Maximum likelihood estimation of biophysical parameters of synaptic receptors from macroscopic currents

Andrey Stepanyuk1,2*, Anya Borisyuk1,2 and Pavel Belan1,2

1 Laboratory of Molecular Biophysics, Bogomoletz Institute of Physiology, Kiev, Ukraine
2 State Key Laboratory of Molecular and Cellular Biology, Bogomoletz Institute of Physiology, Kiev, Ukraine

INTRODUCTION

Intrinsic biophysical properties of synaptic receptor channels are important for determining of both efficacy of synaptic transmission and activation of dendritic voltage-gated channels underlying active properties of dendrites. For example, synaptic NMDA receptors directly contribute to non-linear depolarizing drive in dendrites and control dendritic firing patterns and local dendritic Ca2+ concentration transients (Major et al., 2013). Changes in the postsynaptic receptor number, unitary conductance, and kinetics may affect dendritic integration (Magee, 2000) and lead to alteration in synaptic strength and memory function (Li and Stevens, 1977; Traynelis and Jaramillo, 1998; Milescu et al., 2005; Moffatt, 2007) all of them do not get over the quantal variability of postsynaptic currents. The new method also improves the accuracy of evaluation of unitary current as compared to the peak-scaled non-stationary fluctuation analysis, leading to a possibility to precisely estimate this important parameter from a few postsynaptic currents recorded in steady-state conditions. Estimation of unitary current with this method is robust even if postsynaptic currents are generated by receptors having different kinetic parameters, the case when peak-scaled non-stationary fluctuation analysis is not applicable. Thus, with the new method, routinely recorded postsynaptic currents could be used to study the properties of synaptic receptors in their native biochemical environment.

Keywords: unitary current, synaptic currents, peak-scaled non-stationary fluctuation analysis, maximum likelihood, semiseparable matrix, kinetic model, Markov chain Monte Carlo
By overcoming computational complexity that emerges due to
quantal variability of postsynaptic currents, a maximum likeli-
hood non-stationary fluctuation analysis (ML NSFA) suggested in
this work makes it possible to estimate unitary currents, number of
channels bound with a neurotransmitter, peak open proba-
bility, and some kinetic constants for synaptic channels in their
native biochemical environment from the experimentally feasible
number of macroscopic postsynaptic currents.

MATERIALS AND METHODS

KINETIC MODEL

In this work we consider simulated macroscopic synaptic currents
generated by a varying number of synaptic receptor channels. The
channels are assumed to be independent and identical. We assume
that the synaptic channel gating is a Markov process and \( p_{ij} \)
the probability of channel transition from state \( j \) to state \( i \) at
time \( \Delta t \). The rate matrix is a \( N_s \times N_s \) matrix \( \mathbf{Q} : q_{ij} = \lim_{\Delta t \to 0} \frac{p_{ij}}{\Delta t} \), \( N_s \)
is the number of states of the synaptic channel model. Each
element of the matrix \( \mathbf{Q} \) gives the rate constant of transition \( j \rightarrow i \) if
the transition is allowed by the model and otherwise \( q_{ij} = 0 \). The
diagonal elements, \( q_{ii} \), are set to \( -\sum_j q_{ij} \), so the sum over each col-
umn is zero. Synaptic release of neurotransmitter is modeled as a
step pulse of its concentration in the synaptic cleft, which leads
to instantaneous change of concentration-dependent transition
probabilities, \( p_{ij} \).

We assume that the kinetic matrix topology (i.e., a set of
allowed transitions) and the number of conducting states are
known. The required model parameters were arranged into the
parameter vector \( \theta = [q, i_{ch}, N_{ch}] \) and they were: rate constants,
\( q_{ij} = \lim_{\Delta t \to 0} \frac{p_{ij}}{\Delta t}, i \neq j \), unitary currents, \( i_{ch} \), and the number of
postsynaptic receptors bound with a neurotransmitter right after
the concentration transient, \( N_{ch} \).

THE LOG-LIKELIHOOD FUNCTION

The likelihood function, \( L_\theta \), that is to be maximized by ML NSFA
in order to find the most likely set of parameters is defined as the
conditional probability to observe \( N \) macroscopic current traces \( c_i \), \( i = 1 : N \), sampled at time points \( t = [1 \ldots N_T] \) given the
model parameters \( \theta \) (Colquhoun and Hawkes, 1977; Celentano
and Hawkes, 2004; Milesceu et al., 2005; Stepanyuk et al., 2011):

\[
L_\theta = P(c|\theta) \xrightarrow{N_{ch} \to \infty} \frac{1}{(2\pi)^{N_T/2}} \prod_{i=1}^{N} \frac{1}{N_{ch}} e^{-\frac{1}{2} \sum_{i=1}^{N} \left( (c_i - \mu_{N_{ch}}) \right)^T \left( \mathbf{c}_{m1} \right)^{-1} \left( c_i - \mu_{N_{ch}} \right)}
\]

Here \( N \) is the number of synaptic macroscopic current traces \( c_i \) (sample size) and \( N_T \) is the number of points in each trace;
\( \mathbf{c}_{m1} \) is a \( N_T \times 1 \) vector and \( \mathbf{c}_{m1} \), an \( N_T \times N_T \) matrix with elements \( \{\mathbf{c}_{m1}\}_{i,r} \), and denote mean and covariance of single
channel current, respectively, and they both are the functions of \( \theta \) : \( \mathbf{c}_{m1} \) is related to the covariance matrix, \( \mathbf{c}_{m} \), of a macroscopic
synaptic current \( c_i \) by the following expression: \( \mathbf{c}_{m} = \mathbf{c}_{m1} N_{ch} \).

Mean and covariance follow equations (Colquhoun and Hawkes,
1977)

\[
\mu = \mathbf{c}_{i} e^{\mathbf{Q}t} p(0)
\]

Here \( Q \) is a rate matrix (Colquhoun and Hawkes, 1977; Celentano
and Hawkes, 2004) and \( p(0) \) is an initial state vector. The elements of \( p(0) \) can be calculated as the equilibrium probabilities
determined by the initial experimental conditions, which are
assumed to last for sufficiently long time \( T \) to allow the channels
reach equilibrium

\[
p(0) = \prod_j e^{\mathbf{Q}t} p(-T)
\]

It is generally accepted to maximize the logarithm of the like-
lihood function \( \log L_\theta \) instead of the likelihood function \( L_\theta \)
itself. Therefore, our objective was to find the most likely model
parameter set, \( \theta_{ML} \), i.e., the parameter set that maximized the
log-likelihood

\[
\theta_{ML} = \arg \text{max}_{\theta} (\log L_\theta)
\]

The log-likelihood function \( \log L_\theta \) can be efficiently estimated
using the fact that \( \mathbf{c}_{m1} \) has a specific structure of semiseparable
matrix (DeWilde and van der Veen, 1998; Stepanyuk et al., 2011).

EFFICIENT ESTIMATION OF THE LOG-LIKELIHOOD FUNCTION FOR
SYNAPTIC CURRENTS WITH NOISE

Efficient log-likelihood estimation used in this article is based on
our previously described method (Stepanyuk et al., 2011). Briefly,
the method was based on the fact that the covariance matrix \( \mathbf{c}_{m1} \),
has a specific structure of semiseparable matrix, namely \( \mathbf{c}_{m1} \) can be represented as Stepanyuk et al. (2011).

\[
\{\mathbf{c}_{m1}\}_{ij} = \sum_{k=1}^{N_{s}} A_{ik} B_{kj}, \quad i \geq j
\]

\[
\{\mathbf{c}_{m1}\}_{ji} = \{\mathbf{c}_{m1}\}_{ij}, \quad i \geq j
\]

where

\[
A_{ik} = e^{\lambda_{k} t_{0}} \sum_{o=1}^{N_{o}} i_{o} U(o, k), 1 \leq k \leq N_{S}
\]

\[
A_{ik} = \mu_{k}, \quad k = N_{S} + 1
\]

\[
B_{kj} = e^{\lambda_{k} t_{j} - \lambda_{k} t_{0}} \sum_{o=1}^{N_{o}} U^{-1}(k, o') p_{o}(t_{j} i_{o'}), 1 \leq k \leq N_{S}
\]

\[
B_{kj} = -\mu_{k}, \quad k = N_{S} + 1
\]

where \( U : e^{\mathbf{Q}t} = \mathbf{Ue}^{\mathbf{D}t}\mathbf{U}^{-1} \) is \( N_{s} \times N_{s} \) matrix of the eigenvectors of \( \mathbf{Q} \), \( \lambda \) are different; \( N_{o} \) is a number of open states
in the channel model. Efficient linear algebra algorithms for
semiseparable matrices (Vandebril et al., 2007; Eidelman and
Gohberg, 2008) allowed us to compute the log-likelihood and provided almost linear scaling of its computational cost with the number of states in a kinetic model for the case of sufficiently large number of currents, ensuring fast, and accurate estimation of model parameters. The number of synaptic channels exposed to neurotransmitter was assumed to be the same for all currents. However, in the case of synaptic currents this number could vary between trials due to quantal variability. As a result, logL_c must be estimated separately for each current, and then summed up, thus increasing the number of operations in N times at least. However, calculations could be substantially simplified if the majority of receptors, which will participate in the current, are found in one particular state immediately after the neurotransmitter concentration transient, as it is expected for the synaptic receptors. To compute logL_c in this case, let denote noisy macroscopic synaptic current with an NT × 1 vector c_l and let denote by n_l an NT × 1 vector of noise imposed on the i-th current. Then −logL_c of the set of parameters θ given macroscopic synaptic currents without noise imposed on them is (we omit here the constant term NN_T log (2π)/2)

\[ - \log L_c (c - n) = \frac{1}{2} \sum_{i=1}^{N} (c_i - n_i - \mu_{N_{ch_i}})^T \frac{c_{m1_i}^{-1}}{N_{ch_i}} \]

\[ + \sum_{i=1}^{N} \left( \log |c_{m1}\rangle \right) \]

where μ is an expectation of synaptic current without noise and logL_c (c - n) denotes the required log-likelihood given the set of macroscopic synaptic currents without noise. Equation (8) can be rewritten as

\[ - \log L_c (c - n) = - \log L_c (c) - \frac{1}{2} \sum_{i=1}^{N} n_i^T c_{m1_i}^{-1} n_i \frac{1}{N_{ch_i}} \]

\[ - \sum_{i=1}^{N} n_i^T c_{m1_i}^{-1} (c_i - n_i - \mu_{N_{ch_i}}) \]

Since the background noise and the macroscopic current are uncorrelated the last term in Equation (9) can be neglected without loss in accuracy given the number of currents, N, is large enough. Therefore, we have

\[ - \log L_c (c - n) = - \log L_c (c) - \frac{1}{2} \sum_{i=1}^{N} n_i^T c_{m1_i}^{-1} n_i \frac{1}{N_{ch_i}} \]

Finally, from Equation (11) we obtain

\[ \sum_{i=1}^{N} n_i^T c_{m1_i}^{-1} n_i \frac{1}{N_{ch_i}} = \sum_{k=1}^{NT} \sum_{j=1}^{NS} \left( c_{m1_k}^{-1} \circ c_{noise} \right) N \left( \frac{1}{N_{ch_i}} \right) \]

where \( \circ \) denote Hadamard multiplication.

Keeping in mind that \( \sum c_{m1_i}^{-1} \circ c_{noise} = tr \left( c_{m1}^{-1} c_{noise} \right) \), we rewrite Equation (10) for logL_c (c - n) as

\[ - \log L_c (c - n) = - \log L_c (c) - \frac{1}{2} \sum_{i=1}^{N} \left( c_i - \mu_{N_{ch_i}} \right)^T c_{m1_i}^{-1} \left( c_i - \mu_{N_{ch_i}} \right) \]

\[ + N_T \log N_{ch_i} + N_T \log |c_{m1}| \]

is the log-likelihood function of macroscopic synaptic currents with noise.

To quickly evaluate \( tr \left( c_{m1}^{-1} c_{noise} \right) \) we note that matrices \( c_{m1}^{-1} \) and \( c_{noise} \) are quasiseparable (as an inverse of semiseparable matrix, Vandebril et al., 2007) and semiseparable matrix, respectively. Semiseparability of noise covariance matrix, \( c_{noise} \), follows from the fact that experimental background noise can be well approximated by a stationary Gaussian process, and the covariance matrix of such process is semiseparable (De Wilde and van der Veen, 1998). Then, the computation of trace of the product of such matrices can be accelerated by representing it as \( tr \left( F \cdot C \right) = 2tr(H \cdot B) + \sum_{k=1}^{NT} F_{kk} d_k \), where H is \((NT - 1) \times NS \) matrix, F is symmetric \( NT \times NT \) semiseparable or quasiseparable matrix and B is defined by Equation (7) (see also Equations A1.26–A1.35 from Text S1 in Appendix in Stepanyuk et al., 2011).

**ESTIMATION OF THE NUMBER OF CHANNELS AND PEAK OPEN PROBABILITY**

To estimate the number of channels \( N_{ch_i} \) (see Results for further definition), we re-write Equation (8) for a single macroscopic synaptic current:

\[ - \log L_c (c_i - n_i) = \frac{1}{2} \left( c_{m1_i}^{-1} c_i + \mu^T c_{m1_i} \mu_{N_{ch_i}} \right) \]

\[ - \sum_{k=1}^{NT} \left( c_{m1_k}^{-1} c_{noise} \right) N \left( \frac{1}{N_{ch_i}} \right) \]

\[ - 2 \mu^T c_{m1_i} \]

\[ + \frac{N_T}{2} \log N_{ch_i} + \frac{1}{2} \log |c_{m1}| \]

In the last expression we neglect the 3-d term, as it was done in Equation (9), and the 5-th and the last terms does not depend
on $N_{\text{ch}}$ at all. Leaving terms that depend on $N_{\text{ch}}$ only we obtain log-likelihood as a function of the number of channels:

$$- \log L_\theta (c_i - n_i) = \frac{1}{2} \left( \frac{c_i^T c_{m1}^{-1} c_i - n_i^T c_{m1}^{-1} n_i}{N_{\text{ch}}} + \mu^T c_{m1}^{-1} \mu N_{\text{ch}} \right) + \frac{N_T}{2} \log N_{\text{ch}}$$

(16)

The number of channels, $N_{\text{ch}}$, can be approximated for each macroscopic synaptic current $c_i$ as a number that gives maximum of the likelihood function when being substituted into Equation (16). Therefore, after differentiation of Equation (16) with respect to $N_{\text{ch}}$

$$\frac{\partial \log L_\theta (c_i - n_i)}{\partial N_{\text{ch}}} = 0 = \frac{1}{2N_{\text{ch}}} (c_i - n_i)^T c_{m1}^{-1} (c_i - n_i) - \frac{N_T}{2N_{\text{ch}}} \mu^T c_{m1}^{-1} \mu + N_{\text{ch}} N_T$$

(17)

we find an approximation for the number of channels, $N_{\text{ch}}$, for each macroscopic synaptic current

$$N_{\text{ch}} = \frac{-N_T + \sqrt{N_T^2 + 4c_i^T c_{m1}^{-1} c_i \cdot \mu^T c_{m1}^{-1} \mu}}{2\mu^T c_{m1}^{-1} \mu}$$

(18)

Here $c_i$ is not the whole decaying phase of each current but only those part where signal-to-noise ratio is high and therefore noise term can be neglected (usually from peak of the current to 0.1–0.5 of the peak). Therefore, before calculating the log-likelihood, we first estimate $N_{\text{ch}}$ for each macroscopic synaptic current $c_i$, then substitute $N_{\text{ch}}$ into Equations (13) and (14) and calculate the log-likelihood of the set of parameters $\theta$ given the set of simulated macroscopic synaptic currents. Accordingly, $N_{\text{ch}}$ is estimated automatically when the maximization is finished.

The peak open probability, $P(o, \text{peak})$, was defined as a probability that a channel is opened at the peak of the macroscopic current given that this channel was bound with a neurotransmitter immediately after the end of concentration transient, which was assumed to be sufficiently short (0.1–0.2 ms) with respect to the time interval (1–4 ms) from the moment of stimulation to the starting point of the analyzed fragment of current. $P(o, \text{peak})$ can be expressed as a function of rate constants: $P(o, \text{peak}) = \max (e^{\theta^T p(0)})$, where $p(0)$ is an initial state probability vector assumed to be zero for all states except for the RG2 state in the case when currents were simulated with 7-state GABA$_A$R scheme or RL state in the case when currents were simulated with simple 3-state kinetic scheme (see descriptions of both schemes below).

Summing up, M1 NSFA can be used for the fast estimation of rate constants, unitary current of synaptic ion channel, the number of synaptic channels bound with a neurotransmitter right after the concentration transient for each synaptic current and peak open probability from the set of macroscopic synaptic currents under Gaussian colored background noise.

**THE LOG-LIKELIHOOD MAXIMIZATION PROCEDURE**

We search for the log-likelihood global maximum to obtain the required model parameters from a set of macroscopic synaptic currents. In order to do this, we minimize the negative log-likelihood with a variant of graduated optimization technique using SQP algorithm embedded in finicon function in MATLAB Optimization toolbox. Initial estimates of each parameter were chosen randomly and uniformly from the logarithmic scale interval, $[\theta_0/10, \theta_0 \cdot 10]$, where $\theta_0$ is a vector composed of the true values of each parameter (rate constants and unitary current), i.e., of values utilized by the macroscopic current generator (see below). During the search of a minimum, all parameters were bounded within the interval $[\theta_0/50, \theta_0 \cdot 50]$.

In our version of graduated optimization technique, the whole minimization procedure was divided into sequential minimization steps. On the first step the negative log-likelihood was minimized given the first 2 or 3 currents regularly sampled at 50 points each. On each consequent minimization step the number of points and the number of currents was increased. The parameter estimates, $\theta_{ML}$, obtained on each previous step were taken as initial parameters $\theta_0$ for each next minimization step. For all calculations in this work each minimization was rerun 5 (3-state scheme) or 10 (7-state scheme) times, each time starting from the different initial parameter set.

**SIMULATION OF MACROSCOPIC SYNAPTIC CURRENTS**

First series of simulations of macroscopic synaptic currents were based on experimentally derived 7-state kinetic scheme for GABA$_A$ receptor that had one unliganded state, R, two liganded closed states (RG, singly-ligated and RG2, doubly-ligated) and the respective open (O1 and O2) and desensitized (D1 and D2) states (Mozrzymas et al., 2003). The following rate constants were adapted from Mozrzymas et al. (2003): $k_{\text{off}} = 0.13 \text{ ms}^{-1}$, $d_1 = 0.14 \text{ ms}^{-1}$, $d_2 = 1.5 \text{ ms}^{-1}$, $r_1 = 0.02 \text{ ms}^{-1}$, $r_2 = 0.12 \text{ ms}^{-1}$, $a_1 = 1.5 \text{ ms}^{-1}$, $a_2 = 1 \text{ ms}^{-1}$, $b_1 = 0.15 \text{ ms}^{-1}$, $b_2 = 0.8 \text{ ms}^{-1}$; $k_{m1} = 4 \text{ ms}^{-1}$ M$^{-1}$, $k_{m2} = 8 \text{ ms}^{-1}$ M$^{-1}$; Unitary current was the same for singly- and doubly-ligated states and was set to 1 pA. Variability in the amplitude of macroscopic postsynaptic responses was achieved by trail-to-trial Gaussian variation of the number of available synaptic channels (mean = 250; SD = 50). Simulation time step was $\Delta t = 0.2 \text{ ms}$. Synaptic vesicle release was modeled as a square pulse of saturating agonist concentration with a duration equal to the single simulation time-step ($\Delta t = 0.2 \text{ ms}$), which caused transition of all available channels from R to RG2 state. A total of 1000 macroscopic synaptic currents were simulated and colored noise that resembled baseline noise of experimentally recorded IPSCs was added to each current. Colored noise ($SD = 3 \text{ pA}$) was modeled as a sum of 4 AR(1) processes (Qin et al., 2000; Venkataramanan and Sigworth, 2002):

$$\text{noise}_t = \sum_{k=1}^{N_{\text{noise}}} \text{noise}_{t,k} + \text{noise}_{t,k} = \varphi_k \text{noise}_{t-1, k} + \sigma_w \text{noise}_{t,k} \sim N(0,1)$$

(19)
with parameters $\varphi = [0.0067, 0.61, 0.96, 0.999]$ and $\sigma = [0.32, 1.0, 1.42, 0.72]$, pA that were obtained from the approximation of autocorrelation function of experimentally recorded (whole-cell patch clamp) background noise by the sum of 4 exponentials (see Equations 23, 24 in Stepanyuk et al., 2011). The decaying phases of the responses (starting in 1 ms after the end of stimulation pulse) were taken for the consequent log-likelihood maximization.

In a second series of simulations we have used simple 3-state kinetic scheme of an abstract synaptic receptor. The scheme consisted of unliganded state, R, singly-ligated state, RL, and open state, O and had the following rate constants: binding rate, $k_{on} = 6 \text{mM}^{-1}\text{ms}^{-1}$, unbinding rate, $k_{off} = 0.025 \text{ms}^{-1}$, opening rate, $b = 0.25 \text{ms}^{-1}$. The closing rate constant, $a$, was 2.5 ms$^{-1}$ for Model R and Model N and 1.25 ms$^{-1}$ for Model A (see Section ML NSFA Distinguishes Between Changes in the Channel Gating and Changes in the Number of Receptors Bound with a Neurotransmitter in Results). Unitary current was set to 1 pA. Variability in the amplitude of macroscopic postsynaptic responses was achieved by trail-to-trial Gaussian variation of the number of available synaptic channels ($\text{mean} = 400; SD = 50$ for Models R and A; $\text{mean} = 800; SD = 71$ for Model N). Simulation time step was $\Delta t = 0.1$ ms. Synaptic vesicle release was modeled as a square pulse of saturating agonist concentration with a duration equal to two simulation time-steps (0.2 ms), which caused transition of all channels from R to RL state.

**PEAK-SCALED NON-STATIONARY FLUCTUATION ANALYSIS**

Accuracy of single-channel current estimates obtained with ML NSFA method presented here was compared to those obtained by PS NSFA. In PS NSFA, variance in currents arising from the stochastic gating of the ion channel is isolated from variance arising from sources such as quantal variability by scaling the mean simulated current waveform to the peak amplitude of each individual simulated current and then subtracting the two waveforms (Traynelis et al., 1993).

$$i_{\text{peak-scaled}} = I_i - \langle I \rangle \frac{\max(I)}{\max(I)}$$

To estimate the accuracy of unitary current estimates with PS NSFA, it was applied to $n = 1000$ bootstrap samples each of which contained either $N = 5, 10, 20, 30, 40, 100$ currents simulated with a 7-state GABA$_A$ receptor scheme (see Section Simulation of Macroscopic Synaptic Currents above). For each bootstrap sample the ensemble variance, $\sigma^2$, was plotted against the ensemble mean, $\langle I \rangle$, and then fitted with parabola:

$$\sigma^2 = i_{\text{ch}} \langle I \rangle - \frac{(\langle I \rangle^2)}{N_{\text{ch}}} + \sigma_0^2$$

where $\sigma_0^2$ is the variance of the background noise. Accuracy of unitary current estimates was calculated as described above, and was then compared with the accuracy of estimates obtained with ML NSFA. To ensure the best accuracy possible with PS NSFA, the ensemble mean current ($\langle I \rangle$) and variance, $\sigma^2$, were calculated for each data point and the rising phase of variance vs. mean curve was fitted with parabola using weighted (with weights $\omega_i = 1/\text{var}(\sigma_i^2)$) least squares method.

**ESTIMATION OF UNITARY CURRENT FROM A SINGLE MACROSCOPIC CURRENT**

Sampling from a likelihood distribution of model parameters that were estimated from a single macroscopic synaptic current was done by the slice sampling Markov chain Monte Carlo algorithm (Neal, 2003), implemented in "MCMC Methods for MLP and GP and Stuff" toolbox by Toni Auranen and Aki Vehtari (available at http://www.lce.hut.fi/research/compinf/mcmcsuffit/).

**RESULTS**

**ML NSFA APPLICABILITY TO ESTIMATION OF UNITARY CURRENT AND KINETIC CONSTANTS OF POSTSYNAPTIC RECEPTOR CHANNELS**

Postsynaptic architecture restricts direct electrophysiological access to individual receptors in native synaptic environments, with only occasional exceptions when channel openings and closings can be resolved on the very tail of postsynaptic currents (Silver et al., 1992). Both these exceptions and application of PS NSFA (Traynelis et al., 1993) do not allow estimating any parameters of synaptic receptors except their unitary conductance and the number of receptors open at the peak of synaptic current (Hartveit and Veruki, 2006).

In this part of the work we tested how ML NSFA estimates the unitary current and kinetic constants of GABA$_A$ receptors from stochastically simulated macroscopic currents having a trial-to-trial Gaussian variation in the number of available receptors ($N_{\text{ch}} = 250 \pm 50$). Currents were simulated with a 7-state model of this receptor (Mozrzymas et al., 2003, see Methods) having one unbound, two liganded closed, two open and two desensitized states (Figure 1A). Synaptic release of GABA was modeled as a brief (0.2 ms) step of saturating GABA concentration resembling GABA release in real synaptic connections (Perrais and Ropert, 1999, 2000; Hájos et al., 2000; Nuşser et al., 2001; Biró et al., 2006).
1000 macroscopic currents generated in response to this stimulation had the mean amplitude of $184 \pm 35\ pA$ and decay kinetics of $43.4 \pm 3.6\ ms$ (Figure 1B) and resembled postsynaptic currents routinely recorded in cortical GABAergic synapses (Nadkarni et al., 2010). Background colored noise ($SD = 3\ pA$, see Section Simulation of Macroscopic Synaptic Currents in Methods) was added to the simulated currents.

Samples consisting of $N = 5, 10, 20, 30, 40$ macroscopic currents were randomly chosen from initially generated set of 1000 currents and analyzed using ML NSFA. In order to assess the accuracy of estimates for the unitary current and kinetic rates, parameter search was performed for 60 samples obtained in such a way and log-likelihood maximization was run 10 times for each sample in order to achieve the global maximum (see Section Accuracy of the Estimates in Methods). For each run, the initial parameter values were chosen randomly and uniformly in the logarithmic scale from the range $[\theta_0/10, \theta_0 \cdot 10]$, where $\theta_0$ denotes true parameter values, i.e., those used for simulation of currents.

The unitary current was estimated with good accuracy even from samples consisting of only 10 simulated postsynaptic currents (Figure 1C, 8.1% relative error) whereas it was estimated with almost 2-fold better accuracy when the number of currents in the sample was increased from 10 to 40 (4.3% relative error). Three rate constants: unbinding rate ($k_{off}$), desensitization ($d_1$) and resensitization ($d_2$) rate from double-liganded state could also be estimated (Figures 1D–F). For samples consisting of 10 and 40 currents the relative errors of estimates were: $k_{off} — 49.0\%$ and 19.1%; $d_1 — 28.3$ and 14.6%; $d_2 — 8.9$ and 4.7%, respectively. Some of rate constants associated with single-liganded states were estimated in order of magnitude ($a_1$) or bounded from below ($\theta_1, d_1$).

Thus, we demonstrate that ML NSFA could reliably estimate the unitary current of synaptic receptor channel and several kinetic constants of synaptic receptor model from the very limited number of postsynaptic currents ($5–40$). These results indicate that ML NSFA may allow analysis of kinetic models of synaptic receptors in their native biochemical environment using routinely recorded macroscopic postsynaptic currents.

### ML NSFA Accuracy in Estimation of Unitary Current Compared to PS NSFA

The number of currents necessary for a particular algorithm to secure a given accuracy of unitary current estimate is an important practical issue. With many hundreds or even thousands of simulated macroscopic currents accuracy of PS NSFA in estimating the unitary current is fairly good (Markova et al., 2005; Hartveit and Veruki, 2006). At the same time it is hard to collect more than about 100 of evoked postsynaptic currents in steady-state conditions in routine electrophysiological experiments.

![FIGURE 1 | Estimation of unitary current and kinetic constants from simulated GABAergic synaptic currents. (A) 7-state kinetic scheme of GABA$_A$ receptor that was used to simulate macroscopic synaptic currents (Mozrzymas et al., 2003; see Section Simulation of Macroscopic Synaptic Currents in Methods). The scheme has one unbound state, $R$, two liganded states (single-liganded, RG, and double-liganded, RG2) and related open (O1 and O2) and desensitized (D1 and D2) states. Rate constants were adapted from Mozrzymas et al. (2003) and were as follows: $k_{off} = 0.13\ ms^{-1}$, $d_1 = 0.14\ ms^{-1}$, $d_2 = 1.5\ ms^{-1}$, $r_1 = 0.02\ ms^{-1}$, $r_2 = 0.12\ ms^{-1}$, $a_1 = 1.5\ ms^{-1}$, $a_2 = 1\ ms^{-1}$, $b_1 = 0.15\ ms^{-1}$, $b_2 = 8\ ms^{-1}$; $k_{on1} = 4\ mM^{-1}\ ms^{-1}$, $k_{on2} = 8\ mM^{-1}\ ms^{-1}$. Unitary currents for the states O1 and O2 were equal and were set to $n = 2 = 1\ pA$. The number of channels exposed to GABA varied from trial to trial ($N_{ch} = 250$, $SD = 50$, Gaussian variation).](#)
Thus, to see whether ML NSFA gives any advantage with respect to the number of required traces we calculated a relative error of unitary current estimates obtained with ML NSFA from the above described samples of different sizes (5, 10, 20, 30, 40, and 100 currents; 60 samples were analyzed in each case to estimate the error) and compared this error with one estimated with PS NSFA applied to 1000 samples of similar sizes.

Figure 2A demonstrates that the error of unitary current estimates obtained with both methods decreases with the number of currents taken for the analysis. However, the unitary current can be estimated with as low as 10.8, 8.1, and 4.9% relative error from only 5, 10, and 20 simulated synaptic currents, respectively, whereas PS NSFA resulted in about 2-fold lower accuracy (23.0, 14.7, and 10.4 relative error, respectively). The estimates obtained with ML NSFA from the analysis of samples of 30 and 40 currents had relative error of 4.6 and 4.3%, whereas PS NSFA gave 8.6 and 7.2% error for these cases. At last, accuracies of unitary current estimates obtained from 100 simulated currents were high for both methods and were comparable (Errors: 2.9% for ML NSFA vs. 4.5% for PS-NSFA; Figure 2A).

Thus, for some complex models ML NSFA allows evaluation of the unitary current with good accuracy using experimentally realistic number of macroscopic currents and substantially outperforms PS NSFA in terms of accuracy when only a few (5–30) postsynaptic currents are available for estimating the unitary current.

**ML NSFA ESTIMATES THE NUMBER OF SYNAPTIC RECEPTORS BOUND WITH NEUROTRANSMITTER AND PEAK OPEN PROBABILITY**

PS NSFA was specifically designed to be independent of variations in the number of postsynaptic receptors exposed to neurotransmitter and peak open probability for the sake of more accurate estimation of a unitary current (Silver et al., 1996) from postsynaptic current fluctuations. Unfortunately, this method could not be used for the estimation of the total number of receptors in the synapse. To the contrary, ML NSFA presented here allows estimation of the number of receptors bound with neurotransmitter by the end of neurotransmitter concentration transient in each macroscopic current (liganded channels, \( N_{ch} \)). It is assumed that this transient time course is known or sufficiently brief, meaning that it could be approximated by delta function in the latter case. Indeed, GABA concentration in the synaptic cleft decreases by a factor of 10 during less than 0.1 ms after synaptic vesicle release (Scimemi and Beato, 2009) resulting in almost instantaneous concentration transient. For such a brief concentration transient and for a given GABA receptor model (Figure 1A) ML NSFA would estimate the number of receptors bound with two GABA molecules by the end of concentration transient in all synapses of particular synaptic connection independently upon receptor saturation in the case when most of the current is mediated by the receptors in double-liganded states.

The open probability \( P(o) \) at any given time is determined as a mean current divided by a product \( i_{ch}N_{ch} \), and it is a function of rate constants: \( P(o) = e^{Qt}p(0) \) (see Section Estimation of the Number of Channels and Peak Open Probability in Methods). Thus, \( P(o) \) as a function of time is automatically estimated at the end of log-likelihood maximization procedure. The peak open probability is simply a maximum of this function, \( P(o_{peak}) = \max (e^{Qt}p(0)) \). Peak open probability estimated by ML NSFA is, thus far, a ratio of the number of receptors open at the peak of postsynaptic current to the number of double-liganded receptors by the end of neurotransmitter concentration transient.

![Figure 1](image-url)

**FIGURE 1** | ML NSFA is more accurate than PS NSFA in estimating of unitary current. Estimation of the number of receptors bound with a neurotransmitter and peak open probability with ML NSFA. (A) Statistical plots demonstrating accuracy of unitary current estimates obtained with ML NSFA (blue boxes) and PS NSFA (black boxes) from simulated macroscopic synaptic currents with trial-to-trial Gaussian variation in the number of receptors \( N_{ch} = 250, SD = 50 \); see kinetic scheme in Figure 1A. On each plot, the central mark (red) is the median, the edges of the box are the 25th and 75th percentiles, the whiskers extend to the most extreme data points not considered outliers, and outliers are plotted individually by red crosses. Green line indicates a true value of unitary current. ML NSFA and PS NSFA were performed using \( n = 60 \) and \( n = 1000 \) samples consisting of \( N = 5, 10, 20, 30, 40 \) or 100 simulated currents, respectively. Note that the accuracy of estimates obtained with ML NSFA using a few (5–20) currents was 2-times better than one obtained with PS NSFA. (B) An example of variance vs. mean plot (gray dots) obtained with PS NSFA for \( N = 30 \) simulated macroscopic currents having trial-to-trial Gaussian variation in the number of receptors \( N_{ch} = 250, SD = 50 \) and a parabolic fit of its rising phase (red). Note that variance-mean relationship (gray dots) is skewed rather than parabolic and therefore the number of receptors could not be estimated with PS NSFA. (C,D) Statistical plots for the estimates of the number of channels bound with a neurotransmitter right after the concentration transient, \( N_{ch} \), and peak open probability, \( P(O_{peak}) \) obtained with ML NSFA. Green line in (C) indicated the true value of the number of channels estimated as mean peak current amplitude (averaged over \( N = 1000 \) currents) divided by the true value of \( P(o_{peak}) \) and by the true value of unitary current (1 pA) and in (D) green line indicates the true value of \( P(o_{peak}) \) estimated as \( P(o) = e^{Qt}p(0) \). Other colors and notations are the same as in Figure 1C.
Figures 2C,D demonstrate that the error of \(N_{ch}\) and \(P(\theta, \text{peak})\) estimates obtained with ML NSFA decreases with the number of currents taken for the analysis. The number of liganded receptor channels, \(N_{ch}\), was calculated as an average over all currents in the sample and was estimated with 24.5 and 12.4% relative error from samples consisting of only 5 and 10 simulated macroscopic synaptic currents, respectively. The respective estimates of accuracy for the peak open probability, \(P(\theta, \text{peak})\), had 14.4 and 9.8% relative error, respectively. Both \(N_{ch}\) and \(P(\theta, \text{peak})\) were estimated with even better accuracy from samples consisting of 100 simulated currents (10.0 and 4.3% relative error, respectively).

At the same time PS NSFA applied to the same samples resulted in a variance vs. mean curve that was profoundly skewed (Figure 2B, gray dots) and, therefore, could not give an estimate of the number of liganded channels, \(N_{ch}\).

**ESTIMATION OF UNITARY CURRENT AND KINETIC CONSTANTS OF RECEPTORS HAVING MULTIPLE CONDUCTANCE LEVELS**

Most ligand-gated channels are described by kinetic schemes with multiple, non-identical open states often having different conductance levels (Jin et al., 2003; Mozrzymas et al., 2003; Robert and Howe, 2003; Wyllie et al., 2006; Keramidas and Harrison, 2010; Mortensen et al., 2010). In practice some open states should be considered rare and excluded from the fitting of experimental results in order to estimate at least some parameters of receptor kinetic schemes (Mortensen et al., 2010). Unfortunately, PS NSFA is also not applicable to examination of receptors having multiple conductance levels giving values of unitary current and channel number having no obvious physical interpretation (Hartveit and Veruki, 2006). Thus, at the present moment single-channel recordings are virtually the only approach that allows identifying multiple conductance levels of ligand-gated receptors and this approach is also not applicable for studying of synaptic receptors.

We next wanted to investigate if ML NSFA suggested in this work is applicable to analysis of ion channels and ligand-gated receptors with multiple conductance levels, described by kinetic schemes with non-identical open states. 7-state kinetic model of GABA\(_A\) receptor (Mozrzymas et al., 2003) having two open states \(O1\) and \(O2\) with identical unitary current \((i1 = i2, \text{see Figure 1A})\) was modified to have the unitary current \(i1 = 2\) pA and \(i2 = 1\) pA for the states \(O1\) and \(O2\), respectively (Figure 3). Rate constants of the model were modified in such a way that the contribution of single- and double-ligated open states to the total macroscopic current became comparable. Modified constants were (in ms\(^{-1}\)): \(b2 = 4, b1 = 1.2, d1 = 1, r1 = 1, d2 = 0.15, r2 = 1\). Colored NSFA were applied to samples of 50 macroscopic currents (\(n = 15\) and \(n = 250\) bootstrap samples for MS NSFA and PS NSFA, respectively) simulated as described in (A) and having true values of \(n_{O1} = 2\) pA and \(n_{O2} = 1\) pA, respectively (indicated by green lines). On each plot, the central mark (red) is the median, the edges of the box are the 25th and 75th percentiles, the whiskers extend to the most extreme data points not considered outliers, and outliers are plotted individually by red crosses. Note that ML NSFA accurately distinguishes both unitary current levels, whereas PS NSFA gave some value of the unitary current that was close to \(i2\). (C) Statistical plot for the estimates of kinetic rates of transitions from and to a single-ligated state obtained by ML NSFA (in ms\(^{-1}\)): unbinding rate, \(k_{off}\) = 0.13 ± 0.01, desensitization rate, \(d1 = 0.89 ± 0.34\), re-sensitization rate \(r1 = 1.02 ± 0.08\), closing rate, \(a1 = 1.55 ± 0.06\), opening rate, \(b1 = 1.17 ± 0.24\); \(N = 50\) currents simulated as described in (A). The estimates were in good agreement with their true values (green lines). See a legend to panel (B) for further description.

**FIGURE 3** | Estimation of unitary currents and kinetic constants of receptors having two open states with different conductance levels. (A) Upper panel. Example of 50 synaptic currents simulated with a 7-state kinetic scheme of GABA\(_A\) receptor having two open states (Figure 1A, some rate constants were modified: \(b2 = 4, b1 = 1.2, d1 = 1, r1 = 1, d2 = 0.15, r2 = 1\)). Unitary currents were set to \(i1 = 2\) pA and \(i2 = 1\) pA for open states \(O1\) and \(O2\), respectively. The number of channels varied from trial to trial \((N_{ch} = 500 ± 50\); Gaussian variation). Lower panel. Representative example of single simulated macroscopic current components mediated by single-ligated open state \(O1\) (blue trace) and double-ligated open state \(O2\) (green trace) demonstrating comparable contribution of \(O1\) and \(O2\) to the total macroscopic current. (B) Statistical plots for the estimates of unitary currents obtained with PS NSFA (leftmost bar, \(i = 1.86 ± 0.03\) pA) and ML NSFA (two bars on the right; \(i1 = 2.0 ± 0.11\) pA and \(i2 = 0.89 ± 0.08\) pA). (C) Statistical plot for the estimates of kinetic rates of transitions from and to a single-ligated state obtained by ML NSFA (in ms\(^{-1}\)): unbinding rate, \(k_{off}\) = 0.13 ± 0.01, desensitization rate, \(d1 = 0.89 ± 0.34\), re-sensitization rate \(r1 = 1.02 ± 0.08\), closing rate, \(a1 = 1.55 ± 0.06\), opening rate, \(b1 = 1.17 ± 0.24\); \(N = 50\) currents simulated as described in (A). The estimates were in good agreement with their true values (green lines). See a legend to panel (B) for further description.
background noise with \( SD = 3 \) pA was added to the simulated currents (Figure 3A, upper panel).

Representative examples of the simulated current components associated with either state O1 or state O2 are shown in Figure 3A, lower panel, by blue and green lines, respectively. When 250 samples consisting of \( N = 50 \) simulated currents (Figure 3A, upper panel) were analyzed by PS NSFA the unitary current estimates were close to the unitary current of single-ligated open state O1 (1.86 ± 0.03 pA vs \( i_1 = 2 \) pA for the state O1). At the same time, ML NSFA gave reasonable estimates for both conductance levels (Mean ± SE \( i_1 = 2.00 ± 0.11 \) pA and \( i_2 = 0.89 ± 0.08 \) pA; \( n = 15 \) samples of \( N = 50 \) simulated currents; Figure 3B). ML NSFA also reliably estimated kinetic rates for single-ligated state transitions (\( k_{\text{off}} = 0.13 ± 0.01 \), \( d_1 = 0.89 ± 0.34 \), \( r_1 = 1.02 ± 0.08 \), \( a_1 = 1.55 ± 0.05 \), \( b_1 = 1.17 ± 0.24 \) ms\(^{-1} \), Figure 3C) and the mean number of liganded channels (\( N_{\text{hl}} = 557 ± 53 \) vs. 500 ± 50 used in simulation).

Thus, contrary to PS NSFA, ML NSFA can reliably estimate kinetic schemes with several open states having different conductance levels and gives precise values of unitary currents, some kinetic rates, and the mean number of liganded receptors in a given synaptic connection.

ML NSFA distinguishes between changes in the channel gating and changes in the number of receptors bound with a neurotransmitter

We next attempted to explore ML NSFA capability to identify which postsynaptic parameters were changed in the case when mean amplitude of simulated currents was increased without changes in macroscopic current waveform and unitary current.

To this end three distinct groups of 1000 macroscopic currents were generated using a simple 3-state scheme of synaptic channel (Figure 4A, see Section Simulation of Macroscopic Synaptic Currents in Methods). A similar increase in mean current amplitude was achieved by changes in either receptor gating or receptor number. A reference kinetic scheme (Model R; Figure 4A, red) had the closing rate, \( a = 2.5 \) ms\(^{-1} \) and the total number of channels \( N_{\text{hl}} = 400 ± 50 \) and was used to generate a group of macroscopic currents before putative remodeling of synaptic connection (Figure 4B). In the second kinetic scheme (Model A; Figure 4A, blue) mimicking remodeling of receptor gating the closing rate, \( a \), was changed from 2.5 ms\(^{-1} \) to 1.25 ms\(^{-1} \) resulting in almost 2-fold increase of average current amplitude (Figure 4C, blue) without substantial changes in current waveform (Figure 4D, blue vs. red). Conversely, in the third model (Model N; Figure 4A, black) the number of available channels, \( N_{\text{hl}} \), was increased from 400 ± 50 to 800 ± 71 without any changes in the kinetic constants, which led to similar changes in current amplitude (Figure 4C, black) as for Model A without any changes in current waveform (Figure 4D, black vs. red). Therefore, currents generated with Models A and N had similar amplitudes and when normalized, appeared to have the same waveforms as reference currents generated by Model R (Figures 4B–D).

ML NSFA was run with \( n = 20 \) bootstrap samples consisting of \( N = 100 \) currents (see Section Accuracy of the Estimates in Methods) for each of the 3 groups of simulated currents in order to evaluate the receptor model parameters and the respective errors. Log-likelihood maximization was run 5 times for each bootstrap sample in order to achieve the global maximum. When the parameter estimates obtained from currents generated with Model R were compared to those obtained from currents generated with Model A (Figure 4E, red vs. blue boxes) the difference, \( \Delta_{\text{RA}} \), between mean values of each parameter estimates except the closing rate, \( a \), and peak open probability, \( P(o, \text{peak}) \), was small and was within the standard error (SE) range of the respective estimates: \( k_{\text{off}}: \Delta_{\text{RA}} = 1.7\% (SE = 2.2\%) \), \( b: 8.5\% (13.0\%) \), \( i_{\text{ch}}: 0.6\%(2.4\%) \), \( N_{\text{hl}}: 8.8\% (14.0\%) \). At the same time, \( \Delta_{\text{RN}} \) was 49.7\% for the closing rate, \( a \) and 70.1\% for the peak open probability, \( P(o, \text{peak}) \) and did not fall within the narrow ranges of the respective SE's (2.4\% and 14.1\%, respectively). The mean values of the respective estimates were \( a = 2.51 ± 0.04 \) for Model R and 1.26 ± 0.03 for Model A, \( P(o, \text{peak}) = 0.08 ± 0.01 \) for Model R and 0.14 ± 0.02 for Model A. Therefore, we could infer that these were the parameters that altered. These results directly indicate that ML NSFA may reliably determine changes in receptor gating, which leads to an increase in peak open probability.

When estimates obtained from currents generated with Model R and Model N were compared, we observed insufficient differences, \( \Delta_{\text{RN}} \), between mean values of all parameter estimates except the number of receptors, \( N_{\text{hl}} \), which was changed from 419 ± 62 for Model R to 942 ± 228 for Model N (Figure 4E, compare red vs black boxes). \( \Delta_{\text{RN}} \) for \( N_{\text{hl}} \) was 124.9\% and did not fall within the range of its SE (24.2\%). At the same time, \( \Delta_{\text{RN}} \) for other parameters fell within the respective standard error (SE) range: \( k_{\text{off}}: \Delta_{\text{RN}} = 0.3\% (SE = 1.7\%) \), \( a: 1.3\% (1.7\%) \), \( b: 7.1\% (18.0\%) \), \( i_{\text{ch}}: 0.8\%(1.6\%) \), \( P(o, \text{peak}): 8.3\% (18.9\%) \) and it was possible to conclude that the number of receptors was the only altered parameter in this case.

Thus, with ML NSFA it becomes possible to distinguish between alteration in receptor channel gating and receptor number, which nonetheless resulted in visually indistinguishable postsynaptic currents.

Estimation of unitary current from macroscopic currents generated by receptors having different kinetic schemes

The key assumption of the PS NSFA is that all receptors in a particular synaptic connection under study have identical kinetic properties (Silver et al., 1996). As a result, all variance in the currents could be attributed to the stochastic nature of the channel gating rather than to the variability in their kinetics. In fact, this assumption could be violated since receptors in the synaptic connection could have different subunit composition or could be differentially modulated (Popescu and Auerbach, 2003) and a set of receptors contributing to each postsynaptic current could vary from trial to trial. In this case PS NSFA overestimates the unitary current and this overestimation could be quite significant even if the difference between receptor kinetic rates is so small that it could be hardly noticed from the observation of synaptic currents (see Figure 5A and below).

Using likelihood approximation it is possible in principle to estimate unitary current and other parameters independently for each individual synaptic current. To test this possibility we
have conducted a series of computational experiments. A group of 1000 synaptic currents was simulated using 7-state kinetic scheme of GABA<sub>A</sub> receptor channel (Mozrzymas et al., 2003, see scheme in Figure 1A) and the other 1000 currents were simulated using similar scheme in which several parameters (closing rate, $k_{\text{off}}$, desensitization rate, $d_2$, and resensitization rate, $r_2$) varied between trials randomly and uniformly in the range of ±20% of parameter values that were used to generate the first group of currents. In both cases the unitary current was set at 1 pA and colored background noise ($SD \pm 3$ pA) was added to the simulated currents (see Section Simulation of Macroscopic Synaptic Currents in Methods for details).

Figure 5A demonstrates that both groups of currents had similar waveforms and their decay times were almost identical although variability of decay times in the second group was slightly higher (Mean ± SD: 43.6 ± 3.7 ms vs. 43.9 ± 6.1 ms, $N = 1000$ currents). Nevertheless, variance vs. mean curves for these two groups of currents differed significantly (Figure 5B) and for the second group unitary current appeared to be 1.9-fold overestimated by PS NSFA (Mean ± SE was 1.01 ± 0.03 pA for the group of currents without variation of parameters vs 1.92 ± 0.05 pA for the group of currents with variation of $k_{\text{off}}$, $d_2$, and $r_2$; $N = 250$ currents; true value was 1 pA).

To the contrary, when ML NSFA was applied to the group of currents with varying rate constants and log-likelihood of each
current in the group was optimized independently, a reasonably accurate estimate of unitary current was obtained (Mean ± SD = 0.89 ± 0.23 pA, N = 50 currents). Standard error of mean unitary current estimate was very low (SE = 0.053 pA), but bias from the true value (1 pA) was significant. We have noticed that the cause of this bias is the skewed shape of the likelihood distribution of a single simulated synaptic current, which means that for the case of single current the maximum likelihood value of unitary current is not the most common value. An example of the typical distribution of unitary current obtained by sampling from the likelihood distribution for a single simulated macroscopic current using the slice sampling Markov chain Monte Carlo method (MCMC, 2000 samples) is shown in Figure 5C (upper panel). It can be seen that the distribution maximum significantly differs from the distribution mean (red vertical line). Therefore, in order to obtain "typical" values of unitary current, mean values of unitary current were also estimated by slice sampling from the likelihood distributions obtained for individual synaptic currents (1000 MCMC samples for each current) from the same group of 50 currents. The resulting distribution of unitary current estimates obtained by accumulation of all 50 distributions for individual currents is represented in Figure 5C (lower panel). The final estimate of unitary current was obtained by averaging over N = 50 mean unitary currents and was in perfect agreement with its true value (Mean ± SD = 0.97 ± 0.39 pA, red vertical line in Figure 5C, lower panel; SE = 0.056 pA). Figure 5D shows statistics of the mean unitary current estimates obtained with MCMC applied to likelihood distributions of individual currents (left box, N = 50) in comparison with the same statistics obtained with PS NSFA applied to individual currents as described above (right box, N = 250, n = 50 bootstraps). It is clearly seen that, contrary to MCMC, PS NSFA significantly overestimates unitary current (green line indicates true value, 1 pA). Among the other model parameters only the number of liganded channels, N_{ch}, and the desensitization rate, r_{2,}, were estimated with MCMC with relatively high accuracy. The desensitization rate, d_{2}, and GABA unbinding rate, k_{off}, were estimated in order of magnitude. The median of the absolute difference between estimates of model parameters and their true values for k_{off}, d_{2}, r_{2,}, i_{ch}, and N_{ch} was 191, 188, 22, 31, and 35% of their true values, respectively.

We conclude that the mean values for several parameters of the synaptic receptor model, such as the unitary current, the number
of channels and the peak open probability, can be estimated with a reasonable accuracy using ML NSFA or MCMC sampling from the likelihood distribution of each individual current in the group of currents even if these currents were mediated by receptors having different kinetic models.

**DISCUSSION**

In this study we have further developed a new maximum likelihood method that we suggested earlier (Stepanyuk et al., 2011) and applied it to analysis of simulated macroscopic currents, in which the number of receptors exposed to a neurotransmitter varied from trial to trial. In the newly developed method, ML NSFA, the number of liganded receptors was first optimized for each macroscopic current and then these estimates were used to maximize the log-likelihood in order to obtain a set of kinetic model parameters as it was described earlier (Stepanyuk et al., 2011).

We explored the performance of ML NSFA with several different kinetic schemes of varying complexity and varying conditions relevant for real synaptic transmission. It was shown that contrary to PS NSFA (Traynelis et al., 1993) ML NSFA could estimate not only the unitary current of synaptic receptor channel but also multiple conductance levels, the number of liganded receptors, peak open probability and some kinetic constants from the experimentally realistic number of simulated postsynaptic currents. We have also evaluated the accuracy of ML NSFA compared to PS NSFA with respect to estimating the unitary current and found it 2-fold more accurate for a few (5–30) macroscopic currents. ML NSFA estimation of the unitary current was robust even when currents were generated by receptors having different kinetic parameters, the case when PS NSFA is not applicable. Thus, our results demonstrate that ML NSFA that takes into account correlations between different time points of a macroscopic currents and computationally scales linearly with the number of channel states (Stepanyuk et al., 2011) quantitatively and qualitatively outperforms currently available approaches for analysis of kinetic schemes of synaptic receptors.

**ML NSFA APPLICABILITY TO ANALYSIS OF SYNAPTIC RECEPTOR PROPERTIES**

Noise analysis of macroscopic currents remains a useful tool for determining the properties of different ligand- and voltage-operated channels (Traynelis and Jaramillo, 1998). Moreover, PS NSFA, the most frequently used noise analysis approach, is the only approach that can be applied to analysis of channels with an unusually low unitary conductance (Swanson et al., 1997) and receptor channels localized at synapses (Traynelis and Jaramillo, 1998). At the same time the unitary current is virtually the only parameter that can be reliably obtained from this type of analysis (Traynelis et al., 1993; Silver et al., 1996). To the best of our knowledge, kinetic rates have never been estimated for any synaptic receptors in their intrinsic environment. Peak open probability of receptors and the number of receptors bound with a neurotransmitter could not be also directly analyzed by any current approach. Possibility to estimate the unitary current and some kinetic rates using a few simulated postsynaptic currents demonstrated in this study allows for the first time to follow a time course of receptor remodeling in one and the same synaptic connection. Having in mind that estimation of some receptor parameters with accuracy of 10% can be obtained from 10 macroscopic currents (Figures 1, 2), which can be collected in routine electrophysiological experiments for about 30 s, dynamics of receptor remodeling can be followed with a time course of several measurements per minute. It can be, for example, used for studying of modal gating, which refers to low probability rearrangements in receptor structure producing a substantial change in the overall pattern of channel opening (Popescu, 2012). Modal switches can be observed in single channel recordings of most ionotropic ligand-gated channels (Popescu, 2012) but it has never been directly demonstrated for synaptic receptors located in their intrinsic environment in a response to synaptic release of neurotransmitter. Modal gating may result not only in the different unitary conductance of receptors but also in changes in their gating and peak open probability (Popescu, 2005; Lema and Auerbach, 2006; Zhang et al., 2008; Poon et al., 2010; Prieto and Wollmuth, 2010). Moreover, in many cases, especially for the instance of NMDA receptors, substantial changes in gating, and peak open probability is observed without changes in the unitary conductance (Popescu, 2005; Zhang et al., 2008). Thus, such remodeling of synaptic receptors cannot be, in general, revealed by PS NSFA, while ML NSFA should certainly uncover it due to intrinsic ability to estimate some kinetic constants and peak open probability (Figures 1, 2). The modal gating is slow (Popescu, 2005; Zhang et al., 2008; >5 min) and agonist- and stimulus-sensitive (Armstrong and Gouaux, 2000; Poon et al., 2010). Thus, it looks potentially plausible to synchronize synaptic receptor switching between different modes for a set of synaptic receptors in a given synaptic connection and to study the modal gating of synaptic receptors in their intrinsic environment by means of ML NSFA. For example, multiple conductance levels observed in modal gating of GluA2 AMPA receptors (Prieto and Wollmuth, 2010) or different open channel probabilities found for the type 2A isoform of NMDA receptors (Popescu and Auerbach, 2003) can be resolved from the respective postsynaptic currents (Figures 2–4).

Moreover, different types of AMPA receptor regulation that occur during LTP or LTD expression, such as changes in receptor trafficking (Huganir and Nicoll, 2013), in interaction of AMPARs with auxiliary subunits (Khodosevich et al., 2014) or adapter proteins that could lead to changes in receptor kinetics (Studniarczyk et al., 2013), phosphorylation-evoked changes in unitary current and peak open probability (Traynelis and Wahl, 1997; Derkach, 2003) could be potentially resolved with ML NSFA applied to the respective postsynaptic currents. Studies of developmental, pathological, plastic, and tissue specific modifications of synaptic receptors (Kittler et al., 2004; Lüthi et al., 2004; Palmer, 2006; Stubblefield and Benke, 2010) including changes in receptor subunit composition and trafficking (Ruiz et al., 2005; Patten and Ali, 2007) that have been earlier analyzed by PS NSFA may now also obtain a second wind due to a possibility to evaluate many parameters of the respective synaptic receptors.

Conclusions about mechanisms of synaptic receptors modulation that are based solely on the analysis of the amplitude of postsynaptic currents or unitary current might be misleading.
Indeed, stable unitary conductance might be accompanied by changes in receptor gating that may lead to an increase in the total charge transferred via a single synaptic receptor (Figure 4). At the level of macroscopic current it would result in an increase of current amplitude without substantial changes of its waveform (Figure 4). Together with absence of changes in the unitary conductance reported by PS NSFA it would be interpreted as presynaptic modification or an increase in the number of postsynaptic receptors. At the same time ML NSFA would certainly reveal changes in postsynaptic receptor gating.

The new approach also allows separate estimation of kinetic parameters of synaptic and extrasynaptic receptors expressed in the same neuron. For that, a set of postsynaptic currents necessary for evaluation of synaptic receptor model parameters must be initially recorded. Then strong presynaptic stimulation that can activate the whole set of synaptic terminals innervating the neuron under study should be performed in the presence of an irreversible use-dependent inhibitor of the respective synaptic receptors (e.g., picrotoxin for GABA_A (Olsen, 2006) or MK-801 for NMDA (McAllister and Stevens, 2000) receptors, respectively). Next, several different agonist concentrations should be sequentially applied to the preparation in order to activate the extrasynaptic receptors and to record the respective transmembrane currents. Analysis of these macroscopic currents by ML NSFA or some of previously developed approaches (Milescu et al., 2005; Moffatt, 2007; Stepanyuk et al., 2011) would give kinetic parameters of extrasynaptic receptors.

**ML NSFA applicability to analysis of synaptic receptor number and peak open probability**

PS NSFA provides only an estimate of unitary current (Traynelis et al., 1993). In spite of this, estimation of N_ch and P(o, peak) was performed for single mossy fiber synapses of hippocampal granule cells having saturating glutamate concentration induced by synaptic vesicles release (Silver et al., 1996). In this case variance due to quantal variability is negligible and conventional NSFA can estimate these parameters. Although saturation of postsynaptic receptors is not rare in central synapses (Auger and Marty, 1997; Perras and Ropert, 1999, 2000; Hájos et al., 2000; Nusser et al., 2001; Biró et al., 2006) estimation of N_ch and P(o, peak) could not be performed for the synaptic connections with multiple release sites by conventional NSFA due to trial-to-trial variability in the number of released vesicles and, as a result, in the number of receptors exposed to neurotransmitter. Moreover, in most of the central synapses neurotransmitter does not saturate postsynaptic receptors making all current methods void in determining N_ch and P(o, peak). On the other hand ML NSFA suggested in this study can directly evaluate the number of receptors, N_ch, bound with neurotransmitter by the end of fast transient of neurotransmitter concentration in a synaptic cleft and P(o, peak) defined as a fraction of liganded receptors N_ch, opened at the peak of macroscopic current (Figure 2). Moreover, N_ch could be separately evaluated for each postsynaptic current (Equation 18) and open probability as a function of time, which, in particular, includes P(o, peak) (Figure 2) could be obtained from estimated kinetic rate constants (Figures 1, 3, 5). Assumptions underlying ML NSFA suggest that estimations of kinetic rates as well as N_ch and P(o, peak) are correct if all synaptic receptors are subjected to the same and fast neurotransmitter profile or if the receptors are saturated. For some kinetic schemes (Figure 1A) saturation or the same concentration profile for all receptors are not obligatory and fast (compared to some kinetic rates) neurotransmitter profile is the only necessary assumption for ML NSFA applicability.

ML NSFA might be generally applicable to studies of synaptic and extrasynaptic NMDA receptors, glutamate receptors that directly contribute to active properties of dendrites. In the case of synaptic AMPA and NMDA receptors the ability of ML NSFA to analyze currents with variable kinetics could be important due to significant variability of glutamate transients in the excitatory synapses, low saturation levels of both receptor types (McAllister and Stevens, 2000) and complexity of their kinetic schemes (Popescu and Auerbach, 2004).

In conclusion we would like to note that more accurate estimation of unitary current compared to PS NSFA together with possibilities to distinguish multiple conductance levels and evaluate the number of liganded receptors, peak open probability and some kinetic constants position ML NSFA as a powerful tool to study synaptic receptor properties in their native environment using experimentally recorded postsynaptic macroscopic currents.

**Acknowledgments**

This work was supported by NASU Biotechnology and Functional Genomics and Metabolomics Grants and DFFD F46.2/001 and F47/066 Grants.

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 30 June 2014; paper pending published: 10 August 2014; accepted: 09 September 2014; published online: 02 October 2014.

Citation: Stepanyuk A, Borisyk A and Belan P (2014) Maximum likelihood estimation of biophysical parameters of synaptic receptors from macroscopic currents. Front. Cell. Neurosci. 8:303. doi: 10.3389/fncel.2014.00303

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