Approximate Bayesian Inference for a Mechanistic Model of Vesicle Release at a Ribbon Synapse

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Abstract

The inherent noise of neural systems makes it difficult to construct models which accurately capture experimental measurements of their activity. While much research has been done on how to efficiently model neural activity with descriptive models such as linear-nonlinear-models (LN), Bayesian inference for mechanistic models has received considerably less attention. One reason for this is that these models typically lead to intractable likelihoods and thus make parameter inference difficult. Here, we develop an approximate Bayesian inference scheme for a fully stochastic, biophysically inspired model of glutamate release at the ribbon synapse, a highly specialized synapse found in different sensory systems. The model translates known structural features of the ribbon synapse into a set of stochastically coupled equations. We approximate the posterior distributions by updating a parametric prior distribution via Bayesian updating rules. We show that model parameters can be efficiently estimated for synthetic and experimental data from in vivo two-photon experiments in the zebrafish retina. Also, we find that the model captures complex properties of the synaptic release such as the temporal precision. Our framework provides a viable path forward for linking mechanistic models of neural activity to measured data.

1 Introduction

The activity of sensory neurons is noisy — a central goal of systems neuroscience has therefore been to devise probabilistic models that allow to model the stimulus-response relationship of such neurons while capturing their variability [1]. Specifically, linear-nonlinear (LN) models and their generalizations have been used extensively to describe neural activity in the retina [2,3]. However, these type of models cannot yield insights into the mechanistic foundations of the neural computations they aim to describe, as they do not model their biophysical basis. On the other hand, mechanistic models on the cellular or subcellular level have been rarely used to model stimulus-response relationships: they require highly specialized experiments to estimate individual parameters [4,5], making it difficult to employ them directly in a stimulus-response model; alternatively, they often result in an intractable likelihood, making parameter inference challenging [6].

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Here we make use of recent advances in approximate Bayesian computation (ABC) [6, 7, 8, 9, 10, 11] to fit a fully stochastic, biophysically inspired model of vesicle release from the bipolar cell (BC) axon terminal to functional two-photon imaging data from the zebrafish retina (Fig. 1). It includes a linear-nonlinear stage to model the stimulus dependency, and a set of stochastically coupled equations modeling biophysical properties of the BC synapse. At this so-called “ribbon synapse”, a specialized protein complex, the “ribbon”, acts as a conveyor belt that “tethers” and “loads” vesicles onto active zones for future release [12]. It organizes vesicles into multiple pools: the “docked” (or readily releasable) pool consists of a number of vesicles located directly above the plasma membrane, while the “ribboned” pool consists of vesicles attached to the ribbon further from the cell membrane. The docked vesicles are thus primed for immediate release and can be released simultaneously (so called multivesicular release, MVR). The ribboned vesicles are held in reserve to refill the docked pool as it is depleted by exocytosis [13, 14]. The transitions of vesicles between those pools can be modeled by a set of coupled differential equations [15, 4], which we extend to a stochastic treatment. In addition to photoreceptors and bipolar cells in the retina [16], ribbon synapses are featured in many other sensory systems, such as in auditory hair cells and the vestibular system [17].

Thus, our proposed Bayesian framework links stimulus-response modeling to a biophysically inspired, mechanistic model of the ribbon synapse. This may contribute to a better understanding of sensory computations across levels of description, with applications in a diverse range of sensory systems.

2 Previous work

Models of neural activity  Variants and extensions of LN models have been widely used to model the activity of retinal neurons [2, 18, 13]. In these descriptive models, the excitatory drive to a cell is modeled as the convolution of a receptive field kernel with the stimulus, followed by a static nonlinearity. The result of this computation sets the rate of a stochastic spike generator, most commonly using either a Binomial or Poisson distribution. These basic models have also been used to approximate BC activity [19], however they do not explicitly model the dynamics of vesicle release at the ribbon synapse. Existing mechanistic models of synaptic release often require highly specialized experiments to estimate parameters [20] or make only indirect inferences based on the spiking activity of post-synaptic cells [21, 22]. In addition, they have not been used to perform system identification. The linear-nonlinear kinetics (LNK) model [23] attempts to address this issue. After an initial LN stage, the LNK model passes this information into a “kinetics block” consisting of a first-order set of kinetic equations implicitly representing the availability of vesicles. However, the LNK model treats the states of the pools as rescaled Markov process and cannot easily account for discrete vesicle release or MVR at the given noise level of single synapses.
Table 1: Variables, parameters and distributions of the model.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
<th>Parameter</th>
<th>Movement distribution</th>
<th>Distribution</th>
</tr>
</thead>
<tbody>
<tr>
<td>time stretch of the kernel</td>
<td></td>
<td>( \gamma )</td>
<td>( N(\mu, \sigma^2) )</td>
<td></td>
</tr>
<tr>
<td>non-linearity</td>
<td></td>
<td>( k, h )</td>
<td>( N(\mu, \Sigma) )</td>
<td></td>
</tr>
<tr>
<td>correlation of exocytosed vesicles</td>
<td></td>
<td>( \rho )</td>
<td>( N(\mu, \sigma^2) )</td>
<td></td>
</tr>
<tr>
<td>( d_t )</td>
<td>exocytosed vesicles</td>
<td>( p_{d_t} )</td>
<td>Beta-Bin</td>
<td></td>
</tr>
<tr>
<td>( D )</td>
<td>vesicles at dock</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( r )</td>
<td>vesicles ribbon → dock</td>
<td>( p_{r} )</td>
<td>res. Binomial</td>
<td>( N(\mu, \sigma^2) )</td>
</tr>
<tr>
<td>( R )</td>
<td>vesicles on ribbon</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>( c )</td>
<td>vesicles cytoplasm → ribbon</td>
<td>( \lambda_c )</td>
<td>res. Poisson</td>
<td>( \Gamma )</td>
</tr>
</tbody>
</table>

We address these issues by proposing a model that combines LN modeling for system identification with a probabilistic, biophysically inspired model for the ribbon synapse, with the capability to model discrete, multi-vesicular release. In contrast to classical LN models, the parameters of this model are readily interpretable as they directly refer to biological processes.

Approximate Bayesian Computation  Many mechanistic models in computational neuroscience only provide means to simulate data and do not yield an explicit likelihood function. Therefore, their parameters cannot be inferred easily. In such simulator-based models, Bayesian inference can be performed through techniques known as Approximate Bayesian Computation or likelihood-free inference [8]. The general inference problem can be defined as follows: given some experimental data \( x_0 \) and a mechanistic, simulator-based model \( p(x|\theta) \) parametrized by \( \theta \), we want to approximate the posterior distribution \( p(\theta|x = x_0) \). The simulator model allows us to generate samples \( x_i \) given any parameter \( \theta \), but the likelihood function cannot be evaluated. Often, \( x_i \) is first mapped to a low dimensional space (so called “summary statistics”), in which a loss function is computed. This mapping defines the features the model is trying to capture [10].

There are two main approaches to solve the inference problem: (1) approximate the likelihood \( p(x_0|\theta) \) and then sample (e.g. via MCMC) to get the posterior [8, 10]. In this approach, guided sampling is often used to generate new samples and either train a neural network or update other parametric models for the likelihood [8, 9]. One disadvantage of this approach is that there is a second sampling step necessary to obtain the posterior, which can be as time consuming as the inference of the likelihood. (2) approximate the posterior \( p(\theta|x = x_0) \). In principle, inference via rejection sampling could be applied, but is often inefficient. Thus, recently proposed methods use parametric models (like a mixture of Gaussian) to approximate the posterior over several sampling rounds [6]. In our work, we use an ABC method of type (2) with parametric prior distributions and Bayesian updating rules to approximate the posterior distribution \( p(\theta|x = x_0) \). We show that it efficiently learns the parameters of the proposed release model.

3 Linear Non-Linear Release Model

Our model consists of two main parts (Fig. [1]): a linear-nonlinear (LN) stage models the excitatory drive to the BC and a release (R) stage, models the vesicle pools as dependent random variables (see Appendix [A] for pseudocode). Therefore, we refer to the model as LNR-model.

3.1 Linear-Nonlinear stage

The first stage of the LNR model is a standard LN model, in which a light stimulus \( l(t) \) is convolved with a receptive field \( w_\gamma \) to yield the surrogate calcium concentration \( ca(t) \) in the synaptic terminal which is then passed through a static nonlinearity:

\[
ca(t) = \int_{\tau=0}^{T} l(t-\tau)w_\gamma(\tau)d\tau .
\]
We assume \( w_{\gamma} \) to be a biphasic kernel in order to model the signal processing performed in the photoreceptor and the BC [15, 24] (Figure A, B). A single parameter \( \gamma \) was used to stretch/compress the kernel on the time axis to estimate the receptive field (see Appendix C). This could be generalized to allow for more flexible receptive field shapes in principle. We used a sigmoidal non-linearity to convert the calcium signal to the release probability:

\[
p_{d_t}(t) = 1 / (1 + \exp(-k(c_0 - h))),
\]

where the parameters for the slope \( k \) and the half activation \( h \) are inferred from the data. We add a small positive offset to the non-linearity and renormalized it to allow for spontaneous release.

### 3.2 Release stage

The second stage of the LNR model consists of a model for the synaptic dynamics based on the structure of the BC ribbon: we define variables representing the number of vesicles present in each pool of the ribbon and the number of vesicles moving between pools (see Table 1). We use capital letters to define the number of vesicles in a specific pool, and lowercase letters to indicate the moving pool of the ribbon and the number of vesicles moving between pools (see Table 1). We use capital letters to define the number of vesicles in a specific pool, and lowercase letters to indicate the moving pool of the ribbon and the number of vesicles moving between pools (see Table 1). We use capital letters to define the number of vesicles in a specific pool, and lowercase letters to indicate the moving pool of the ribbon and the number of vesicles moving between pools (see Table 1). We use capital letters to define the number of vesicles in a specific pool, and lowercase letters to indicate the moving pool of the ribbon and the number of vesicles moving between pools (see Table 1). We use capital letters to define the number of vesicles in a specific pool, and lowercase letters to indicate the moving pool of the ribbon and the number of vesicles moving between pools (see Table 1). We use capital letters to define the number of vesicles in a specific pool, and lowercase letters to indicate the moving pool of the ribbon and the number of vesicles moving between pools (see Table 1). We use capital letters to define the number of vesicles in a specific pool, and lowercase letters to indicate the moving pool of the ribbon and the number of vesicles moving between pools (see Table 1). We use capital letters to define the number of vesicles in a specific pool, and lowercase letters to indicate the moving pool of the ribbon and the number of vesicles moving between pools (see Table 1).

**Vesicle Release** To model the correlated release of docked vesicles, we use a beta binomial distribution. This is a binomial distribution for which the parameter \( p \) is itself a random variable, leading to correlated events [25]. The release probability \( p_{d_t} \) is assumed to be the output of the LN stage according to equation 1. To achieve a correlation \( \rho \) and a release probability of \( p_{d_t} \), the parameters for the beta binomial distribution are:

\[
\alpha_t = p_{d_t} \cdot \left( \frac{1}{\rho} - 1 \right) \quad \text{and} \quad \beta_t = \alpha_t \cdot \left( \frac{1}{p_{d_t}} - 1 \right),
\]

if \( p_{d_t} \neq 0 \).

Thus, in each time step, we first draw the parameter \( \hat{\rho}_t \) for the binomial distribution according to a beta distribution: \( \hat{\rho}_t \sim Beta(\alpha_t, \beta_t) \) and then sample the number of released vesicles \( d_t \) from a binomial distribution with parameters \( n = D_{t-1} \) (the numbers of vesicles at the dock) and \( \hat{\rho}_t \):

\[
p(d_t | D_{t-1}) = \begin{cases} 0 & \text{if } p_d = 0, \\ \binom{D_{t-1}}{d_t} \hat{\rho}_t^d (1 - \hat{\rho}_t)^{(D_{t-1} - d_t)} & \text{otherwise.} \end{cases}
\]

**Movement to the dock** We assume that \( r_t \) vesicles located at the ribbon move to the dock in each time step. Because there is a maximum number of vesicles \( D_{max} \) that can be docked, such that \( r_t + D_{t-1} \leq D_{max} \), we use a restricted binomial distribution to model stochastic vesicle docking:

\[
p(r_t | D_{t-1}, D_t) = \begin{cases} \binom{R_{t-1}}{r_t} p_r^r (1 - p_r)^{(R_{t-1} - r_t)} & \text{if } r_t < D_{max} - D_t \\ \sum_{r_t \geq D_{max}-D_t} \binom{R_{t-1}}{r_t} p_r^r (1 - p_r)^{(R_{t-1} - r_t)} & \text{if } r_t = D_{max} - D_t \\ 0 & \text{otherwise.} \end{cases}
\]

The first case is the standard binomial distribution with appropriate parameters, the second case models the assumption that moving more vesicles to the dock than its maximum capacity simply fills the dock.

**Movement to the ribbon** We assume a large number of vesicles available in the cytoplasm (which is not explicitly modeled), such that the number of vesicles \( c_t \) moving from the cytoplasm to the ribbon follows a Poisson distribution, again respecting the maximal ribbon capacity \( R_{max} \):

\[
p(c_t | R_t) = \begin{cases} e^{-\lambda \frac{R_t}{c_t}} \frac{(\lambda \frac{R_t}{c_t})^{c_t}}{c_t !} & \text{if } c_t < R_{max} - R_t \\ \sum_{c_t \geq R_{max}-R_t} e^{-\lambda \frac{R_t}{c_t}} \frac{(\lambda \frac{R_t}{c_t})^{c_t}}{c_t !} & \text{if } c_t = R_{max} - R_t \\ 0 & \text{otherwise.} \end{cases}
\]
In each round samples are drawn from the (proposal) prior, the model is evaluated and the response is mapped to its summary statistic. From this, the loss per parameter $\theta$ is calculated and the best samples are accepted and used to update the (proposal) prior via Bayesian updating rules.

4 Bayesian Inference of Model Parameters

In the previous section, we constructed a fully stochastic model of vesicle release from BCs, including an explicit mechanistic model of the ribbon synapse, reflecting the underlying biological structures. The maximal capacity of the dock $D_{\text{max}}$ was set based on measured data to the largest quantal event observed in the functional recordings ($D_{\text{max}} \approx 7 - 8$). $R_{\text{max}}$, the maximal capacity of the ribbon, was set to an estimate on the maximal number of vesicles at the ribbon in goldfish rod bipolar cells [26,27], but decreased to reflect the smaller size of BCs in zebrafish larva ($R_{\text{max}} \approx 50$).

Next, we developed an ABC framework for likelihood free inference to infer the remaining model parameters (Table 1) from functional two-photon recordings. Our method uses parametric prior distributions which are updated in multiple rounds via Bayesian updating rules to provide a unimodal approximation to the posterior (Figure 2). Briefly, in each round we first draw parameters $\theta$ from the (proposal) prior and evaluate several runs of the model $\hat{d}_i$ for each sampled parameter. Due to the stochasticity of the model, each evaluation returns a different trace, for which a summary statistic is calculated. This summary statistic reduces the dimensionality of the simulated trace to a low dimensional vector. Based on this the loss function $L(\theta)$ is calculated by comparing it to the summary statistic of the observed data. The best parameters are used to calculate a posterior, which is then used as a proposal prior in the next round (Fig. 2, pseudocode in Appendix E).

4.1 Prior distributions and inference

As priors, we used normal distributions for all parameters except for $\lambda_c$ (Table 1), where we used a gamma distribution (the conjugate prior to the Poisson distribution). Some parameters were bounded e.g. to the interval $[0, 1]$ and their distributions renormalized to effectively truncate the priors.

In each inference round, we used Bayesian updating rules to calculate the posterior distribution [28,29] based on the $j$ best parameters $\{\theta\}$. For example, in round $n+1$, we updated the hyperparameters for the multivariate normal distribution of the NL parameters, $k$ and $h$, as:

$$
\mu_{n+1} = \frac{\kappa_n}{\kappa_n + j} \mu_n + \frac{j}{\kappa_n + j} \bar{\theta}
$$

$$
\Lambda_{n+1} = \Lambda_n + S + \frac{\kappa_n j}{\kappa_n + j} (\bar{\theta} - \mu_n)(\bar{\theta} - \mu_n)^T,
$$

where $\bar{\theta}$ is the mean over the best parameters and $S = \sum_{i=1}^{j} (\theta_i - \bar{\theta})(\theta_i - \bar{\theta})^T$ the (unnormalized) covariance of these parameters. The mean is thus updated as a weighted average of the prior mean and the mean of the best parameters, with weights specified by $\kappa$, which is updated as $\kappa_{n+1} = \kappa_n + j$. The posterior degrees of freedom $\nu_{n+1}$, which is used to sample the covariance matrix $\Sigma$, is the prior degrees of freedom plus the updating sample size: $\nu_{n+1} = \nu_n + j$. With these updates we end up in a two step sampling procedure: first we draw the covariance $\Sigma_{(n+1)}$, for each sample $i$ of round $n+1$ from the inverse-Wishart distribution $\text{Inv-Wishart}(\Lambda_{n+1}^{-1}, \nu_{n+1})$, and then we draw the samples from the normal distribution $N(\mu_{n+1}, \Sigma_{(n+1)i})$. 

![Figure 2: Overview of the inference method.](image-url)
Figure 3: Results for synthetic data. A. Simulated traces for the synthetic data and simulations with the recovered, fitted parameters in response to a binary light stimulus. B. The time course of the mean and standard deviation of the different one dimensional marginal distributions over several rounds. Notice the asymmetric distribution for $\lambda_c$. See Appendix Fig. 8 for the two dimensional marginals. C. Histogram over the number of vesicles released per release event, error bars indicating ± std. D. Discrepancy of the data and fitted traces. The discrepancy is defined as the difference between the weighted summary statistics of a single data trace and the remaining data (“leave-one-out-procedure”) and accordingly the difference between the weighted summary statistics of a single fitted trace and the data. Error bars indicating ± std. E. The kernel of the linear stage. F. The non-linearity. Although its parameter $k$ is not matched perfectly in (B), there is almost no difference between the fitted and the true non-linearities.

The parameters for the univariate normal distributions as well as for the $\Gamma$-distribution are similarly updated in a Bayesian manner (see Appendix D). The number of drawn and accepted parameters was constant ($20 \cdot 10^3$ and 10) except for the first round, where the number of drawn parameters was doubled.

4.2 Summary statistics and loss function

As a summary statistic, on which the discrepancy between different traces is defined, we used (1) the histogram over the number of vesicles released in each event and (2) the Euclidean distance between the simulated and measured response trace, convolved with a Gaussian kernel (width: 100 ms, inspired by [30]). The former proved especially useful in early rounds of inference. As experiments typically consist of multiple repetitions of the same stimulus, we first calculated the summary statistics $s(d_i)$ for the individual traces $d_i$, normalized each entry by the summary statistic of the data traces and scaled it for its importance. This linear transformation is summarized in a diagonal weight matrix $W$. We used the average euclidean distance of these weighted summary statistics as the loss function $\mathcal{L}(\theta)$ (see also Appendix E and F). For $n$ data traces $d_i$ and a batchsize of $m$ simulations $\hat{d}_j$ per parameter $\theta$, this yields:

$$\mathcal{L}(\theta) = \frac{1}{nm} \sum_{i,j} ||Ws(d_i) - Ws(\hat{d}_j)||_2 .$$

The (weighted) summary statistics can also be used to calculate the variability of the data and compare it to the summary statistics of the simulated data, giving us an estimate of the discrepancy between the different traces (e.g. Fig. 3C, Fig. 5).
Figure 4: Two-photon imaging of in vivo zebrafish BCs allows for counting glutamatergic vesicles. A. Image of a zebrafish BC expressing the Superfolder-iGluSnFR transgene. Dashed circles indicate active zones where glutamate is released. B. Experimental glutamate release traces as $\Delta F/F$ of one OFF BC in two trials and extracted events in response to a binary light stimulus. Notice the high inter-trial variability.

5 Results

5.1 Model inference on synthetic data

Next, we tested whether we can successfully infer the parameters of the mechanistic model with the procedure outlined above. For that, we chose a realistic parameter setting and used the model to generate data. As the sample size per cell is severely limited in experimental data, we generated only four traces of 140 seconds each (Fig. 3A). The light stimulus was the same binary noise stimulus that we used for the experimental recordings.

The inference procedure proved very efficient: most parameters converged quickly to their true values with reasonable uncertainty (Fig. 3B). After inference, it is difficult to differentiate between the true and the fitted traces and the histogram over the number of vesicles released in each time step can be fitted well (Fig. 3C). Indeed, simulations from the model where as similar to the data as different data trials to one another (Fig. 3D). Our procedure identified the time scale of the linear receptive field as well as the non-linearity effectively (Fig. 3E and F). We validated the efficacy of our method also for other choices of parameters (not shown).

5.2 Model inference on BC recordings from zebrafish retina

We acquired two-photon imaging measurements of the glutamate release from BC axon terminals in the zebrafish retina ($n = 2$ cells, see Fig. 4). Briefly, linescans (1 x 120 pixels) were recorded at 1 kHz across the terminal of a BC expressing the glutamate reporter Superfolder-iGluSnFr, while showing a 140 second light stimulus (discrete Gaussian or binary noise) with a frame rate of 10 Hz. For each recording, a region of interest (ROI) was defined and the time series extracted as the weighted average of the spatial profile. Baseline drift was corrected, the traces were converted to $\Delta F/F$ and deconvolution was done with the kinetics of the reporter. Release events were identified as local maxima above a user-defined threshold in the imaging trace. The number of vesicles in each release event was estimated using a mixture of Gaussian model. For more details see [31].

Figure 5A shows the LNR model fitted to four recordings from one OFF BC (total duration of the recordings: 560 sec). We find that model parameters both for the LN-stage as well as the release stage of the model can be inferred efficiently. Posteriors converged quickly (Fig. 5B and Appendix Fig. 7). Interestingly, parameters such as the ribbon-to-dock-transition rate $p_r$, which model not directly observable properties of the system, also had larger uncertainty estimates. Similar to the synthetic data, the histogram over the number of vesicles released in each event was matched well overall (Fig. 5C). In contrast to the synthetic data, the discrepancy among data traces was a bit smaller than the discrepancy between the model fit and the data traces (Fig. 5D). This is likely due to the fact that some events were missed and the data contained more large amplitude events than predicted by the model (Fig. 5A, C).

We finally tested whether the simple model captured two known properties of release events: the temporal precision of events and the maximal release rates of the system. Interestingly, events with many released vesicles were temporally more precise for both the data and the fitted model (Fig. 5E, F). As no summary statistic explicitly measured the temporal precision of different release event types

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Figure 5: Results for experimental data. A. Two experimental data traces and simulated traces with corresponding fitted parameters in response to a binary noise stimulus. B. Some parameters are more restricted than others by the model. (See Appendix Section I for all parameters, the two-dimensional marginals, and the corresponding kernel and non-linearity) C. Histogram over event types. D. Discrepancy as in Fig. 3C. E. Temporal jitter of different event types in response to a binary light stimulus (mean±std, see Appendix Fig. 7 for the Gaussian noise stimulus). F. Cumulative release in response to a “calcium step” (see Appendix Fig. 11 for a comparison to experimental data).

at this resolution, this can be seen as evidence that our model captures crucial aspects of processing in BCs. Additionally, when comparing release rates with those recorded from electrically stimulated cells, we find the shape of cumulative vesicle release matches well with previously published results (Fig. 5F and Appendix J). This indicates that the model also extrapolates well to new stimuli.

6 Discussion

Here we developed a Bayesian inference framework for the LNR model, a probabilistic model of vesicle release from BCs in the retina, which combines a systems identification approach with a mechanistic, biophysically inspired component. In contrast to purely statistical models, the parameters of the LNR model are readily interpretable in terms of properties of the ribbon synapase. Specifically, we show that its parameters can be fitted efficiently on synthetic data and two-photon imaging measurements. The latter is remarkable, as mechanistic models often require highly specialized experiments to determine individual parameters. In this proof-of-principle study, we show that the parameters of the LNR model can be simply inferred from the functional measurements, opening possibilities for inferring mechanistic models from large-scale imaging experiments e.g. for comparison across cell types.

We found that the data overall was able to constrain the parameters very well, for both the LN-stage and the release-stage of the LNR model. Parameters that referred to parts of the model that were not directly observed in our measurements (such as the transition probability from the ribbon to the dock \( p_r \)) were fitted with somewhat higher uncertainty, indicating that a larger range of parameter values was compatible with the measurements. In addition, the LNR model captured MVR (the inferred correlation between vesicles is \( \rho \approx 0.35 \)), despite the inherent variability at the level of the single synapse. In addition, the LNR model captured trends in temporal precision within MVR events as well as release rates to non-physiological stimuli such as electrical stimulation - neither of which were used during inference.

Our proposed framework for Bayesian inference in the LNR model is comparable to recent likelihood-free inference methods (like \[6\] \[11\]). In contrast to those, we do not use a mixture density network (MDN) to approximate the posterior distribution, but rather parametric distributions and analytic Bayesian updating rules. In practice, MDNs can lead to unstable behavior for very small or large weights and sometimes have non-optimal extrapolation properties (but see \[32\]). Due to its simplicity, our method is less susceptible to such problems, but provides only a unimodal approximation of the
posterior $p(\theta | x = x_0)$. For the LNR model, we rarely observe multimodality in the posterior, so our methods yield a good and very fast approximation to the true posterior.

We combined a biophysically inspired mechanistic model with an efficient likelihood free inference method. This eases the development of more accurate and interpretable models, without the necessity of having closed-form likelihoods. At the same time, we could show that our model is able to capture the variability inherent to the neural system we studied. Taken together, the presented methods will allow for further investigations into more complex systems, gaining mechanistic insight into how neurons cope with noise.

**Acknowledgments**

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**References**


Appendix

A Pseudocode of the linear-non-linear release model

Algorithm 1: Pseudocode for the LNR-model

Calculate output of LN-block: \( p_{dt} = \text{NL}(w_{\gamma}(t) \ast l(t)) \);
Initialize pools \( D_0, R_0 \);
for \( t \in (1, \text{timesteps}) \) do
  Draw \( d_t \sim \text{beta Binomial}(D_{t-1}, \rho, p_{dt}) \);
  \( D_t = D_{t-1} - d_t \);
  Draw \( r_t \sim \text{restricted binomial}(R_{t-1}, p_r | D_t) \);
  \( D_t = D_t + r_t \);
  \( R_t = R_{t-1} - r_t \);
  Draw \( c_t \sim \text{restricted Poisson}(\lambda_c | R_t) \);
  \( R_t = R_t + c_t \)
end

B Experimental Methods

The data were collected and analyzed as described in [31], in accordance with redacted guidelines and with the approval of the Anonymous University local ethical committee. RibeyeA:Gal4 transgenic zebrafish were crossed with 10 x UAS:Superfolder iGluSnFR fish [33]. The resulting larva were imaged at 5-7 days post fertilization in vivo after embedding in 5 percent low-melting agarose. Imaging was done with a Scientifica microscope using a Coherent laser tuned to 915 nm. All stimuli were delivered full-field with an LED (\( \text{lmax} = 590 \text{ nm} \), Thorlabs), filtered through a 590/10 nm BP filter (Thorlabs), and output through a light-guide, resulting in a mean intensity of approximately 316 nW/mm\(^2\). Two white noise stimuli were used for the experiments: a binary stimulus constructed using an m-sequence protocol, and a discrete Gaussian stimulus, both with a frame rate of 10 Hz. Two cells recorded from two eyes of a single fish were used for the experiment.

C Linear kernel

Figure 6: Example kernels for 100 samples of compress/stretch parameter \( \gamma \), drawn from the prior distribution (\( \gamma \sim \mathcal{N}(1, 0.2) \)), and the corresponding kernels for the mean (red) and \( \pm2 \) standard deviations (dashed red). The remaining parameters were hold constant: \( \tau_r = 0.05, \tau_d = 0.05, \tau_{\text{phase}} = 100, \phi = -\pi/7 \).

The linear part of the model is the convolution of a kernel with the (light) stimulus \( l(t) \):

\[
ca(t) = \int_{\tau=0}^{T} l(t - \tau)w_{\gamma}(\tau)d\tau .
\]
We used a typical kernel for photoreceptors from previous work \cite{15, 24}, which depends on a rise and decay constant ($\tau_r$ and $\tau_d$) and additional phase parameters ($\phi$ and $\tau_{\text{phase}}$). To get different time scales, we included a new parameter $\gamma$ that stretches or compresses the kernel in time and was inferred from the data. Including $\gamma$, the equation of the kernel becomes:

$$w(t, \gamma) = \frac{- (\frac{t}{\gamma \tau_r})^3}{1 + (\frac{t}{\gamma \tau_r})^2} \cdot \exp \left( - \left( \frac{t}{\gamma \tau_d} \right)^2 \right) \cdot \cos \left( \frac{2\pi t}{\gamma \phi} + \tau_{\text{phase}} \right).$$

Figure 6 shows example kernels for different draws of $\gamma$.

### D Detailed updating rules

We follow the standard Bayesian updating rules to calculate the posterior \cite[page 73]{28}.

**Normal distributions** As already stated, we use a two dimensional multivariate normal distribution with unknown mean and covariance as distribution for the non-linearity parameters $k$ and $h$. For the remaining parameters, we take independent one dimensional normal distributions with unknown mean and variance (except for $\lambda_c$), such that the updating rules are basically the same: in round $n+1$, we take the $j$ best parameters $\{\theta\}$ and update as follows:

$$\mu_{n+1} = \frac{\kappa_n}{\kappa_n + j} \mu_n + \frac{j}{\kappa_n + j} \bar{\theta},$$

$$\kappa_{n+1} = \kappa_n + j,$$

$$\nu_{n+1} = \nu_n + j,$$

$$\Lambda_{n+1} = \Lambda_n + S + \frac{\kappa_n j}{\kappa_n + j} (\bar{\theta} - \mu_n)(\bar{\theta} - \mu_n)^T,$$

where $\bar{\theta}$ is the mean of the best parameters and $S = \sum_{i=1}^{j} (\theta_i - \bar{\theta})(\theta_i - \bar{\theta})^T$ the (unnormalized) estimation for the covariance of the best parameters.

To sample from the posterior distribution, we first draw $\Sigma$ from the inverse-Wishart distribution $\Sigma \sim \text{Inv-Wishart}(\Lambda_n^{-1}, \nu_{n+1})$, and then we draw samples from $\mathcal{N}(\mu_{n+1}, \Sigma)$.

For the one dimensional normal distributions we follow the same updating rules, except that the variance $\sigma_{n+1}^2$ is updated as following:

$$\sigma_{n+1}^2 = \frac{1}{\nu_n} (\nu_n \sigma_n^2 + S + \frac{\kappa_n j}{\kappa_n + j} (\bar{\theta} - \mu_n)^2)$$

To sample from the posterior we first draw $\sigma^2 \sim \text{Inv-}\chi^2(\nu_{n+1}, \sigma_{n+1}^2)$ and then draw samples from $\mathcal{N}(\mu_{n+1}, \sigma^2)$.

**Gamma distribution** The parameter $\lambda_c$ for the Poisson distribution has a $\Gamma$-distribution as conjugate prior, which is as well updated in a Bayesian way. We update the hyperparameters $k$ and $\theta$ of the $\Gamma$ distribution for the $j$ best parameters $\{\lambda_i\}_{i=1,\ldots,j}$ as:

$$k_{n+1} = k_n + \sum_{i=1}^{j} \lambda_i,$$

$$\theta_{n+1} = \frac{\theta_n}{1 + j \cdot \theta_n}.$$
Algorithm 2: Parametric approximation of the posterior via Bayesian updating

Input: 
- Simulator \( p(x|\theta) \); data traces \( \{x_{0i}\}_{i \in I} \);
- Summary statistics \( s(x) \) with weights \( W \);
- prior distribution \( p(\theta) \);
- number of rounds \( R \);
- simulations per round \( N \);
- batchsize \( B \);
- number of accepted parameters \( n_{\text{best}} \).

\[ \hat{p}_1(\theta) := p(\theta) ; \]

for \( r \in (1, R) \) do
  for \( n \in (1, N) \) do
    Draw \( \theta_{r,n} \sim \hat{p}_r(\theta) ; \)
    for \( b \in (1, B) \) do
      Simulate \( \hat{x}_{r,n,b} \sim p(x|\theta_{r,n}) ; \)
      end
    \end{array} \]
  end

end

\[ L(\theta_{r,n}) := \frac{1}{NB} \sum_{i,b} W s(x_{0_i}) - W s(\hat{x}_{r,n,b}) ||_2 ; \] // compute loss

end

\[ \theta_{\text{sorted}} := \text{sort}(\{\theta_{r,n}\}_{n \in N}) ; \]

\[ \theta_{\text{best}} := \theta_{\text{sorted}}[0 : n_{\text{best}}] ; \] // sort by loss function

\[ \hat{p}_{r+1}(\theta) := \hat{p}_r(\theta_{x_0}, \theta_{\text{best}}) ; \] // Bayesian updating of (proposal) prior

end

Return: \( \hat{p}_{R+1}(\theta) \)

F Details for the inference method

For the presented data we run 100 rounds with \( 20 \cdot 10^3 \) samples per round (except for the first round, where we used \( 40 \cdot 10^3 \) samples) and a batchsize of 4 simulated data traces per sampled parameter. We accepted the 10 best parameters per round (the ones producing the lowest loss) and used these to update our (proposal) prior.

For the prior distributions we used the following hyperparameters:

For the multivariate distribution for the non-linearity we used

\[ \mu_0 = (20, 0.5) \]
\[ \kappa_0 = 4 \]
\[ \nu_0 = 4 \]
\[ \Lambda_0 = \text{Id}_2 \cdot \left( \begin{array}{cc} 400 & 0.1 \end{array} \right) , \]

and truncated the distribution to the domain \((0, 50) \times (-2, 3)\).

For the univariate distribution for \( p_r(\rho, \gamma) \) we used

\[ \mu_0 = 0.3 \ (0.5, \ 1) \]
\[ \kappa_0 = 3 \]
\[ \nu_0 = 3 \]
\[ \sigma_0^2 = 0.05 \ (0.05, \ 0.2) , \]

and truncated the distribution to the interval \((0, 1) ((0, 1), (0.05, 2))\).

For the Gamma distribution \( \Gamma(k, \theta) \) we specified \( k_0 = 2 \), \( \theta_0 = 0.25 \ (E[\lambda] = k\theta) \) and bounds \((0, 1)\) as additional prior information.

As described in the main text we took the histogram over the different event types as summary statistics as well as the euclidean distance of the traces convolved with a Gaussian kernel \( g \). As additional entry we took the overall released vesicles. As scalings for for the summary statistics we took \( \text{Id}_x \cdot (5, 5, 5, 5, 2, 2, 4, 2)^t \) in the order \(|d_i * g - d_j * g| ||_2, \text{total release}, \text{total } 1_{d_j = 1}, \ldots, \text{total } 1_{d_j = 6}) \),

where \( d_i \) are the data traces and \( d_j \) is the simulated data. The weight matrix \( W \) is the product of the normalizing constants, which is the mean summary statistic of the data traces, with these scalings.
Figure 7: **Temporal jitter for the response to the Gaussian noise stimulus.** Same as Fig. 5E, but here for a Gaussian noise stimulus. As in previous studies [31], the lower contrast stimulation of the Gaussian noise stimulus results in higher temporal jitter when compared to the binary noise stimulus. This is well captured by the model. The exact numbers differ slightly to previous studies where a sinusoidal stimulus was used instead of binary and binned Gaussian noise.

G Temporal jitter

In order to define the temporal precision of release events, we begin by computing the vector strength for each quantal event type, $V S_q$:

$$VS_q = \frac{1}{N_q} \left[ \left( \sum_{i=1}^{N_q} \cos \left( \frac{2\pi t_{q_i}}{T} \right) \right)^2 + \left( \sum_{i=1}^{N_q} \sin \left( \frac{2\pi t_{q_i}}{T} \right) \right)^2 \right],$$

where $N_q$ is the number of $q$-quantal events, $t_{q_i}$ is the phase of the $i$-th $q$-quantal event with respect to the stimulus, and $T$ is the stimulus period. This metric takes values from zero to one, with zero representing complete independence between event times and stimulus phase, while a value of one indicates perfect phase-locking with all events occurring at exactly the same phase.

Temporal precision is more commonly expressed as temporal jitter, the standard deviation in milliseconds of the phase of the event times, as this yields a more intuitive interpretation. Vector strength can be mapped to temporal jitter by a one-to-one function:

$$TJ_q = \frac{\sqrt{2(1 - VS_q)}}{2\pi f},$$

where $f$ is the stimulus frequency.
H Results for the synthetic data

Figure 8: **Approximated marginals for the synthetic data.** One and two dimensional marginals of the posterior distribution for all parameters inferred from synthetic data in response to a binary noise stimulus.
1 Results for the experimental data

Figure 9: Marginals of the posterior for the experimental data (binary noise stimulus). A. The time course of the mean and standard deviation of the one dimensional marginal distributions. B. One and two dimensional marginals of the posterior.
Figure 10: **LN portion of the model for experimental data.** A. Linear kernel according to the prior and the posterior for the response to the binary noise stimulus. B. Non-linearity according to the prior and the posterior for the response to the binary noise stimulus.

**J Comparison to experimentally measured release rates**

Figure 11: **Inferred parameters reproduce the overall shape of vesicle release, as compared with previously published results.** A. Cumulative release rates for the model fit to experimental data bypassing the LN block (same as Fig. 5F). B. Estimates of the cumulative release from a goldfish rod bipolar cell in response to electric stimulation (taken from [34]).

Previous work has estimated maximum release rates from ribbon synapses by electrically stimulating the cell and estimating the cumulative number of vesicles released [34]. In order to compare these results with those estimated by fits of our model on experimental data, we bypassed the LN portion of our model, instead setting the release probability parameter \( p_d \) to mimic electrical stimulation. \( p_d \) was set to 0.8 during the period to simulate electrical stimulation, and 0.2 during the 250 ms period after the stimulation to mimic lingering increases in internal calcium (based upon experimental recordings [34]). All other parameters were set as fit with experimental data.

While the cumulative release between previous reports and our model vary by a constant scaling factor, this is likely due to the stimuli used to fit the model to experimental data. The maximal release rates estimated using physiological stimuli differ from those estimated using direct electrical stimulation.