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Abstract. Among electrophysiological signals, Local Field Potentials (LFPs) are extensively used to study brain activity, either in vivo or in vitro. LFPs are recorded with extracellular electrodes implanted in brain tissue. They reflect intermingled excitatory and inhibitory processes in neuronal assemblies. In cortical structures, LFPs mainly originate from the summation of post-synaptic potentials (PSPs), either excitatory (ePSPs) or inhibitory (iPSPs) generated at the level of pyramidal cells. The challenging issue, addressed in this paper, is to estimate, from a single extracellularlyrecorded signal, both ePSP and iPSP components of the LFP. The proposed method is based on a model-based reverse engineering approach in which the measured LFP is fed into a physiologically-grounded neural mass model (mesoscopic level) to estimate the synaptic activity of a sub-population of pyramidal cells interacting with local GABAergic interneurons. The method was first validated using simulated LFPs for which excitatory and inhibitory components are known a priori and can thus serve as a ground truth. It was then evaluated on in vivo data (PTZ-induced seizures, rat; PTZinduced excitability increase, mouse; epileptiform discharges, mouse) and on in clinico data (human seizures recorded with depth-EEG electrodes). Under these various conditions, results showed that the proposed reverse engineering method provides a reliable estimation of the average excitatory and inhibitory post-synaptic potentials originating of the measured LFPs. They also indicated that the method allows for monitoring of the excitation/inhibition ratio. The method has potential for multiple applications in neuroscience, typically when a dynamical tracking of local excitability changes is required.

1. Introduction

Local field potentials (LFPs) refer to extracellularly-recorded electrophysiological signals generated by neuron assemblies (Buzsáki 2004). LFPs are extensively used to monitor and analyze brain function in brain research as well as in clinical studies in neurology. LFP recording requires a simple differential amplification coupled with a filtering montage and a surgical procedure to implant electrodes in or close to the brain regions

of interest. Therefore, the LFP provides critical information on local neuronal activity. LFPs are used in a wide spectrum of applications, from basic neuroscience to clinics (Destexhe & Bedard 2013) (e.g., using the LFP as a control signal to trigger stimulation in closed-loop (Priori et al. 2013), which requires a more sophisticated hardware than a standard LFP recording setup, due to the stimulation montage and the control processing).

Physiologically, LFP signals reflect ongoing excitation- and inhibition-related processes in recorded networks (Buzsáki et al. 2012). Indeed, it is known that the major contribution to LFPs arises from post-synaptic potentials (PSPs) generated at the level of pyramidal cells within spatially extended neuronal assemblies. As described by bioelectromagnetic models (Malmivuo et al. 1995, Ramachandran 2002) and experimental studies (Buzsáki et al. 2012), the summation of PSPs, either excitatory (ePSPs) or inhibitory (iPSPs) generated at the level of pyramidal cells located in the cerebral cortex is the major contribution to LFPs. This PSP summation is explained by two major factors: (1) synaptic activation leads to the formation of a sink and a source at the level of neurons, which can then be viewed as elementary current dipoles (Buzsáki et al. 2012), and (2) when neurons are geometrically aligned (Buzsáki et al. 2012, Harris et al. 2003) (such as pyramidal cells organized "in palissade" in cortical structures), then dipole contributions tend to sum up instead of cancelling out (Buzsáki et al. 2012). In addition, due to the frequency-filtering properties of cerebral tissue, slow post-synaptic currents are more likely to be recorded further away from the electrode contacts, as opposed to fast transmembrane ionic currents (involved in action potentials) which attenuate much faster with space (Bédard et al. 2004). Overall, LFPs mainly result from an averaging process of synaptic currents, both excitatory (glutamatergic) and inhibitory (GABAergic). Therefore, LFPs may, in theory, give access to underlying synaptic currents and, subsequently, provide insights about the excitability level of recorded neuronal local networks. To reach this goal, the challenging issue is to estimate from a single extracellularly-recorded signal both ePSP and iPSP components of the LFP. In this paper, we propose a novel model-based approach to solve this problem. The method is based on model-based reverse engineering approach in which a neural mass model (NMM) is used to deconstruct the extracellularly-recorded field activity and reveal its main excitatory and inhibitory components. This reverse engineering method involves two main steps. First, the classical NMM was revisited to solve an ill-posed problem, since two parameters (ePSPs and iPSPs amplitude) are estimated from a single signal (LFP recording) measured over a finite time window. Second, an optimization procedure was developed allowing for time tracking of ePSPs and iPSPs over a shortduration sliding window. From a theoretical viewpoint, our approach makes use of the neural mass model structure as a constraint to solve an ill-posed problem, i.e. the identification of two parameters (namely EXC and INH) from one signal measurement only.

As demonstrated from both simulated and experimental data, the proposed approach can reliably estimate and monitor the average post-synaptic potentials and the densities Reconstruction of post-synaptic potentials by reverse modeling of local field potentials 3 of action potentials (AP) arising from underlying excitatory and inhibitory subsets of neurons.

It is worth noting that several methods have been proposed to quantify the balance between excitation and inhibition (excitation to inhibition ratio, EIR). For instance, a study used the slope of the power spectrum at low-frequencies (Gao et al. 2017) and identified that the latter was significantly associated with changes in the EIR. Despite its interest in terms of understanding how excitability shapes subtle features of the LFP power spectrum, this approach relies mainly of the power spectrum slope at low frequencies to estimate the EIR, and excitatory/inhibitory components were indirectly estimated through a multivariate regression model. Another study attempted to relate the EIR with the transfer function of a neural mass model, (Moran et al. 2007), based on an approximation using a linearized version of the system equation, without accounting for fine dynamics due to the hypothesis of stationarity. Our method overcomes these difficulties and provides a direct access to quantified indexes such as post-synaptic potentials (excitatory/inhibitory, from which the EIR can be derived). Therefore, the proposed method can be utilized in a wide range of applications that make use of mesoscopic LFPs, from basic neuroscience (tracking of excitability changes in neuronal networks) to clinics (analysis of depth-EEG recordings in patients with epilepsy).

2. Materials and methods

2.1. Principle of the model-based LFP reconstruction

The method exploits a priori knowledge about the processes (type and kinetics) underlying LFP generation in order to constrain LFP decomposition. LFPs are mostly composed of two sub-components (excitatory and inhibitory), and their extraction is challenging since 1) reconstructing two signals from only one is by definition an ill-posed problem; and 2) their frequency range is similar. In order to overcome this roadblock, we used a physiologically relevant NMM as a constraint to enable this decomposition process. NMM are a well-established computational class of models of neuronal population activity considering synaptic interactions between the neuronal assemblies involved in each LFP component: pyramidal cells (excitation) and interneurons (inhibition). We then used an LFP (experimental or simulated) to constrain the NMM response, and exploited this NMM to extract information contained in the LFP, especially to reconstruct excitatory and inhibitory processes. p-values were calculated using the Welch's t-test, suited for unequal variances and sample sizes.

2.2. Revised version of the neural mass model (NMM)

The NMM used in this study is an established and validated neural population model (Jansen & Rit 1995) including two sub-populations corresponding to pyramidal cells for excitation, and interneurons for inhibition. The corresponding block diagram of the

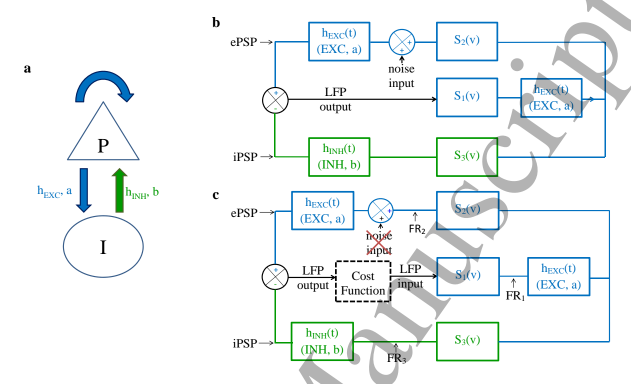


Figure 1. Illustration of the reverse modeling approach. (a) Neural mass model block diagram including two sub-populations: pyramidal cells (P) and interneurons (I). Excitatory (blue) and inhibitory (green) links between sub-populations are represented with arrows. (b) Neural mass model diagram where sigmoid functions were modified and connectivity parameters from the original NMM were removed. Excitatory and inhibitory loops are represented respectively in blue and green. LFP is the summation of synaptic inputs onto the pyramidal population (c) Reverse modeling approach where the new neural mass model is used to estimate PSPs contained in an LFP signal, the latter being set as an input of the model. A cost function based on the RMS error between estimated $\widehat{\text{LFP}}$ (LFP output) and input LFP was introduced to improve identification of EXC (excitatory) and INH (inhibitory) gain parameters. ePSPs: excitatory post-synaptic potentials, iPSPs: inhibitory post-synaptic potentials, h: synaptic impulse response functions, S_n : sigmoid functions, 1/a and 1/b: dendritic average time constant.

model is presented in figure 1, where excitatory/inhibitory processes are represented in blue/green, respectively. In order to improve the physiological relevance of the NMM, we adapted the parameters of the "wave-to-pulse" sigmoid functions, converting the mean depolarization of sub-populations into an output firing rate, according to available neurophysiological data from the literature. We also reduced the number of parameters to be identified through the reverse modeling process by removing the connectivity parameters, which were instead integrated in the new sigmoid functions and in the synaptic gain parameters of the transfer functions. The main difference with the original Jansen-Rit's model (Jansen & Rit 1995) is therefore the absence of connectivity parameters. Table 2 summarizes model parameters values. The resulting equations were derived following the method presented by Touboul et al. (Touboul et al. 2011). The synaptic impulse response functions used in the model were under the

Pyramidal cells

References	Sigmoid function parameters		
	e0	r	v0
Reid et al. (Reid et al. 2014)	48.9	0.534	5.12
Chen et al. (Chen & Bazan 2005)	26.7	0.721	5.81
Malik et al. (Malik & Chattarji 2011)	20.5	0.435	5.81
Malik et al. (Malik & Chattarji 2011)	112	0.356	6.62
Wang et al. (Wang et al. 2015)	19.1	0.550	6.67
Mean	45.4	0.519	6.00

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References	Sigmoi	d function	parameters		
	e0	r	v0		
Gibson et al. (Gibson et al. 2006)	173	0.166	17.7		
Ferguson et al. (Ferguson et al. 2013)	128	0.239	12.0		
Fidzinski et al. (Fidzinski et al. 2015)	195	0.320	11.1		
Fidzinski et al. (Fidzinski et al. 2015)	154	0.210	13.1		
Erisir et al. (Erisir et al. 1999)	66.3	0.376	10.6		
Mean	143	0.262	12.9		

Table 1. Identified sigmoid function parameters.

standard form: $h_{EXC}(t) = EXC \cdot t \cdot e^{\frac{-t}{\tau_e}}$ for excitation, and $h_{INH}(t) = INH \cdot t \cdot e^{\frac{-t}{\tau_i}}$ for inhibition, leading to a set of 6 differential equations forming the full NMM:

$$\dot{y}_{0}(t) = y_{3}(t)
\dot{y}_{1}(t) = y_{4}(t)
\dot{y}_{2}(t) = y_{5}(t)
\dot{y}_{3}(t) = EXC \cdot aS_{1}(y_{1}(t) - y_{2}(t)) - 2ay_{3}(t) - a^{2}y_{0}(t)
\dot{y}_{4}(t) = EXC \cdot a[p(t) + S_{2}(y_{0}(t))] - 2ay_{4}(t) - a^{2}y_{1}(t)
\dot{y}_{5}(t) = INH \cdot bS_{3}(y_{0}(t)) - 2by_{5}(t) - b^{2}y_{2}(t)$$
(1)

Table 1 lists the studies that were used to identify sigmoid function parameters corresponding to pyramidal cells and interneurons. Since, in these studies, the pulse-to-wave relation was studied with respect to current, the first step was to convert the data with respect to voltage. To do so, a current-to-voltage conversion was used according to Destexhe and Paré's study (Destexhe & Paré 1999), which used Ohm's law with the membrane resistance. Then, a non-linear least square (Levenberg-Marquardt algorithm)

Parameter	Interpretation	Value
EXC	Average excitatory synaptic gain	60
INH	Average inhibitory synaptic gain	15
$\tau_e = 1/a$	Dendritic average time constant	1/100 s
	in the feedback excitatory loop	
$\tau_i = 1/b$	Dendritic average time constant	1/35 s
	in the feedback inhibitory loop	
V_{0e}	Parameters of the sigmoid	$V_{0e} = 6 \text{ mV}$
e_{0e}	function for pyramidal cells	$e_{0e} = 45.4 \text{ Hz}$
$ m r_e$		r_e =0.519 mV ⁻¹
V_{0i}	Parameters of the sigmoid	$V_{0i} = 12.9 \text{ mV}$
e_{0i}	function for interneurons	$e_{0i} = 143 \text{ Hz}$
r_i		$r_i = 0.262 \ \mathrm{mV^{\text{-}1}}$
p(t)	Excitatory input noise	m = 90 Hz
	(Gaussian white noise)	s = 30 Hz

Table 2. Model parameters.

was used to fit data from the literature, and an average was made between the two available sets to obtain sigmoid function parameters used in our model. The three sigmoid functions that were included in the NMM are:

$$S_{1} = S_{2} = \frac{e_{0_{e}}}{1 + e^{r_{e}(v_{0_{e}} - v)}}, with \begin{cases} e_{0_{e}} = 45.4 \\ r_{e} = 0.519 \\ v_{0_{e}} = 6 \end{cases}$$

$$S_{3} = \frac{e_{0_{i}}}{1 + e^{r_{i}(v_{0_{i}} - v)}}, with \begin{cases} e_{0_{i}} = 143 \\ r_{i} = 0.262 \\ v_{0_{i}} = 12.9 \end{cases}$$

$$(2)$$

2.3. Estimation of excitatory and inhibitory post-synaptic currents

We verified that this novel NMM could reproduce the four possible types of activity of the original model, namely: background activity, sporadic epileptic interictal discharges, rhythmic discharges and narrow band activity at theta and alpha frequencies (see Figure_S1_ActivityMap in Supplementary materials). The reverse modeling approach uses LFPs (simulated or recorded experimentally) that are injected in the NMM. The originality in our approach is to force the NMM state at the node of the associated block diagram (see figure 1(c)), corresponding to where LFPs are generated ("LFP input"), forcing other model components to use the LFP that has been injected as an input. By comparing the block diagrams in figure 1(b) and 1(c), the LFP input of the reverse modeling approach (figure 1(c)) corresponds to the LFP output of the NMM (figure 1(b)).

The injected LFP was first normalized to ensure that its magnitude was similar to simulated LFPs. The physical unit of LFPs, as well as ePSP and iPSP, is in terms of Volts, and the normalization process is used to avoid considering potential differences in the amplification gain of the recording hardware. Furthermore, the normalization was done only once for the entire time course of the analyzed signal, even if the LFP was split into windows afterwards as explained below. The reverse modeling approach involved removing the noise input (s=0) to avoid adding a stochastic component to the process, and also to correctly adapt EXC and INH parameters to fit optimally the input LFP with the estimated LFP. Since the model is then in an "open loop" form, the noise variance would just add noise onto ePSP. The method only allows the EXC and INH parameters to vary, since all other parameters relate to intrinsic tissue properties. Therefore, the dendritic average time constants (1/a and 1/b) and the sigmoid function parameters (V_{0e}, e_{0e}, r_e, V_{0i}, e_{0i}, and r_i) were fixed based on the literature. In order to ensure that NMM dynamics corresponds to the injected LFP, a key step consists in comparing the NMM output LFP (estimated LFP computed in the reverse modeling approach) with the injected LFP, and to identify NMM gain parameters maximizing the match between LFP and LFP. The identified gain parameters are EXC for the excitatory loop and INH for the slow inhibitory loop. This identification was performed using a gradient descent, where the cost function was chosen as the root mean square (RMS) error between LFP and LFP in magnitude (M_{RMS}) and derivative (d_{RMS}) :

$$M_{RMS} = \sqrt{\frac{\sum_{n=1}^{N} \left(LFP_n - \widehat{LFP}_n \right)^2}{N}},$$

$$d_{RMS} = \sqrt{\frac{\sum_{n=1}^{N} \left(\frac{dLFP_n}{dt} - \frac{d\widehat{LFP}_n}{dt} \right)^2}{N}},$$
(3)

$$d_{RMS} = \sqrt{\frac{\sum_{n=1}^{N} \left(\frac{dLFP_n}{dt} - \frac{d\widehat{LFP}_n}{dt}\right)^2}{N}},$$
(4)

$$RMSE = M_{RMS} + d_{RMS}, (5)$$

where N is the number of samples in the LFP.

Using both the difference in magnitude and derivative between LFP and LFP in the cost function enables evaluating the similarity in magnitude, but also in tendency. In the gradient descent method, both EXC and INH parameters were set to initial values (in a 2D parameter space) and the estimated LFP was computed for different set of parameters around the initial ones (on an ellipse where both radii were defined on the 2D parameter space d_1 and d_2). The gradient descent algorithm ended when no error lower than the current parameter set was found, and current EXC and INH parameters were then identified as the optimal parameters. It should be noted that this method does not identify all the basins of attraction in a 2D parameter space. As an attempt to verify that the set of identified parameters was unique, we also checked if there was, for a wide range of parameter values for EXC and INH, a unique solution under the form of a basin of attraction within the parameter space. This is a critical requirement for our method, since it would be problematic to have multiple potential

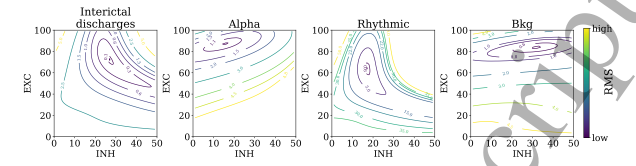


Figure 2. Illustration of the uniqueness of basins of attraction for parameter reconstruction using different LFP modes. RMSE computed within a 2D grid parameter space (EXC and INH). Note that only one basin of attraction is found for each type of activity. Interictal discharges are epileptiform events not associated with seizure symptoms, alpha corresponds to an alpha rhythm with frequency between 8 and 12 Hz, rhythmic corresponds to periodic ictal periods, and bkg corresponds to background activity.

parameter values for a single LFP to analyze. Parameters were set to form a 2D grid where EXC values varied from 0 to 100, and INH values varied from 0 to 50. The step for EXC was 0.25 and the step for INH was 0.125, leading to a 400x400 grid. For each parameter set, the RMS value between LFP and LFP was computed. We present in Figure 2 the corresponding result for qualitatively different LFPs (interictal discharges, alpha, rhythmic and background activity), which highlight the uniqueness of the parameters set towards which the method converges. Note that this does not prove that it is impossible to have several basins of attraction, however we only identified one basin of attraction in all of our tests. The four types of LFP tested were: Interictal discharges (epileptiform events not associated with seizure symptoms), alpha rhythm (frequency content between 8 and 12 Hz), rhythmic discharges (periodic ictal periods), and background activity.

Custom software was designed in Python, integrating a Qt interface (See Video_S1_Software_demo in Supplementary materials), dedicated to fitting at best NMM parameters on an injected LFP along short sliding windows on the time axis (i.e., NMM gain parameters are identified once per sliding window). Excitatory (EXC) and inhibitory (NH) gains could therefore be tracked during the LFP time course, along with the excitation to inhibition ratio (EIR, computed as the EXC/INH ratio). Obviously, the NMM provides full access at any time to the intermediate states for each sub-population, such as the average firing rate of action potentials and average post-synaptic potentials at the level of pyramidal cells and interneurons. Such signals are challenging to extract from LFPs with standard signal processing methods, especially since LFPs originate from the summation of PSPs onto pyramidal cells, and are not directly related with APs at the soma level. Using our approach, FRs can be directly deduced from a LFP with the reverse modeling approach, since FRs are intermediate states in the NMM.

2.4. Goodness-of-fit

The method used to compute the goodness-of-fit between an LFP and a reconstructed LFP was the zero normalized cross correlation (ZNCC (Nakhmani & Tannenbaum 2013)). This method provides an indication of similarity between two signals, named γ , which is bound [-1;1]. This indicator is 1 when both signals are identical, -1 when one signal is the opposite of the other, and is close to 0 when both signals are far from similar. This indicator neither depends on the length of the LFP nor the amplitude scales of different LFP measures (different amplifier gains for instance). Therefore, the ZNCC is especially suited to compare results for different LFP time/amplitude scales.

2.5. Simulated epileptic activity using another published model

In order to compare the performance of our model-based reconstruction of excitatory and inhibitory post-synaptic potentials, we used an intrinsically different computational model of LFP, known as the " Z^6 model" (Kalitzin et al. 2010). The Z^6 model is a mathematical phenomenological model, that has been previously used in the context of epileptic signals interpretation (Koppert et al. 2016). Although it is more abstract than the NMM, it includes one parameter ('c') that allows for controlling the excitability of the system, which provides us a ground truth for assessing the excitation/inhibition ratio (EIR) estimated with the proposed method. When $-1 \le c < 0$, this model generates steady-state behavior for values close to -1 (background activity), or a limit cycle for values close to 0 (ictal activity). All other parameters were fixed: a = -2; b = 2; $\omega_{mean} = 1$; $\omega_{sd} = 0.2$; $\eta_{mean} = 0$; $\eta_{sd} = 0.2$.

2.6. In vivo recordings

Electrophysiological recordings from N=7 mice were performed in accordance with the European Community Council Directive of November 24th 1986 (86/609/EEC), and were approved by the local ethics committee from the University of Rennes (agreement No 7872-2017031711448150). In all cases, bipolar electrodes (tips 400 μ m apart) were implanted bilaterally in the hippocampus (-2 mm anteroposterior, -1.5 mm mesio-lateral, -2 mm dorso-ventral from the Bregma) of C57B6j/Rj mice (80 days old). A reference electrode (monopolar) was placed at the cerebellum level. Mice were made epileptic following an intra-hippocampal injection of kainic acid that triggered an approx. 4-week epileptogenesis phase preceding the chronic epileptic phase. LFPs were sampled at 2048 Hz and hardware highpass filtered (0.16 Hz cutoff frequency). Recordings were made after the 4-week epileptogenesis period (chronic epilepsy stage). After protocol completion, mice were euthanized using the CO2 gradient method.

Electrophysiological recordings in rats (N=1) were obtained from the company Biotrial (http://www.biotrial.com). Ethics approval was obtained for these recordings, which involved injection of convulsive PTZ doses. A single bipolar electrode was implanted in the rat cortex, while a reference electrode was placed at the cerebellum

level. A surgically implanted wireless transmitter enabled recording throughout the entire experiment without any physical manipulation of the rat, avoids movement artifacts. PTZ injection was performed through perfusion at the dose of 75 mg/kg.

2.7. In clinico recordings

In the context of pre-surgical planning, epileptic patients candidate to surgery undergo stereoencephalography (sEEG), involving the implantation of multiple intracranial electrodes aiming at improving the identification of epileptogenic zones. sEEG recordings from N=5 patients in the context of clinical evaluation were obtained from the Epileptology Unit of Rennes University Hospital (CHU Pontchaillou, Rennes, France). sEEG recordings were sampled at 2048 Hz with a Deltamed EEG acquisition system. Seizure epochs, involving several minutes pre- and post-seizures, were extracted for each patient to enable LFP reconstruction pre-, per- and post-seizure.

All the data are available on the following public link https://zenodo.org/deposit/1285630 and with the DOI: 10.5281/zenodo.1285630

3. Results

3.1. In silico validation

The reverse modeling method was first tested on simulated data. In this first validation step, the input LFP was simulated by the same NMM as the one used in the reverse modeling algorithm. The unique advantage is the ground-truth: estimated parameters (denoted by ePSP and iPSP) can be directly compared to actual components (ePSP and iPSP) of the simulated LFP used as an input signal.

Figure 3 presents the method performance on simulated data. Scatter plots in figure 3(a) display estimated parameter values ($\widehat{\text{EXC}}$, $\widehat{\text{INH}}$ and $\widehat{\text{EIR}}$: excitation to inhibition ratio) versus reference parameter values (denoted as EXC, INH and EIR) for a set of 35 simulated LFPs. The reverse modeling approach provides an excellent identification of parameters, since points in the scatter plot are close to the ideal result (represented by black lines when identified and reference parameters are equal). The method also provides an excellent $\widehat{\text{EIR}}$ estimation, represented by the $\widehat{\text{EXC}}/\widehat{\text{INH}}$ ratio. Panels in figure 3(b) to 3(g) present four qualitatively different types of LFPs generated from the model, and the corresponding reconstruction computed from the proposed approach.

Panels in figure 3(b) display LFPs, while panels in figure 3(c) display ePSPs, and panels in figure 3(d) display iPSPs. As depicted in figure 3(b), results revealed an excellent LFP reconstruction from estimated synaptic components, as shown by the goodness-of-fit indexes close to 1 (γ =0.998 for interictal discharges, γ =0.968 for alpha, γ =0.999 for rhythmic and γ =0.894 for background). For background activity, the goodness-of-fit index was lower (γ =0.894) due to the strong contribution of the stochastic noise input (p(t)) for this particular type of activity. Reconstructed LFPs

were obtained from estimated postsynaptic potentials, either excitatory (ePSP) or inhibitory (iPSP), respectively shown in panels in figure 3(c) and 3(d). As depicted, these estimated components perfectly fit the reference components (ePSPs and iPSPs) that were summed up to obtain the simulated LFP signal used as an input for the reverse modeling approach. As for average PSPs, average firing rates (FRs) of neuronal sub-populations are explicit variables of the NMM. Therefore, it is possible to access to these signals. As an example, FRs are presented in figure 3(e) for FR1, figure 3(f) for FR2 and figure 3(g) for FR3. As depicted, both reconstructed and reference FRs are quasi-identical.

In the previous example, the same model has been used both to generate LFPs and as the core for the reverse modeling approach. In order to perform a validation using another model based on a different formalism, we present below the results for LFPs generated with the Z^6 model ((Kalitzin et al. 2010), see Methods section). We generated 150 s long LFP signals, with different values of the parameter c (from -1 to -0.01). Our method was applied to 4 s windows with 1 s shift. Results are displayed in figure 4. The overall goodness-of-fit was γ =0.948, which shows an excellent agreement between the generated LFPs and reconstructed LFPs. Results point at a small decrease of EXC and a more pronounced decrease of INH for increasing c values, leading to an EIR increase. Note that the EIR increases when the c parameter decreases, which in both cases results in an increase of excitability. In addition, our method also captures the non-linearity of the Z^6 model, as shown in figure 4(c) where an exponential variation is seen. Since the c parameter controls the excitability of the system, we conclude that our method is able to capture the same excitability increase from LFP signals generated with another model of epileptiform activity. In other words, the performance of our method does not rely on the type of computational model used to generate epileptiform signals.

In the following section, we move from the validation on simulated data to an experimental validation of the proposed method in the specific context of epileptic activity. This choice was motivated by the fact that the major qualitative changes observed in LFPs (during the interictal to ictal transition for example) involve drastic changes in neuronal excitability, potentially quantified by the proposed approach. Typically, seizures are associated with a significant increase of the excitation to inhibition ratio (Fritschy 2008).

3.2. In vivo and in clinico evaluation of the proposed approach

3.2.1. PTZ-induced seizure (rat). In vivo results obtained in rats following PTZ (pentylenetetrazol) intra-peritoneal injection are reported in figure 5. Results indicated that the proposed method leads to a reconstructed $\widehat{\text{LFP}}$ signal (red line, figure 5(a)) that was in excellent agreement (normalized goodness-of-fit between 0.85 and 0.97) with the experimentally-recorded LFP (black line, figure 5(a)) for the entire recording duration (130 seconds), as depicted on the 4 selected epochs (figure 5(a)). This illustrates the

robustness of our estimation method, which provides accurate estimates for epochs featuring qualitatively different dynamics. In addition to close replication of LFP dynamics, the estimation of the excitation/inhibition ratio (EIR) presented in figure 5(b), was in agreement with the expected excitability changes during the seizure time course: the EIR transiently increased at the onset of the PTZ-induced seizure, and then dramatically increased again during the seizure. It gradually decreased back to baseline as the seizure terminated. Interestingly, a decrease in the excitation level is observed during the seizure (figure 5(b)), which is however more modest than the drastic drop in inhibition, therefore still resulting in an increased EIR. Furthermore, since PSPs are computed in the NMM, they can be estimated through the proposed method as presented in figure 5(c). The gradual decrease in inhibitory post-synaptic potentials associated with the seizure is clearly visible from epoch 1 to epoch 3, with a slight recovery in epoch 4 (figure 5(c)). Overall, there is satisfactory qualitative and quantitative agreement of LFP dynamics on the one hand, and of estimated EIR and PSPs on the other hand. This supports the validity of our model-based reconstruction of excitatory and inhibitory components in data obtained in vivo. In addition, our method points at a result that is non-intuitive, namely the slight excitation decrease at the seizure onset.

3.2.2. Interictal activity in the kainate model of epilepsy (mouse). The proposed method was also tested in epileptic mice (kainate model, see Materials and Methods section). Based on our method applied to 53 epochs of interictal events (hippocampal paroxysmal discharges, HPD) in N=7 mice, a reconstruction of excitatory and inhibitory components along with the EIR was performed. As depicted in figure 6(a), the modelbased reconstruction of the LFP (red line) was in excellent agreement (goodness-offit index γ between =0.89 and 0.94) with the experimental LFP (black line) for the four considered epochs (see numbered panels in figure 6(a)). The reconstructed LFP accurately matched the experimental LFP not only during background activity (panel 3 in figure 6(a)), but also during HPD (panel 4 in figure 6(a)). In addition, the time course of the estimated EIR (figure 6(b)) was consistent with the occurrence of HPD within the signal: the EIR remained low during background activity, transiently increases during HPD, before returning rapidly to background values (figure 6(b)). Interestingly, as can be seen in figure 6(b), there was both a reduction of EXC and INH components (and not a decrease of INH only as could be intuitively deducted in this case), which was however greater for INH, still resulting in a major EIR increase. This is similar to the results obtained in the case of the PTZ-induced seizure (see Section 3.2.1). In addition, figure 6(c) presents reconstructed post-synaptic potentials (ePSP and iPSP) corresponding to each of the four LFP epochs in figure 6(a). The reconstructed iPSP points at a drastic inhibition decrease during an interictal discharge (epoch 2 in figure 6(c)) or HPD (epochs 1 and 4 in figure 6(c)), as compared to background (epoch 3 in figure 6(c)). The slight decrease in the reconstructed excitatory PSPs during interictal events (interictal discharges or HPD) is also apparent (panels 1, 2 and 4 as compared

to panel 3 in figure 6(c)). Figure 6(d) presents results at the group level (N=53 epochs) regarding the excitation and inhibition levels during HPD as compared to background activity epochs. We observed significant decreases in excitation and inhibition (p < 0.001) during HPD as compared to background activity, with a lesser excitation decrease as compared to the inhibition decrease, which still results in a drastic \widehat{EIR} increase.

3.2.3. LFP recordings after injection of a sub-convulsive dose of PTZ (mouse). In the third in vivo experiment, a non-epileptic mouse was injected with a sub-convulsive dose of PTZ, aiming at validating our model-based method when LFP changes are subtler (i.e., where no seizures are present). The LFP recorded over the entire experiment (approx. 2200 seconds long) is presented in figure 7(a). Estimated EXC and INH components of the LFP are presented in figure 7(b) and 7(c), and illustrate a significant increase (p < 0.001) of the EIR (figure 7(d)) shortly after injection, as compared to pre-injection levels. Consistent with the PTZ-induced rat seizure, the EIR increased (figure 7(d)) due to a joint decrease of EXC and INH components, which is more pronounced for the INH component than EXC. Furthermore, the EIR followed an exponential decay past the first 500 seconds after injection (the 500 s immediately after injection are discarded to avoid contamination by movements of the mouse due to manipulation by the experimenter), which is consistent with a pharmacokinetic response. The decay curve of the EIR was fitted with an exponentially decaying curve, and the time constant optimizing the fit was estimated to be 370 seconds, in agreement with the PTZ pharmacokinetic time constant found in the literature (Wendling et al. 2016, Mandhane et al. 2007).

In clinico validation of the method in epileptic patients. Our method was validated using intracranial recordings from N=5 epileptic patients undergoing stereoencephalography (sEEG) in the context of pre-surgical evaluation. Among the available recordings, we extracted a total of 47 spontaneous seizures, which featured artifact-free epochs pre- and post-seizures, so that the EIR evolution could be tracked over the entire seizure time course. Figure 8 presents an example of human seizure and associated LFP reconstruction (figure 8(a)) along with EIR estimation (figure 8(b)), and reconstruction of PSPs (figure 8(c)). LFP reconstruction is illustrated in figure 8(a), with the four zoomed panels emphasizing qualitatively different portions of the signal, illustrating the excellent agreement between the recorded intracranial signal (black line) and the reconstructed LFP (red line), as quantified by the goodness-of-fit index ranging between 0.92 and 0.98. In the seizure presented as an example in figure 8(a), the estimated EIR over the seizure time course (see figure 8(b)) was physiologically plausible, with a drastic EIR increase during the seizure itself, with a return to pre-ictal values after seizure termination. The reconstructed post-synaptic components (figure 8(c)) for each of the four panels presented in figure 8(a) also pointed at a gradual decrease in the inhibition level as the seizure progresses, consistently with the in vivo PTZ-induced seizure studied (figure 5). Figure 8(d) presents the group results (N=5) and compares

reconstructed excitation and inhibition levels during interictal and ictal epochs. The estimated $\widehat{\text{EIR}}$ increase during seizure was driven by a significant $\widehat{\text{EXC}}$ increase (p < 0.001), combined with a significant $\widehat{\text{INH}}$ decrease (p < 0.001). Reconstructed $\widehat{\text{LFPs}}$ for this database matched closely actual sEEG signals (analogous to LFPs), with an average goodness-of-fit γ =0.893 ±0.56.

4. Discussion

We developed a novel model-based method enabling the plausible reconstruction of excitatory and inhibitory post-synaptic potentials involved in LFP generation as measured experimentally by extracellular electrodes. This method can be applied in a wide range of applications (in in vivo/in vitro studies conducted in animals or in patients with implanted electrodes), typically when local neuronal network dynamics and excitability need to be assessed from recorded electrophysiological signals. To our knowledge, this is the first study solving the ill-posed inverse problem of LFP to estimate its main synaptic components based on a neural mass model. In this regard, it differs from two previously-published studies (Einevoll et al. 2007, Grativ et al. 2011), where the authors introduce a method called laminar population analysis (LPA) to estimate firing rates and evoked synaptic activity from multi-electrode laminar recordings. First, the LPA is based on a detailed description of the laminar structure of the cortex. Second, it assumes that the observed LFP can be decomposed as a linear sum over LFP population templates and that synaptic connection patterns can be extracted from the fitted weights of the linear sum. Here, the neural mass nonlinear input-output functions are kept in the proposed reverse modeling method. Third, LPA was applied to stimulus-averaged data that consist in evoked responses which are relatively stereotyped and which differ, in this regard, from the spontaneous patterns analyzed in this study which show higher diversity.

The method is based on a neural mass model, since this modeling approach captures crucial physiological components such as the dendritic average time constants, or wave-to-pulse sigmoid non-linear functions. While obviously such models do not capture all biophysical aspects (e.g., neuron orientation with respect to the recording electrode as mentioned in (Einevoll et al. 2013) or cell morphology (Gratiy et al. 2011)), it stills features the two main excitatory (Glutamate) and inhibitory (GABA) post-synaptic currents, which are generic and widespread among cortical structures. Furthermore, this model is able to generate LFP signals with only a limited number of parameters. The proposed method was tested at several levels: from in silico (using the method on LFPs generated using a NMM, providing a ground truth, as well as LFPs generated with another computational model, the Z^6 model), to in vivo (rat and mouse LFPs), and finally to in clinico (sEEG recordings in epileptic patients). In both in silico tests, the method was able to track excitability changes controlled by parameters EXC and INH for NMM-based LFPs, and by parameter c for Z^6 model. Regarding experimental data, we found plausible and relevant results emphasizing an EIR increase during the generation

of interictal events and during seizures. Furthermore, the reconstructed LFP, using our model-based decomposition, matched accurately experimentally recorded LFPs, suggesting that the assumption according to which excitatory and inhibitory postsynaptic potentials constitute the major contribution to LFP is reasonable. In addition, using a subconvulsive dose of PTZ in mice, we were able to provide an estimate of the PTZ time constant consistent with the literature (Wendling et al. 2016, Mandhane et al. 2007). Therefore, a major advantage of our technique is that it enables the detection of subtle variations in neuronal excitability; such as those induce by subconvulsive doses of PTZ. The increase of EIR is mainly explained by the major drop of GABAergic inhibition, already observed in computational modeling study (Wendling et al. 2002), animal model of epilepsy (Trevelyan et al. 2007, Wenzel et al. 2017) and in human epilepsy (Schevon et al. 2012). In addition, it is worth noting that the proposed method consistently pointed at a counter-intuitive decrease of excitation (minor as compared to the decrease of inhibition) in all in vivo recordings (PTZ-induced seizure, epileptiform events in the kainate model). At least the PTZ-model directly affects the function of type A GABAergic receptors and is likely to interfere with the inhibitory synaptic barrage. The slight decrease of excitation, predicted by the model, was not found during human seizures from our available database. A possible explanation is that the model focused only on post-synaptic potentials but not the overall excitation (e.g extrasynaptic ligand-receptors activation, potassium induced direct depolarization, paroxysmal depolarization shift, and other mechanisms). Moreover, slightly different pathophysiological mechanisms might be involved in the generation of epileptiform activity in the PTZ/Kainate in vivo models as compared to human seizures.

The possibility to reconstruct, from an experimentally recorded LFP, the time course of excitatory (glutamatergic) and inhibitory (slow GABAergic) post-synaptic contributions, is especially appealing. Our method provides full access to fundamental variables such as ePSPs and iPSPs for each subpopulation. These signals are challenging to extract from LFPs, since the problem is underdetermined. The identification of NMM parameters to optimize the fit of simulated LFPs with experimental LFPs has been addressed in several studies, using methods based on genetic algorithms (Wendling et al. 2005), Kalman filter (Bellanger et al. 2005), synaptic conductance estimation (Rudolph et al. 2004), or Hodgkin-Huxley parameters estimation (Zheng et al. 2012). However, none of these methods provides access to PSP signals from experimentally recorded LFPs. Our method should have applications in numerous areas of neuroscience, since LFP is a very common electrophysiological recording modality, from in vivo to in clinico, and ranging from neurophysiology to diagnostic applications. In addition, the method is sensitive enough to capture subtle changes in excitability, as shown in the case of the sub-convulsive PTZ dose experiment, and was able to estimate successfully PTZ pharmacokinetics.

One limitation is that our method currently features only slow, dendrite-targeting GABAergic synapses, and neglects fast soma- or axon-targeting GABAergic synapses; which limits the possible frequency range of LFPs that can be reconstructed. This

simplification was done for three reasons. First, we aimed at providing a proof-ofconcept that a method based on a physiologically-based NMM can provide meaningful excitatory and inhibitory components from an experimentally recorded LFP. Second, due to its very generic structure, the NMM used should be general enough to analyze LFPs from any brain region. Third, if a sub-population of fast GABAergic interneurons was added in the model, the proposed reverse modeling approach would require to identify three parameters respectively related to ePSPs, slow iPSPs and fast iPSPs by optimization in a 3D parameter space. In this case, the computational cost associated with the search of the global minimum would dramatically increase and an optimization algorithm different from the gradient descent should be used. This absence of fast GABAergic activity explains why the epochs where higher frequencies (beta band -13 to 30 Hz- and above) were present in the LFP were more challenging to reconstruct using our approach. This specific point explains why the goodness-of-fit is better in the case of HPD in mice (figure 6) as compared to epochs of relatively fast activity during the rat PTZ-induced seizure (figure 5): HPD are mainly characterized by slow activity, as compared to the relatively fast activity occurring at the onset of seizures. Therefore, agreement between reconstructed and experimentally measured LFPs is lower when higher frequencies (beta range and above, i.e. > 13 Hz) are present in the signal. Finally, it should also be mentioned that in the case where extracellular micro-electrodes (like microwires or micro-array electrodes) are being used to record multi-unit activity, the proposed method can only be applied to the LFP component of recorded signals (that can be obtained from low-pass filtering of recorded signals), regardless of the electrode geometry. This limits its range of application if the goal is to analyze action potentials in addition to recovering the post-synaptic components of the LFP.

Using our model-guided approach to unveil the time course of excitatory and inhibitory post-synaptic currents is promising, especially since the algorithm provides excitability tracking with a performance in real-time (Video_S1_Software_demo in Supplementary materials): on average, 2 s of LFP sampled at 1024 Hz requires 0.7 s using a single core on a PC equipped with an Intel Xeon E5-2637 3.5 GHz CPU and 64 GB of 1866 MHz RAM. Furthermore, there is still ample room for computation time optimization, since the algorithm is currently programmed in Python, which is known for not being optimal in terms of computation time. The possibility to automatically track neuronal excitability in real-time could be implemented on-chip, for example, in a neuromodulation device, to trigger stimulation when excitability exceeds a certain threshold. Another immediate application would be the monitoring of epileptic patients undergoing sEEG for pre-surgical evaluation.

Author Contributions

Conceived and designed the experiments: P.B., DJ.M., F.W. Computational model: M.Y., J.M., F.W. Performed the experimental study: P.B., J.M. Manuscript preparation: M.Y., P.B., J.M., DJ.M., F.W.

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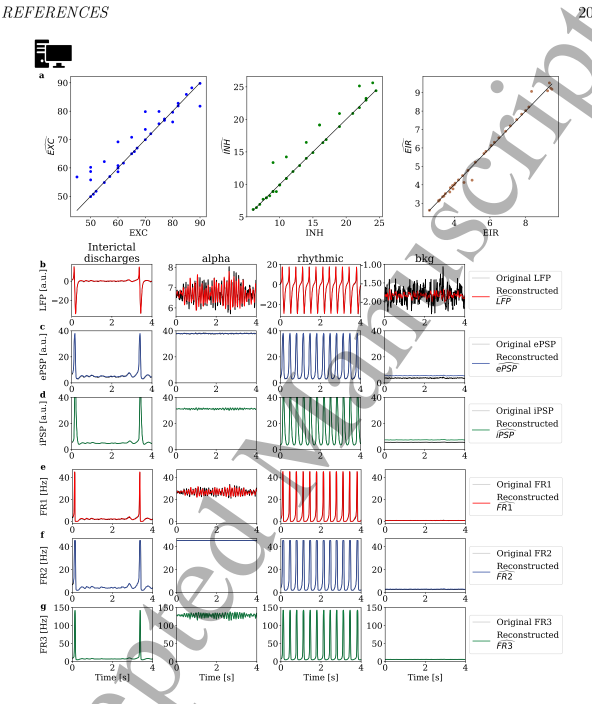


Figure 3. Results for simulated LFPs obtained using the same model as the reverse modeling approach. (a) Scatter plots for 35 simulated LFPs where ideal parameters vs identified parameters are displayed. Columns of combined panels b to e represent four types of activities. (b) LFP signals generated with the model (black) and LFP signals reconstructed with the reverse modeling approach (red). Goodness-of-fit indexes are: γ =0.998 for interictal discharges, γ =0.968 for alpha, γ =0.999 for rhythmic and γ =0.894 for background (bkg), (c) ePSP signals generated with the model (black) and ePSP signals extracted from the reverse modeling approach (blue). (d) iPSP signals generated with the model (black) and iPSP signals extracted from the reverse modeling approach (green). (e-g) show respectively FR1, FR2 and FR3 generated with the model (black) and FR1, FR2 and FR3 signals extracted from the reverse modeling approach (in color).

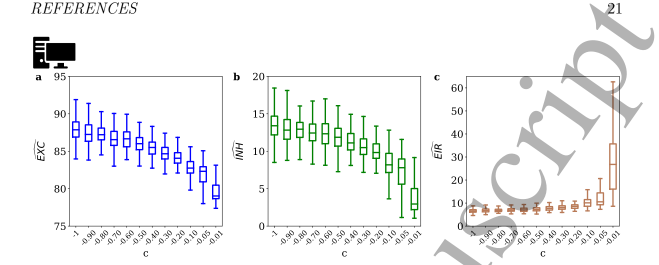


Figure 4. Results obtained with the Z^6 model with different c values (duration: 150 s per LFP signals for each value of c). $\widehat{\text{EXC}}$ and $\widehat{\text{INH}}$ were identified using our reverse modeling approach. (a) Boxplot of $\widehat{\text{EXC}}$, (b) $\widehat{\text{INH}}$, and (c) $\widehat{\text{EIR}}$.



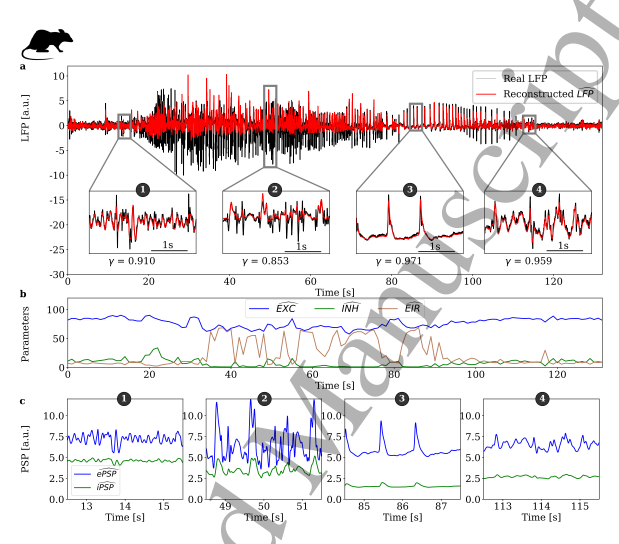


Figure 5. In vivo validation of the model-based LFP reconstruction in a PTZ-induced seizure (rat, N=1). (a) LFP signals recorded during a PTZ-induced seizure in rat (black line) and LFP signals reconstructed using the reverse modeling approach (red line) for different types of activity (panels numbered from 1 to 4). The normalized goodness-of-fit index γ is presented for each activity type. (b) Identified model parameters: excitatory EXC (blue line), inhibitory $\widehat{\text{INH}}$ (green line), and the excitability ratio $\widehat{\text{EIR}}$ (brown line). (c) Reconstructed $\widehat{\text{ePSP}}$ and $\widehat{\text{IPSP}}$ extracted from the NMM for each numbered panel presented in (a).

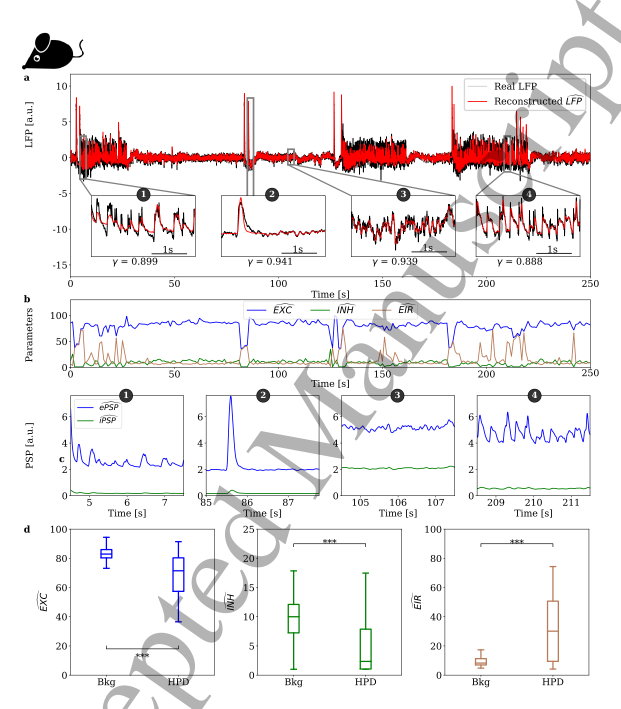


Figure 6. In vivo validation of the model-based LFP reconstruction in kainate mice (N=7). (a) LFP signals recorded in mouse where HPD are present (black line) and LFP signals reconstructed with the reverse modeling approach (red line) for different types of activity (panels numbered from 1 to 4). The normalized goodness-of-fit index γ is presented for each activity type. (b) Identified model parameters: excitatory EXC (blue line), inhibitory $\widehat{\text{INH}}$ (green line), and the excitability ratio $\widehat{\text{EIR}}$ (brown line). (c) Reconstructed ePSP and iPSP extracted from the NMM for each numbered panel presented in (a). (d) Boxplot of $\widehat{\text{EXC}}$, $\widehat{\text{INH}}$, and $\widehat{\text{EIR}}$ over background activity vs. HPD segments of 53 LFPs (\approx 500s long each) in seven mice. *** stands for p<0.001. Statistical significance was assessed using a Welch's t-test (sample size: Bkg=1856 and HPD=2255 values.

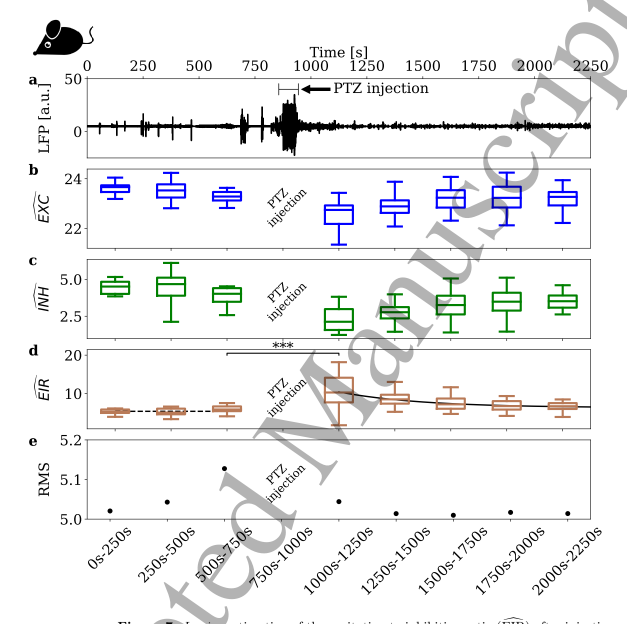


Figure 7. In vivo estimation of the excitation to inhibition ratio ($\stackrel{\frown}{\text{EIR}}$) after injection of a subconvulsive dose of PTZ (mouse, N=1). (a) LFP recording. (b) Estimation of the $\stackrel{\frown}{\text{EXC}}$ parameter. (c) Estimation of the $\stackrel{\frown}{\text{INH}}$ parameter. (d) $\stackrel{\frown}{\text{EIR}}$. Dashed line corresponds to the average of three first boxplots. The continuous line corresponds to an exponential decay model ($ae^{\frac{t-1000}{b}}+c$, with a=4.1, b=-370 and c=6.3) on the five last boxplots. *** stands for p<0.001. Statistical significance was assessed using a Welch's t-test (sample size: 250 values per window). (e) RMS values computed for each time window.

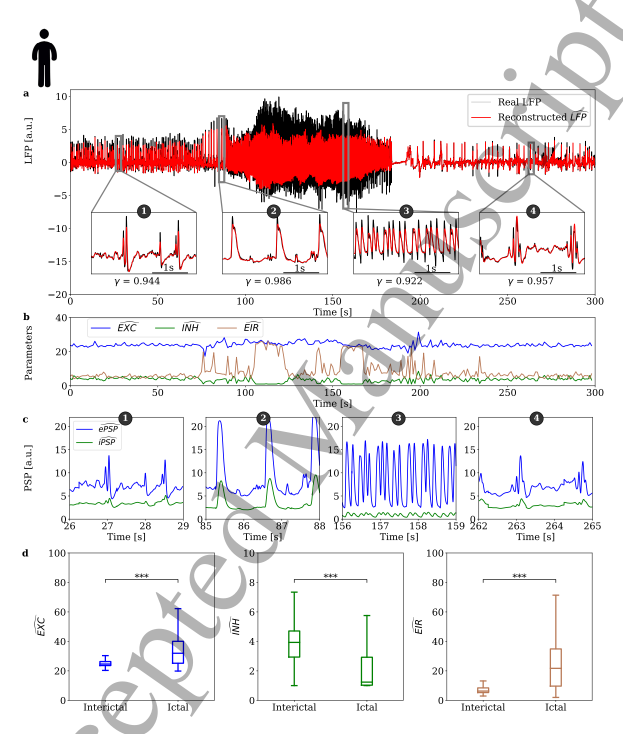


Figure 8. Validation on human seizure data (epileptic patients, N=5) obtained through intracranial recordings. (a) LFP signals recorded in human where a seizure is present (black line) and LFP signals reconstructed with the reverse modeling approach (red line) for different types of activity (panels numbered from 1 to 4). The normalized goodness-of-fit index γ is presented for each activity type. (b) Identified model parameters: excitatory EXC (blue line), inhibitory INH (green line), and the excitability ratio EIR (brown line). (c) Reconstructed ePSP and iPSP extracted from the NMM for each numbered panel presented in (a). (d) Boxplot of EXC, INH, and EIR ratio over interictal vs ictal segments of 47 LFPs in five patients. *** stands for p<0.001. Statistical significance was assessed using a Welch's t-test (sample size: Interictal=2283 values and Ictal=2255.