Firing statistics of inhibitory neuron with delayed feedback. I. Output ISI probability density

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Abstract

Activity of inhibitory neuron with delayed feedback is considered in the framework of point stochastic processes. The neuron receives excitatory input impulses from a Poisson stream, and inhibitory impulses from the feedback line with a delay. We investigate here, how does the presence of inhibitory feedback affect the output firing statistics. Using binding neuron (BN) as a model, we derive analytically the exact expressions for the output interspike intervals (ISI) probability density, mean output ISI and coefficient of variation as functions of model’s parameters for the case of threshold 2. Using the leaky integrate-and-fire (LIF) model, as well as the BN model with higher thresholds, these statistical quantities are found numerically. In contrast to the previously studied situation of no feedback, the ISI probability densities found here both for BN and LIF neuron become bimodal and have discontinuity of jump type. Nevertheless, the presence of inhibitory delayed feedback was not found to affect substantially the output ISI coefficient of variation. The ISI coefficient of variation found ranges between 0.5 and 1. It is concluded that introduction of delayed inhibitory feedback can radically change neuronal output firing statistics. This statistics is as well distinct from what was found previously, [23], by a similar method for excitatory neuron with delayed feedback.

Keywords: Inhibitory neuron, Delayed feedback, Poisson process, Interspike intervals probability density, Coefficient of variation

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1. Introduction

A realistic neuronal network is normally characterized with a complicated system of excitatory and inhibitory interconnections between individual neurons the network is composed of. Statistics of spiking activity of individual neurons can be measured experimentally [9, 15, 18]. It would be interesting to understand how the details of network’s construction might influence statistics of neuronal activity, when the network is driven with some stimulation, or allowed reverberating freely. Exact theoretical analysis of this question in a developed network represents fair mathematical difficulties. At the same time, numerous studies suggest that feedback and delays in the intercomponent communication can be essential factors in determining activity of a composed system, see, e.g. [7, 4, 24].

In a real neural network, constructing elements can be found, which allow exact mathematical treatment. The results of such a treatment can shed light on the nature of transformations the neuronal activity might undergo while spreading within a real neural network. One example is an excitatory neuron which sends its output impulses onto its own dendritic tree — the neuron with excitatory feedback. This type of constructive element has been found in the olfactory bulb [1, 10]. Theoretical study of this construction fed with Poisson stream revealed interesting peculiarities in its output activity statistics [23]. Another natural variant of this construction is a neuron with inhibitory feedback. It seems that self-inhibition happens more frequently in the brain, than self-excitation. Self-inhibition can be slow, due to potassium channels opening [3], or fast, due to chlorine channels [2, 12]. It also can be direct (through autapses) [2, 10, 17], or acting through a single intermediate neuron [12]. Also, it can be evoked not only by means of a spike delivered to corresponding synaptic connection, but also through extended diffusion of some usual [17], or unusual [3] mediator.

In this paper, we consider situation of inhibitory neuron fed externally with excitatory impulses from the Poisson stream. The neuron sends its output impulses to its own input through feedback line with delay. Both input and output streams are treated as point stochastic processes with no diffusion approximation applied. Our purpose is to find the probability density function (PDF) of the output inter-spike intervals and to study its properties, as well as to compare those quantities for two neuronal models, namely the binding neuron and the leaky integrate and fire neuron.
2. Methods

2.1. BN without feedback

The binding neuron model [20] is inspired by numerical simulation [19] of Hodgkin-Huxley-type point neuron, as well as by the leaky integrate-and-fire (LIF) model [16]. In the binding neuron, the trace of an input is remembered for a fixed period of time after which it disappears completely. This is in the contrast with the above two models, where the postsynaptic potentials decay exponentially and can be forgotten only after triggering. The finiteness of memory in the binding neuron allows one to obtain exact mathematical conclusions concerning its firing statistics beyond the diffusion approximation technique. Recently, the finiteness is utilized for exact mathematical description of the output stochastic process if the binding neuron is driven with the Poisson input stream in the case of no feedback, [21], for BN with instantaneous feedback, [22] and for BN with delayed excitatory feedback, [23].

![Figure 1: Binding neuron with feedback (see [20] for details).](image)

Figure 1: Binding neuron with feedback (see [20] for details). $\tau$ is similar to the “tolerance interval” discussed in [8, p. 42]. Multiple input lines with Poisson streams are joined into a single one here.

The BN works as follows (see Fig. 1 with the feedback line removed). All excitatory input impulses have the same magnitude. Each one of them is stored in the BN for a fixed period of time, $\tau$, and then it is forgotten. When the number of stored excitatory impulses, $\Sigma$, becomes equal to the BN’s threshold, $N_0$, the BN fires an output spike, clears its internal memory, and is ready to receive fresh inputs. Thus, the state just after firing corresponds to the resting state of excitable membrane in real neurons, and the presence of impulses in the internal memory of BN corresponds to partially depolarized state. In this work, we take BN with $N_0 = 2$ for analytic derivation. BNs with higher thresholds are studied numerically in Sec. 4.2.3.

Normally, any neuron has a number of input lines. If input stream in each line is Poissonian and all lines have the same weight, all of them can be joined into
a single one, like in Fig. 1, with intensity, $\lambda$, equal to sum of intensities in the individual lines.

The output statistics for BN with $N_0 = 2$ and without feedback was studied before. Here, we will need the ISI probability density function for BN without feedback, $P^0(t)$, where $t > 0$ denotes the output ISI duration, which was obtained in [21, Eq. (3)] as

$$m\tau \leq t \leq (m+1)\tau \Rightarrow P^0(t) = y_m(t), \ m = 0, 1, \ldots,$$

where the functions $y_m(t)$ are defined according to the following recurrent relation:

$$y_0(t) = e^{-\lambda t} \lambda^2 t,$$

$$y_{m+1}(t) = y_m(t) + \frac{\lambda^{m+3}}{(m+2)!} (t - (m+1)\tau)^{m+2} e^{-\lambda t}$$

$$- \frac{\lambda^{m+2}}{(m+1)!} (t - (m+1)\tau)^{m+1} e^{-\lambda t}, \quad m = 0, 1, \ldots$$

$P^0(t)$ is a uni-modal function, which reaches its maximum at $t = \min(1/\lambda; \tau)$.

The first moment, $W^0_1$, of the probability density (1) was found in [21] as

$$W^0_1 \equiv \int_0^\infty t P^0(t) \, dt = \frac{1}{\lambda} \left( 2 + \frac{1}{e^{\lambda \tau} - 1} \right),$$

which will be used later. Further, we also utilize the probability $\Pi(t)$ to get from BN without feedback an output ISI, which is longer than $t$:

$$\Pi(t) \equiv \int_t^\infty P^0(t') \, dt' = 1 - \int_0^t P^0(t') \, dt'.$$

We will need the expression for $\Pi(t)$ only in the domain $0 \leq t \leq \tau$, where it can be found by substituting $y_0(t')$, (2), instead of $P^0(t')$ above:

$$\Pi(t) = (1 + \lambda t) e^{-\lambda t}, \quad 0 \leq t \leq \tau.$$

2.2. Feedback line action

In this work, we consider the situation, when BN receives excitatory input from the Poisson stream and inhibitory impulses from the feedback line.
We assume, that time delay $\Delta$ of impulse in the feedback line is fixed and is smaller than the BN’s memory duration, $\tau$:

$$\Delta < \tau.$$  \hspace{1cm} (6)

It allows us to make analytic expressions shorter.$^1$

If the line is empty, when neuron fires, an output impulse enters the line and after delay $\Delta$ reaches neuron’s input. If the line already keeps an impulse at the moments of BN firing, it does not accept a new one. It means, that at any given moment, the feedback line either conveys one impulse, or it is empty. The state of the nonempty feedback line can be described with a single number, $s$, $s \in [0; \Delta]$, which gives the time to live of the impulse in the feedback line. The time to live determines position of the impulse in the feedback line, see Fig 1, thick part of the feedback line. It equals to the time, which is required for the impulse to reach the end of the feedback line, if starting from a given position. The values of $s$ are used just at the moments of output ISI beginnings (just after BN firings). At these moments, the feedback line is never empty.

The inhibitory action of feedback impulses is modelled in the following way. When the inhibitory impulse reaches BN, it annihilates all excitatory impulses already present in the BN’s memory, similarly as Cl-type inhibition shunts depolarization of excitable membrane, see [14]. If at the moment of inhibitory impulse arrival, the BN is empty, then the impulse disappears without any action, similarly as Cl-type inhibition does not affect membrane’s voltage in its resting state. Such inhibition is "fast" in that sense, that the inhibitory impulses act instantaneously and are not remembered by neuron.

2.3. Derivation outline

It is clear, that both the binding neuron and the feedback line operate in deterministic manner. Nevertheless, the probabilistic description is required for the

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$^1$We were unable to found experimental value for $\Delta$ in an autaptic connection. A crude estimate can be made based on the action potential propagation velocity, $v$, and the connection length, $l$. We put $v = 0.5$ m/s (the smallest known). Expect that the autaptic connection is confined within a cortical minicolumn. The minicolumn diameter is about 50 $\mu$m. This gives for the propagation delay $\Delta_p \sim 3 \cdot l/v = 0.3$ ms. In this case, the delay of the feedback communication, $\Delta$, is mainly due to the EPSP rise time. The rise time usually falls between 1 ms and 10 ms, which to an extent supports (6) if one takes into account that $\tau$ should be comparable with the membrane relaxation constant, $\tau_M$. If one expects that the connection is confined within a cortical hypercolumn, then $\Delta_p \sim 3$ ms, which still does not refute (6). But if an autaptic connection is considerably longer, then condition, which is opposite to (6) may be correct.
output stream because of the stochastic nature of the driving Poisson process.

Let us denote by $P^\Delta(t)$ the ISI probability density function for neuron with delayed inhibitory feedback. In order to calculate $P^\Delta(t)$, we use the procedure, previously utilized for BN with excitatory delayed feedback [23]. For this purpose we define an auxiliary random variable $S$, with values $s \in [0; \Delta]$ are the values of time to live of the impulse in the feedback line at the beginning of an interspike interval. A possibility to introduce such a variable is ensured by the fact that, at the beginning of any ISI, the feedback line is never empty, see previous section.

It is clear that both $S$ and $T$ outcomes are uniquely determined by the unique outcome of the driving Poisson process. Here $T$ denotes the ISI random variable. The difference between $T$ and $S$ is that the $S$ outcome becomes known in physical time before the corresponding outcome of $T$ is known. That is, if at the beginning of an ISI, the $S$ outcome is $s$, then this $s$ together with the consequent outcome of the driving Poisson process determine the outcome $t$, corresponding to this $s$.

Due to this fact it is natural to define the conditional probability density, $P^\Delta(t | s)$.

As the first step, we calculate the conditional probability density, $P^\Delta(t | s)$.

Then, we calculate the probability density, $f(s), s \in [0; \Delta]$ for the variable $S$.

The output ISI probability density can be calculated based on the expressions for $P^\Delta(t | s)$ and $f(s)$, namely:

$$P^\Delta(t) = \int_0^\Delta P^\Delta(t | s) f(s) \, ds. \quad (7)$$

In order to find $f(s)$, we first obtain the transition probability density $P(s' | s), s, s' \in [0; \Delta]$, which gives the probability that at the beginning of some output ISI, the line has an impulse with time to live within the interval $[s'; s' + ds']$, provided that at the beginning of the previous ISI it had impulse with time to live equal $s$. $f(s)$ is then found as normalized to 1 solution of the following equation:

$$\int_0^\Delta P(s' | s) f(s) \, ds = f(s'). \quad (8)$$

3. Main calculation

3.1. Conditional probability density $P^\Delta(t | s)$

In order to derive $P^\Delta(t | s)$, domains $t < s$ and $t \geq s$ should be considered separately.
In the case \( t < s \), the output impulse must be generated without the line impulse involved. Therefore, probability density for such ISI values is the same as for BN without any feedback:

\[
P^\Delta(t \mid s) = P^0(t), \quad t < s.
\]  

Here \( P^0(t) \) is the output ISI probability density for BN without feedback, given in Eq. (1).

At the moment \( t = s \), the inhibitory feedback impulse reaches the BN and BN becomes empty. To trigger the neuron within the infinitesimal interval \([s; s + dt] \), one needs to get two input impulses within this interval. Probability of this event is of order \( dt^2 \). Therefore, \( P^\Delta(t \mid s) = 0 \) at \( t = s \).

In order to obtain ISI \( t > s \), two independent events must occur: (i) BN without feedback fires no spikes during time interval \([0; s] \); (ii) BN without feedback starts empty at moment \( s \) and is firstly triggered at moment \( t \). These events are independent since their realizations are defined by behavior of Poisson input stream on disjoint intervals \([0; s] \) and \([s; t] \). By definition of \( \Pi(t) \), see Eq. (5), the probability to have (i) is \( \Pi(s) \), and (ii) has the probability \( P^0(t - s) \). Therefore,

\[
P^\Delta(t \mid s) = \Pi(s) P^0(t - s), \quad t > s.
\]  

Taking into account Eq. (5), (9) and (10) for the case \( \Delta < \tau \) one obtains \( P^\Delta(t \mid s) \) as follows:

\[
P^\Delta(t \mid s) = \begin{cases} 
\lambda^2 t e^{-\lambda t}, & t \in [0; s], \\
(1 + \lambda s) e^{-\lambda s} P^0(t - s), & t \geq s.
\end{cases}
\]  

The conditional probability density \( P^\Delta(t \mid s) \), given in (11), is normalized: \( \int_0^\infty P^\Delta(t \mid s) dt = 1 \). Also, \( P^\Delta(t \mid s) \) has a jump discontinuity of height \( \lambda^2 s e^{-\lambda s} \) at \( t = s \).

3.2. Transition probability density \( P(s' \mid s) \)

From the definition of the transition probability \( P(s' \mid s) \) given in the last paragraph of Sec. 2.3 it follows that

\[
s' \geq s \quad \text{and} \quad s' \neq \Delta \quad \Rightarrow \quad P(s' \mid s) = 0.
\]

Indeed, consider the pair \((t, s)\), where \( t \) is the ISI duration and \( s \) is the impulse time to live in the feedback line when this ISI starts. If for this pair the inequality \( t \leq s \)
takes place, then \( s' = s - t \) and \( s' < s \) with necessity. In the opposite situation when \( t > s \), the line becomes empty before the end of the ISI \( t \). At the end of \( t \), the neuron starts next ISI by firing a spike, which charges the line with a fresh impulse. This means that in the next pair \((t', s')\), with necessity, \( s' = \Delta \). Therefore, the set of values \((s', s)\), where \( P(s' \mid s) \) still has to be found is defined by the following relations:

\[
\text{or } s' = \Delta.
\]

From the meaning of \( P^\Delta(t \mid s) \) it follows that Eq. (9) allows one to calculate \( P(s' \mid s) \) for \( s' < s \), namely:

\[
P(s' \mid s) = P^\Delta(s - s' \mid s) = P^0(s - s') = e^{-\lambda(s-s')}\lambda^2(s-s'), \quad s' < s \in [0; \Delta],
\]

where (11), (2) and (6) were used.

Consider the exact equality \( s' = \Delta \). It is fulfilled every time, when for previous ISI the inequality \( t \geq s \) holds, and this inequality happens with non-zero probability. Therefore, the probability density \( P(s' \mid s) \) has singularity of \( \delta \)-function type at \( s' = \Delta \). For calculating its mass it is enough to utilize the normalization condition:

\[
\int_{0}^{\Delta} P(s' \mid s) ds' = 1, \quad s \in [0; \Delta],
\]

which gives

\[
P(s' \mid s) = \begin{cases} 
  e^{-\lambda(s-s')}\lambda^2(s-s'), & s' < s \in [0; \Delta], \\
  (\lambda s + 1) e^{-\lambda s} \delta(s' - \Delta), & s' \geq s \in [0; \Delta].
\end{cases}
\]

### 3.3. Delays probability density

In order to find \( f(s) \), one should substitute \( P(s' \mid s) \) from (13) into (8) and solve the obtained equation. As \( P(s' \mid s) \) obtained here is exactly the same as for BN with excitatory delayed feedback [23, Eq. (9)], the equation for \( f(s) \) and \( f(s) \) itself will be the same also. In [23, Eq. (10)], the probability density \( f(s) \) was obtained as

\[
f(s) = a \delta(s - \Delta) + g(s),
\]

where \( g(s) \) – is an ordinary function, which vanishes out of interval \([0; \Delta]\):

\[
g(s) = \frac{a\lambda}{2} \left( 1 - e^{-2\lambda(\Delta-s)} \right), \quad s \in [0; \Delta],
\]
and $a$ – is the dimensionless constant:

$$a = 4e^{2\lambda \Delta} / \left( (2\lambda \Delta + 3)e^{2\lambda \Delta} + 1 \right),$$ \hspace{1cm} (16)

which gives the probability to find the impulse in the feedback line with time to live $\Delta$ at the beginning of any ISI.

3.4. Numerical simulations

Numerical simulations were carried out here for several purposes. The first one was to check numerically correctness of the expressions found analytically. The second one was to obtain ISI distributions for higher thresholds and for threshold 2 with $\Delta > \tau$. The third one was to compare the ISI distributions found here for the binding neuron model with those for the leaky integrate and fire (LIF) model.

A C++ program was developed, which allows us to calculate all the necessary probability distributions. The program includes the BNDF class, which analyzes the input stream and fires in accordance with the rules, described above. The Poisson streams of various intensity were produced with the help of the GNU Scientific Library\footnote{see http://www.gnu.org/software/gsl/}. With the help of our program, output stream samples were produced by calculating up to $N = 30000000$ output spikes. The samples were scanned for interspike intervals of various duration, and the probability density distribution was then calculated by normalization.

The LIF neuron was simulated in its simplest version. Namely, the neuron’s state at any moment of time $\vartheta$ is completely characterized by its membrane voltage at that moment, $V(\vartheta)$. Without stimulation, the $V(\vartheta)$ decays exponentially to the resting state with $V = 0$:

$$V(\vartheta + t) = e^{-t/\tau_M} V(\vartheta),$$

where $\tau_M$ – is the membrane relaxation time. An input impulse advances $V$ by a fixed value, $y_0$, instantaneously:

$$V \rightarrow V + y_0,$$

where $y_0$ mimics the EPSP peak value. If the resulting voltage satisfies the inequality

$$V + y_0 > V_0,$$
where $V_0$ is the firing threshold, then the LIF neuron fires an output spike and appears in the resting state.

For numerical simulations we choose $\tau_M = 10$ ms, $\gamma_0 = 4$ mV, $V_0 = 5$ mV. These values are comparable with those found in the inhibitory interneurons of CA3 hippocampal region, \[13\]. The relation between $V_0$ and $\gamma_0$ ensures that two input impulses are able to trigger the LIF provided they are close in time. For the inhibitory interneurons, this is because of their depolarized resting state, \[6\]. It is reported, \[5\], that even single impulse from a piramidal cell may trigger interneuron of this type. Interesting, that selfinhibition is found in the inhibitory interneurons also, but in the neocortex, \[2\].

4. Results

4.1. ISI probability density

In order to find $P^\Delta(t)$, one should substitute (11) and (14) into Eq. (7), which gives:

$$P^\Delta(t) = aP^\Delta(t \mid \Delta) + \int_0^\Delta P^\Delta(t \mid s)g(s)ds. \tag{17}$$

The explicit expression for $P^\Delta(t)$, which can be obtained by further transformations of (17), will be different for $t$ values belonging to different domains. This is because the exact expression for the $P^\Delta(t \mid s)$ is different for different domains, see top and bottom lines of Eq. (11) and Eq. (1). The boundaries of domains in which $P^\Delta(t)$ does not change its formula are dictated both by the first and the second term in (17). Namely, by taking (11) with $s = \Delta$ and using (1) one concludes that...
the first term in (17) retains the same formula in the adjacent domains separated with points

$$t = 0, \Delta, \tau + \Delta, 2\tau + \Delta, 3\tau + \Delta, \ldots$$

(18)

It appears that the second term in (17) as well retains the same formula in the domain bounded with first two points from (18) allowing one to obtain exact formula for the first domain. Namely, if $t \in [0; \Delta]$, then the first term in (17) turns into

$$aP^{\Delta}(t \mid \Delta) = a\lambda^{2} t e^{-\lambda t},$$

(19)

while integration domain in the second term should be split into two parts with the point $s = t$ and use either top, or bottom line of Eq. (11) in the corresponding part:

$$\int_{0}^{\Delta} P^{\Delta}(t \mid s) g(s) ds$$

$$= \int_{0}^{t} (1 + \lambda s) e^{-\lambda s} \lambda^{2} (t-s)e^{-\lambda (t-s)} g(s) ds$$

$$+ \int_{t}^{\Delta} \lambda^{2} t e^{-\lambda t} g(s) ds.$$  

(20)

By combining Eqs. (19) and (20), one obtains after transformations

$$P^{\Delta}(t) = \frac{2\lambda e^{-\lambda t}}{2\lambda \Delta + 3 + e^{-2\lambda \Delta}} \left( \frac{1}{6} \lambda^{3} t^{3} - \frac{1}{2} \lambda^{2} t^{2} + \lambda t \left( \frac{3}{2} + \frac{1}{4} e^{-2\lambda \Delta} + \frac{1}{4} e^{-2\lambda (\Delta-t)} \right) + \lambda^{2} t \Delta \right), \quad t < \Delta.$$  

(21)
When $t \geq \Delta$, one needs only the bottom line of Eq. (11) for calculating $P^\Delta(t)$, and Eq. (17) turns into the following:

$$
P^\Delta(t) = a (1 + \lambda \Delta) e^{-\lambda \Delta} \rho(0) (t - \Delta) + \int_0^\Delta (1 + \lambda s) e^{-\lambda s} \rho(0) (t - s) g(s) \, ds, \quad t \geq \Delta. \tag{22}
$$

Here, the first term cannot change its formula within any domain defined by boundaries (18). This is not the case for the second term. We analyze the behavior of the second term and obtain explicit expression for $P^\Delta(t)$ for any $t > 0$ in the Appendix. Example graph of $P^\Delta(t)$ found is given at Fig. 2 left. Compare with p.d.f. for BN without feedback, Fig. 3 left, and BN with excitatory feedback, right.

4.2. Properties of the ISI probability density

Notice, that the explicit expressions for $P^\Delta(t)$ given in (21) and in the Appendix are not used here. All the properties discussed below are derived from representation (7) and expressions (11), (14)–(16).

4.2.1. Mean interspike interval

The mean output ISI, $W_1^\Delta$, can be defined as the first moment of the ISI probability density:

$$
W_1^\Delta = \int_0^\infty t P^\Delta(t) \, dt.
$$
Taking into account Eq. (7), one obtains:

\[ W_{1}^{\Delta} = \int_{0}^{\Delta} dt \int_{0}^{\Delta} P^{\Delta}(t \mid s) f(s) ds = \int_{0}^{\Delta} ds f(s) \int_{0}^{\Delta} t P^{\Delta}(t \mid s) dt, \]

which taken together with Eq. (11) gives:

\[ W_{1}^{\Delta} = \int_{0}^{\Delta} ds f(s) \left( \int_{0}^{s} t^2 e^{-\lambda t} \lambda^2 dt + + (1 + \lambda s) e^{-\lambda s} \int_{s}^{\infty} t P^{0}(t-s) dt \right) \]
\[ = \frac{1}{\lambda} \int_{0}^{\Delta} ds f(s) \left( 2 + e^{-\lambda s} (\lambda W_{1}^{0} - 2 + (\lambda W_{1}^{0} - 1) \lambda s) \right), \]

where \( W_{1}^{0} \) is given in (4). Use here (14) and (15), which gives after transformations:

\[ W_{1}^{\Delta} = a(\Delta + W_{1}^{0}), \quad (23) \]

where \( a \) is given in (16).

Note that for \( \Delta = 0 \), Eq. (23) turns into the following:

\[ W_{1}^{\Delta} \Big|_{\Delta=0} = W_{1}^{0}. \]

This is consistent with the fact that for \( \Delta = 0 \) \( P^{\Delta}(t) \) turns into distribution for neuron without feedback, which is \( P^{0}(t) \) given in Eqs. (11–3), see Eq. (A.9).

The output intensity, \( \lambda_{o} \), defined as the mean number of impulses per time unit, is the inverse \( W_{1}^{\Delta} \):

\[ \lambda_{o}^{\Delta} = \frac{1}{W_{1}^{\Delta}} = \frac{(2\lambda \Delta + 3 + e^{-2\lambda \Delta})(1 - e^{-\lambda \tau})}{4(\lambda \Delta + 2 - (\lambda \Delta + 1)e^{-\lambda \tau})} \lambda, \quad (24) \]

where Eqs. (4), (16) and (23) were used. At large input rates the following relation takes place

\[ \lim_{\lambda \to \infty} \left( \frac{\lambda}{2} - \lambda_{o}^{\Delta} \right) = \frac{1}{4\Delta}, \quad (25) \]

This limiting relation, which is derived directly from Eq. (24), can be understood as follows. At moderate stimulation some input spikes are lost without influencing output due to high probability of long input ISI. At high intensity, every two consecutive excitatory input impulses trigger the BN and send impulse into the
feedback line, provided it is empty. Thus, output intensity should be \( \lambda / 2 \) minus firings, inhibited by the line. The maximum rate of inhibitory impulses, which can be delivered by the feedback line to the neuron’s input, is \( 1 / \Delta \), and this rate is attainable when \( \lambda \to \infty \). Each inhibitory impulse either cancels one excitatory impulse in the neuron, or does nothing if neuron appears empty at the moment of the feedback line dejection. For high input rates, the probabilities to find the neuron at any moment of time either empty, or storing one impulse seem both approaching 0.5. Thus, due to feedback line activity, about \( 1 / (2 \Delta) \) excitatory impulses will be eliminated every second from the input stream, and about half as much from the output stream, which explains (25).

Graphs of \( \lambda_o^\Delta \) vs \( \lambda \) are shown at the Fig. 4.

4.2.2. Coefficient of variation

The coefficient of variation (CV) \( c^\Delta_v \) of output ISIs is defined as dimensionless dispersion:

\[
c^\Delta_v \equiv \sqrt{\frac{W_2^\Delta}{W_1^\Delta} - 1},
\]

where \( W_2^\Delta \) is the second moment of the ISI probability density:

\[
W_2^\Delta \equiv \int_0^\infty t^2 P^\Delta(t)dt = \int_0^\Delta ds f(s) \int_0^\infty t^2 P^\Delta(t \mid s)dt.
\]
Figure 6: *Left* — Coefficient of variation vs $\lambda$ for BN with inhibitory delayed feedback (1), without feedback [22] (2) and with excitatory delayed feedback [23] (3), obtained analytically. Curves (1)–(3): $N_0 = 2$, $\tau = 10$ ms. *Right* — Coefficient of variation vs $\lambda$ for LIF neuron with inhibitory delayed feedback (1), without feedback (2) excitatory delayed feedback (3), found numerically. $\Delta = 2$ ms for (1), (3) in both panels.

By calculating integrals here and taking into account Eq. (4), one obtains:

$$
(c_\Delta^2) = \frac{B_1 e^{2\lambda \tau} + 2 B_2 e^{\lambda \tau} + B_3}{8 (2 + \lambda \Delta) e^{\lambda \tau} - \lambda \Delta - 1} - 1,
$$

where

$$
B_1 = 3 e^{-4\lambda \Delta} - 8 e^{-3\lambda \Delta} + 2 (6 \lambda \Delta + 13) e^{-2\lambda \Delta} - 8 (2 \lambda \Delta + 3) e^{-\lambda \Delta} + 12 \lambda^2 \Delta^2 + 52 \lambda \Delta + 51,
$$

$$
B_2 = -2 e^{-4\lambda \Delta} + 4 e^{-3\lambda \Delta} + 2 (-5 \lambda \Delta + \lambda \tau - 7) e^{-2\lambda \Delta} + 4 (2 \lambda \Delta + 3) e^{-\lambda \Delta} - 12 \lambda^2 \Delta^2 + 4 \lambda^2 \Delta \tau - 34 \lambda \Delta + 6 \lambda \tau - 24,
$$

$$
B_3 = e^{-4\lambda \Delta} + 2 (4 \lambda \Delta + 3) e^{-2\lambda \Delta} + 12 \lambda^2 \Delta^2 + 24 \lambda \Delta + 9
$$

see Fig. 5 *left*, 6 *left*.

It is clear, that for $\Delta = 0$ Eqs. (26), (27) must give the output ISI coefficient of variation of BN without feedback, $c_\Delta^0$. And indeed, substituting $\Delta = 0$ to (26) and (27), one obtains

$$
c^2_{\Delta=0} = \frac{1}{2e^{\lambda \tau} - 1} \cdot \left(2 e^{2\lambda \tau} + 2(\lambda \tau - 1) e^{\lambda \tau} + 1\right) = (c^0_\Delta)^2,
$$

where $c^0_\Delta$ was previously found in [22, Sec. 5.3].
Figure 7: ISI PDF (in s\(^{-1}\)), found numerically. Left — inhibitory BN with delayed feedback for \(N_0 = 2, \tau = 10\) ms, \(\lambda = 50\) s\(^{-1}\); right — inhibitory LIF neuron with delayed feedback. \(\Delta = 18\) ms in both panels.

### 4.2.3. Numerical Simulations

The numerical simulations were performed as described in Sec. 3.4. BN firing statistics is represented in terms of \(P^\Delta(t), f(s), \lambda^\Delta_o\) and \(c^\Delta_v\). In parallel to the analytic expressions, all these quantities were calculated numerically for various sets of parameters \(\tau, \Delta, \lambda\). Numerically calculated curves were then compared with corresponding analytic expressions given in Eqs. (21), (A.4) – (A.8), (14) – (16), (24), (26) and (27). It was found, that numerically obtained curves fit perfectly with mentioned analytic expressions, see example in Fig. 2.

The set of numerical simulations was performed for the case \(\Delta > \tau\) and/or \(N_0 > 2\), which is not covered by the analytic expressions obtained. The curves obtained are given in Fig. 4 right, (2) and (3), Fig. 5 left, (3), right, (1)-(3), Fig. 7 left, Fig. 8.

A number of numerical simulations were performed for the LIF neuron model with parameters given in Sec. 3.4. The data obtained is presented in Fig. 6 right, Fig. 7 right.

### 5. Discussion

The statement of the problem adopted in this work expects transformation of the input ISI PDF into the output ISI PDF. In the contrast to widely used diffusion approximation, which lacks the input ISI PDF, we treat both input and output streams as objects of the same mathematical nature, namely, the point stochastic processes. This is necessary if one expects to study activity of a neuron involved in the interneuronal communication with the time-coding as opposed to the rate-
coding paradigm. This approach allowed us to consider a system, where output impulses are conveyed directly to its input — neuron with feedback — the simplest case of interneuronal communication. The feedback is inhibitory here, which is in concordance with numerous self-inhibitory neurons observed in nature, see [2, 10, 17]. As a result, we obtained exact expressions for several quantities characterizing the activity of binding neuron with feedback, compared those quantities with those calculated numerically for the LIF neuron and made the following conclusions.

5.1. Conclusions

In this paper the binding neuron model with delayed self-inhibition is studied. The neuron is stimulated with point stochastic process — Poisson stream of given intensity. The ISI PDF was found as an exact function of the input intensity, $\lambda$, delay time, $\Delta$ and neuronal internal memory duration time, $\tau$ for the BN neuron with threshold $N_0 = 2$. The ISI PDF for higher thresholds is found numerically. The course of the PDFs found is bimodal due to a trough at $t = \Delta$, see Figs. 2 [7, 8]. The nature of the trough is the same as is the nature of peak in the case of excitatory feedback, which was discussed by L.Ricciardi, [11], and exactly calculated for the BN in [23].

Exact mathematical expressions are found for the mean interspike interval, and coefficient of variation. Those same quantities are found numerically for the leaky integrate and fire model with shunting type delayed self-inhibition. Both models studied deliver qualitatively similar results. We conclude that the character of dependencies found is mainly due to the presence of inhibitory feedback line.
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Appendix A.

Let us introduce a new variable of integration, \( u = t - s \) in the Eq. (22):

\[
P^\Delta(t) = a (1 + \lambda \Delta) e^{-\lambda \Delta} p^0(t - \Delta) \\
+ \int_{t-\Delta}^t (1 + \lambda (t - u)) e^{-\lambda (t-u)} p^0(u) g(t-u) du.
\] (A.1)

Here, due to (1)–(3), both the first term and the second one can change its formula with changing \( t \) value. The first term changes its form every time when \( t - \Delta \) crosses integer multiple of \( \tau \). This gives the boundary points (18). The second term in (A.1) can change its value every time when either \( t \), or \( t - \Delta \) crosses integer multiple of \( \tau \). This gives the following boundary points

\[
t = 0, \Delta, \tau, \tau + \Delta, 2\tau, 2\tau + \Delta, 3\tau, 3\tau + \Delta, \ldots
\] (A.2)

If the \( t \) value varies between two consecutive points from (A.2), then both terms in (22) retain their algebraic expression. In order to find explicit form of that expression at any domain of \( t \) defined by (A.2), we introduce two groups of domains, \( B_m \) and \( C_m \), namely:

\[
B_m = [m \tau + \Delta; (m + 1) \tau], \\
C_m = (m + 1) \tau; (m + 1) \tau + \Delta[,
\]

\( m = 0, 1, \ldots \)

Note, that the full range \( ]0; \infty[ \) of possible ISI values is covered by alternate domains \( B_m \) and \( C_m \), \( m = 0, 1, \ldots \) and the domain \( ]0; \Delta[ \) for which we already have the explicit formula (21).

If \( t \in B_m \), then \( m \tau \leq t - \Delta < t < (m + 1) \tau \), and one should substitute \( y_{m}(t) \) from (3), corresponding to that \( m \), instead of \( P^0(u) \) in the (A.1). If \( t \in C_m \), then \( m \tau < t - \Delta < (m + 1) \tau + \Delta < t \). Therefore, the domain of integration in the Eq. (A.1) should be split into two with point \((m + 1) \tau \), and as \( P^0(u) \) one should substitute either \( y_{m}(t) \), or \( y_{m+1}(t) \).

Appendix A.1. ISI probability density at the domains \( B_m \)

Thus, in the case \( t \in B_m \), one obtains for \( P^\Delta(t) \):

\[
P^\Delta(t) = a (1 + \lambda \Delta) e^{-\lambda \Delta} y_{m}(t - \Delta) \\
+ \int_{0}^{\Delta} (1 + \lambda s) e^{-\lambda s} y_{m}(t - s) g(s) ds, \quad t \in B_m,
\] (A.3)
which after integration gives:

\[ P^\Delta(t) = a (1 + \lambda \Delta) e^{-\lambda \Delta \cdot y_m(t - \Delta)} - \frac{a}{2} e^{\lambda (\Delta - \tau) \cdot y_{m+1}(t - \Delta + \tau)} + \frac{a}{2} e^{\lambda \tau \cdot y_{m+1}(t + \tau)} + \frac{a \lambda}{2} e^{-\lambda t} \sum_{k=1}^{m+1} \sum_{l=0}^{k} K_{kl} \lambda^{k-l} (t - (k-1)\tau)^{k-l} \]

\[ - \frac{a \lambda}{2} e^{-\lambda t} \sum_{k=1}^{m} \sum_{l=0}^{k} K_{kl} \lambda^{k-l} (t - k\tau)^{k-l}, \quad t \in B_m \]  

(A.4)

where

\[ K_{kl} = \frac{1}{2^l+2(k-l)!} \left( \frac{l+1}{(l+2)!} (-2\lambda \Delta)^{l+2} + l + 1 \right. \]

\[ - (l-1) e^{-2\lambda \Delta} - 2 \sum_{i=0}^{l} \frac{(-2\lambda \Delta)^{l-i}}{(l-i)!} \left( 1 + \frac{1 + l + 1 - i \cdot \lambda \Delta}{(l-1)!} \right) \].

Appendix A.2. ISI probability density at the domains \( C_m \)

Consider the case \( t \in C_m \). Taking into account Eqs. (1)–(3), one can rewrite (A.1) as follows

\[ P^\Delta(t) \bigg|_{t \in C_m} = a (1 + \lambda \Delta) e^{-\lambda \Delta \cdot y_m(t - \Delta)} + \int_0^{t-(m+1)\tau} (1 + \lambda s) e^{\lambda s \cdot y_{m+1}(t-s)} g(s) ds \]

\[ + \int_{t-(m+1)\tau}^{\Delta} (1 + \lambda s) e^{-\lambda s \cdot y_m(t-s)} g(s) ds \]

\[ = a (1 + \lambda \Delta) e^{-\lambda \Delta \cdot y_m(t - \Delta)} + \int_0^{\Delta} (1 + \lambda s) e^{\lambda s \cdot y_m(t-s)} g(s) ds \]

\[ + \frac{\lambda^{m+3}}{(m+2)!} e^{-\lambda t} \int_0^{t-(m+1)\tau} (1 + \lambda s) (t-s-(m+1)\tau)^{m+2} g(s) ds \]

\[ - \frac{\lambda^{m+2}}{(m+1)!} e^{-\lambda t} \int_0^{t-(m+1)\tau} (1 + \lambda s) (t-s-(m+1)\tau)^{m+1} g(s) ds. \]

It is useful to denote as \( P^\Delta_{B,m}(t) \) the right-hand side of Eq. (A.3) defined for all \( t \):
\[ P_{B,m}^\Delta(t) = a (1 + \lambda \Delta) e^{-\lambda \Delta y_m(t-\Delta)} + \int_0^\Delta (1 + \lambda s) e^{-\lambda \Delta y_m(t-s)} g(s) ds, \quad t > 0. \]

With this notation, one obtains:

\[ P_{\Delta(t)} \bigg|_{t \in C_m} = P_{B,m}^\Delta(t) + \frac{a \lambda}{2} e^{-\lambda \Delta} \cdot \rho_{m}^\Delta(\lambda (t - (m + 1) \tau)), \quad (A.5) \]

where

\[ \rho_{m}^\Delta(x) = \frac{2}{a \lambda (m + 2)!} \int_0^x (1 + v) (x - v)^{m+2} g \left( \frac{v}{\lambda} \right) dv \]

\[ - \frac{2}{a \lambda (m + 1)!} \int_0^x (1 + v) (x - v)^{m+1} g \left( \frac{v}{\lambda} \right) dv, \quad m = 0, 1, \ldots \quad (A.6) \]

Here the dimensionless variable of integration \( v = s \) was introduced.

Performing integration in (A.6) one obtains:

\[ \rho_{m}^\Delta(x) = \sum_{l=0}^{m+4} K_l x^l - e^{-2\lambda \Delta} + 2 e^{-2\lambda \Delta} = \sum_{l=0}^{m+3} D_l x^l, \quad (A.7) \]

where

\[ D_l = \frac{1}{2^{m+4-l} \cdot l!} \sum_{i=0}^{l} \frac{(-1)^{l-i} \cdot (m-1-i)!}{i!} \]

\[ K_l = \frac{(m-1-l)!}{2^{m+4-l} \cdot l!} e^{-2\lambda \Delta}, \quad l = 0, \ldots m+1, \]

\[ K_{m+2} = \frac{1}{4 \cdot (m+2)!} e^{-2\lambda \Delta} - \frac{1}{(m+2)!}, \quad (A.8) \]

\[ K_{m+3} = \sum_{i=1}^{m+2} \frac{(-1)^i i}{(m+2-i)! (i+1)!} + \frac{1}{(m+3)!}, \]

\[ K_{m+4} = \sum_{i=0}^{m+2} \frac{(-1)^i (i+1)}{(m+2-i)! (i+2)!} \]

\[ D_{m+2} = \frac{1}{4} \sum_{i=0}^{m+2} \frac{(-1)^i \cdot (i+1)}{i! (m+2-i)!}, \quad D_{m+3} = \frac{1}{2} \sum_{i=0}^{m+2} \frac{(-1)^i}{i! (m+2-i)!}. \]
Note, that in the case $\Delta = 0$, ISI probability density is completely defined by Eq. (A.4), which turns into probability distribution for BN without feedback given in (1):

$$P^{\Delta}(t)_{\Delta=0} = P^{0}(t), \quad t > 0.$$  \hspace{1cm} (A.9)

This indeed should be the case, because when $\Delta = 0$, inhibitory impulses always enter empty neuron and, therefore, the feedback line have no chance to affect the output stream. Naturally, the output ISI distribution for $\Delta = 0$ coincides with that found for BN without feedback.