity over the recent years.

VI. FURTHER APPROACHES

In this section, we briefly review some recently proposed approaches which do not fit conveniently into any of the three previously discussed subgroups, but may be considered as bridgings between usual HP-related tasks and other mathematical subfields.

A. Sparse Gaussian Processes

By building over the modeling in [64], which considers the triggering function of the HP as a Gaussian Process, the work in [63] proposes an approach involving Sparse Gaussian Processes for optimizing over a dataset. In this, the optimization of the likelihood is being taken not over the samples, but over a set of much fewer so-called inducing points, which are also taken as latent variables, so as to result in a final model which is both expressive enough to capture the complexity of the dataset but also tractable enough to be useful and applicable to reasonably-sized datasets.

B. Stochastic Differential Equation

Another way of modeling HPs is through a Stochastic Differential Equation, as proposed in [21]. In them, the decay of
the underlying process is expected to generate as
sure, which is defined as the total number of events the
Forests, to obtain an associated so-called “Popularity” mea-
with other Machine Learning techniques, such as Random

E. Popularity Prediction

N
N

modulated by the size of the available population, as below:
a disease by introducing a HP intensity function which is
a time event as an infection, it models the diffusion of
traditional epidemic models over populations. By considering
D. Epidemic HPs

The work in [39] blends the HP excitation effect with
time event as an infection, it models the diffusion of
a disease by introducing a HP intensity function which is
modulated by the size of the available population, as below:

\[ \lambda(t) = \left(1 - \frac{N_t}{N}\right)\{\mu + \sum_{t_i<t} \phi(t-t_i)\}, \]

where \(N_t\) denotes the counting process associated with the
HP, while \(N\) is the total finite population size.

E. Popularity Prediction

The work in [31] proposes the use of HP modeling blended
with other Machine Learning techniques, such as Random
Forests, to obtain an associated so-called “Popularity” mea-
sure, which is defined as the total number of events the
underlying process is expected to generate as \(t \to \infty\).
This measure is treated as an outcome derived from features
associated with some entity (e.g., social network user), s.a.
number of friends, total number of posted statuses and the
account creation time.

In the next section, we will discuss models in which
one not only wants to capture the temporal dynamics, but
also wishes to influence it towards a certain goal, implicitly
defined through a so-called reward function.

VII. STOCHASTIC CONTROL, REINFORCEMENT
LEARNING AND IMITATION LEARNING OF HPs

In this section, we briefly review some control strategies
regarding HPs. In some cases, one may wish not only to
be able to model the traces of some event sequences, or
to capture the underlying distribution of said sequences, but
also try to influence their temporal dynamics towards more
advantageous ones.

This concept of advantageous is explicit through the
definition of a so-called Reward Function, which is defined
in terms of specific , and sometimes application-specific,
properties of the sequences. Most works related to the subject
deal with Social Network applications, and one example of
Reward can be the total time the post of a user stays at the top
of the feed of his/her followers. One type of reward which
is not domain-specific is the dissimilarity among two sets
of sequences, computed through mappings such as “kernel
mean embeddings” [34].

One example of this control approach is described in [61],
in which the variable to be controlled is the times to post of
a given user, implicitly defined as an intensity function, so
as to maximize the reward function, here computed as the
total time that this user’s posts stay at the top of the feed of
his/her followers.

This “when-to-post” problem can be formulated as:

\[
\min_{u(t_0,t_f)} \mathbb{E}_{(N_i,M_i)(t_0,t_f)} \left[ \phi(r(t_f)) + \int_{t_0}^{t_f} l(r(\tau), u(\tau)) d\tau \right]
\]

subject to \(u(t) \geq 0, \forall t \in [t_0, t_f]\),

(143)

where:
• \(i\) is the index of the broadcaster of the posts;
• \(N_i(t)\) is the Counting Process of the \(i\)-th broadcaster,
  with \(N(t) = \{N_i(t)\}_{i=1}^n\) being an array of Counting
Processes along all the \(n\) users of the network;
• \(A \in \{0,1\}^{n \times n}\) is the Adjacency Matrix of the network;
• \(M_i(t) = A^T N(t) - A_i N_i(t),\) which means that \(M_i(t)
  is the sum of the Counting Process of all users con-
nected to user \(i\) excluding user \(i\) him/herself;
• \(t_0\) and \(t_f\) are, respectively, the starting and ending times
  of the problem horizon taken in consideration;
• \(u(t) = \mu_i(t),\) the controlled variable, is the baseline in-
tensity of user \(i,\) to be steered towards the maximization
of the reward function;
• \(\phi(r(t_f))\) is an arbitrarily defined penalty function;
• \(\mathbb{E}_{(N,M)(t,t_f)}\) is a nondecreasing convex loss function
  defined w.r.t. the visibility of the broadcaster’s posts in
each of his/her followers’ feeds.

The approach used in the problem is by defining an optimal
cost-to-go \(J(r(t_f), \lambda(t), t)\):

\[
J(r(t_f), \lambda(t), t) =
\min_{u(t,t_f)} \mathbb{E}_{(N,M)(t,t_f)} \left[ \phi(r(t_f)) + \int_{t}^{t_f} \mathcal{L}(r(\tau), u(\tau)) d\tau \right],
\]

(145)
and find the optimal solution through the Bellman’s Principle of Optimality:

\[
J(r(t), \lambda(t), t) = \min_{u(t,t+dt)} \{ E[J(r(t + dt), \lambda(t + dt), t + dt)] + \frac{1}{2} \| \dot{r}(t), u(t) \| \}
\]

For example, in the case of a broadcaster with one follower \((r(t) = r(t))\), if the penalty and loss functions are defined as:

\[
\phi(r(t)) = \frac{1}{2} r^2(t_f)
\]

and

\[
\frac{1}{2} s(t) r^2(t) + \frac{1}{2} q u^2(t),
\]

for some positive significance function \(s(t)\) and some trade-off parameter \(q\), which calibrates the importance of both visibility and number of posts, we set the derivative of \(J(r(t), \lambda(t), t)\) over \(u(t)\) to 0 and solve it to get to an analytical solution

\[
u^*(t) = \frac{q}{r^2}\left[ J(r(t), \lambda(t), t) - J(0, \lambda(t), t) \right],
\]

which is thus the optimal intensity a broadcaster must adopt to maximize visibility, constrained on the cost associated to the number of posts, along this one follower’s feed. Further derivations are provided to the more natural and general case, in which the broadcaster may have multiple followers.

An earlier version of this type of Stochastic Optimal Control-based approach to influence activity in social networks can be found in [60]. In this, the goal is to maximize the total number of actions (or events) in the network. Analogously to the previously discussed algorithm, one may solve the Continuous Time version of the Bellman Equation through defining a optimal cost-to-go \(J(\lambda(t), t)\), which here depends only on the intensities of the nodes and the time.

The control input vector \(u(t)\) acts on the network by increasing the original vector of uncontrolled intensities

\[
\lambda(t) = \mu_0 + A \int_0^t \kappa(t - s) dN(s)
\]

with the equivalent rates of an underlying Counting Process vector \(dM(s)\), such that the new controlled intensity vector \(\lambda^*(t)\) is now described by

\[
\lambda^*(t) = \mu_0 + A \int_0^t \kappa(t - s) dN(s) + A \int_0^t \kappa(t - s) dM(s),
\]

where \(\kappa(t) = e^{-\omega t}\) in the model.

Then, in the same way, by differentiating the equivalent \(J(\lambda(t), t)\) over the control input, setting the corresponding expression to 0, then defining

\[
\phi(\lambda(t)) = -\frac{1}{2} \lambda^T(t) Q \lambda(t) + \frac{1}{2} u^T(t) S u(t)
\]

and

\[
\frac{1}{2} \dot{\lambda}(t_f) = -\frac{1}{2} \lambda^T(t_f) F \lambda(t_f),
\]

with previously defined symmetric weighting matrices \(Q, F\) and \(S\), we arrive to a closed-form expression for the optimal control intensity value

\[
u^*(t) = -S^{-1} \left[ A^T g(t) + A^T H(t) \lambda(t) + \frac{1}{2} \text{diag}(A^T H(t) A) \right],
\]

where \(H(t)\) and \(g(t)\) can be computed by solving the differential equations

\[
\dot{H}(t) = (\omega I - A)^T H(t) + \text{H}(\omega I - A) + H(t) A S^{-1} A^T H(t) Q
\]

\[
\dot{g}(t) = \left[ \omega I - A^T + H(t) A S^{-1} A^T \right] g(t) - \omega H(t) \mu_0 + \frac{1}{2} \text{diag}(H(t) A S^{-1} - I) \text{diag}(A^T H(t) A),
\]

with final conditions \(g(t_f) = 0\) and \(H(t_f) = -F\). The solution is constant between two consecutive events and gets recomputed at each event arrival.

These SOC-based approaches have two main drawbacks:

- The functional forms of the intensities and mark distributions are constrained to be from a very restricted class, which does not include the state-of-the-art RNN-based HP models;
- The objective function being optimized is also restricted to very specific classes of functions, so as to keep the tractability of the problem.

For circumventing these drawbacks, there have been some proposed approaches which combine more flexible and expressive HP models with robust stochastic optimization procedures independent from the functional form of the objective function.

On of these methods, entitled "Deep Reinforcement Learning of Marked Temporal Point Processes" [45], works by, given a set of possible actions and corresponding feedbacks, which are both expressed as temporal point processes jointly modeled by a RNN-based intensity model \(\lambda^*_\theta(t)\):

\[
\lambda^*_\theta(t) = \exp(b_\theta + w_i(t - t_i) + V_\theta h_i),
\]

with

\[
h_i = \text{tanh}(W_i h_{i-1} + W_1 \tau_i + W_2 y_i + W_3 z_i W_4 b_i + b_h),
\]

where

\[
\tau_i = W_i (t_i - t_i-1) + b_t,
\]

\[
b_i = W_s (1 - b_i) + W_f b_i + b_b,
\]

\[
y_i = W y_i + b_y, \text{ if } b_i = 0
\]

\[
z_i = W z_i + b_z, \text{ if } b_i = 1.
\]

The term \(b_i\) is an indicator function to whether the \(i\)-th event is an action or a feedback. By taking all the \(W\)’s and \(b\)’s as a parameter vector \(\theta\), what the algorithm wishes to do is to update this vector with the gradients of each parameter over an expected Reward Function \(J(\theta)\):

\[
\theta_{t+1} = \theta_t + \alpha_t \nabla_\theta J(\theta)|_{\theta=\theta_t}
\]
\[ \nabla_{\theta} J(\theta) = E_{A_T} p_{A,\theta}(\cdot|A_T) p_{F,\theta}(\cdot) [R^*(T) \nabla_{\theta} \log \mathbb{P}_\theta(A_T)], \]

(159)

where

\[ p_{A,\theta} = (\lambda_\theta^*, m_\theta^*) \]

(160)

is the joint conditional intensity and mark distribution for action events, and

\[ p_{F,\phi} = (\lambda_\phi^*, m_\phi^*) \]

(161)

is the joint conditional intensity and mark distribution for feedback events. The Reward \( R(T) \) is defined over some domain-specific metric, which may involve the responsiveness of followers in a social network setting, or the effectiveness of memorization of words in a foreign language, in a spaced-repetition learning setting.

Another way of using these reinforcement learning approach is through a technique entitled "Imitation Learning" [24]. The reasoning behind it is to treat the real-data sequences as generated by an expert and then, using RNN-based sequence generation, try to make these models approximate the real-data sequence as closely as possible.

Thus, the reward function to dictate the proportion in which the gradient of the parameters over each sequence are going to be considered is equal to how much this given sequence is likely to be drawn from the underlying distribution over the real data. This similarity is computed through a Reproducing Kernel Hilbert Space (RKHS).

The theory of RKHS is very extensive, and it is not our goal to give a detailed account of it here. The key idea is that, to compute similarities among items of a given space, you compute inner products between them. Taking two items, \( x_1 \) and \( x_2 \), and computing a so-called Positive Definite Kernel (PDS) \( K(x_1, x_2) \) is equivalent to computing a inner product among these two items in a high-dimensional, and potentially infinite-dimensional, vector space. The PDS used in the corresponding paper is the Gaussian kernel.

The reward function is then defined as:

\[ \hat{\tilde{p}}^*(t) \propto \frac{1}{L} \sum_{i=1}^{L} \sum_{l=1}^{N(l)} \kappa(t_i^{(l)}, l) - \frac{1}{M} \sum_{m=1}^{M} \sum_{l=1}^{N(m)} \kappa(t_i^{(m)}, l), \]

(162)

where:

- \( L \) is the number of expert trajectories;
- \( M \) is the number of trajectories generated by the model;
- \( \kappa(\cdot, \cdot) \) is the reproducing kernel operator;
- \( t_i^{(l)} \) is the i-th time coordinate of the l-th expert trajectory;
- \( t_i^{(m)} \) is the i-th time coordinate of the m-th model-generated trajectory.

The parameters of the model are then updated through a Gradient Descent-based approach towards convergence, in which the model is expected to generate sequences indistinguishable from the real-world process.

VIII. REAL-WORLD DATA LIMITATIONS

One key aspect of HP modeling which inevitably encompasses all the previously mentioned approaches is that of their applicability to real-world datasets. These may present a series of systematic issues on the training and testing sequences, which may completely hinder the generalization of the models. We discuss some key issues here, along with some recently proposed methods, designed to handle each of them.

A. Synchronization Noise

Temporal data, specially multivariate ones, may have their event streams being extracted from distributed sensor networks. A key challenge regarding them is that of synchronization noise, i.e., when each source is subject to an unknown and random time delay.

In these cases, an inference procedure which neglects these time delays may be ignoring critical causal effects of some events over others, thus resulting in a poorly generalizing model. The work in [44] deals specifically with this aspect of real-data HP modeling, and proposes, for the exponential triggering functions HPs, including the random time shift vector (one entry for each distinct event stream) as a parameter in the HP model, which results in an inference procedure of the following form:

\[ \hat{z}, \hat{\theta} = \arg\max_{z \in \mathbb{R}, \theta \geq 0} \log \mathbb{P}(\hat{t}, z, \theta), \]

(163)

where \( z \) is the random noise vector, and \( \theta \) corresponds to the parameters of the original exponential HP model.

B. Sequences with few events

In many domains, the availability of data is scarce, and the event streams will be composed of too few events, which results in noisiness of the likelihood and, as a consequence, unreliability of the fitted HP model, which calls for strong regularization strategies over the objective functions to be optimized.

For this type of situation, an approach, presented in [40], deals with HPs with triggering functions defined as exponentials and also as a mixture of gaussian kernels, as in [54]. Then, the parameters left to search are the background rates vector \( \mu \) and the tensor of weightings \( W \) for the excitation functions. The optimization is done through a Variational Expectation-Maximization algorithm which takes the distributions over these parameters as gaussians, and optimize, through Monte Carlo sampling, over an Evidence Lower Bound (ELBO) of their corresponding loglikelihood over a set of sequences.

C. Sequences with Missing data

Another issue in HP modeling revolves around learning from incomplete sequences, i.e., streams in which one or more of the events are missing. For this type of problem, two rather distinct approaches were recently proposed:

1) The first one, presented in [42], is applied to exponential and power-law HPs, and consists of a Markov Chain Monte Carlo-based inference over a joint process implicitly defined by the product of the likelihoods of the observed events and of the so-called virtual event
auxiliary variables, which are candidates for unobserved events. This virtual variable is weighted through a parameter \( \kappa \), which is related to the percentage of missing events with respect to the total event count;

2) The second one, introduced in [30], proposes finding the missing events over the sequences through importance weighting of candidate filling event subsequences generated by a bidirectional LSTM model built on top of the Neural Hawkes Process [29].

IX. CONCLUSIONS

Hawkes Processes are a valuable tool for modeling a myriad of natural and social phenomena. The present work aimed to give a broad view, to a newcomer to the field, of the Inference and Modeling techniques regarding the application of Hawkes Processes in a variety of domains.

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