N-Methyl D-Aspartate Receptor Antagonists Amplify Network Baseline Gamma Frequency (30–80 Hz) Oscillations: Noise and Signal
Didier Pinault

To cite this version:
Didier Pinault. N-Methyl D-Aspartate Receptor Antagonists Amplify Network Baseline Gamma Frequency (30–80 Hz) Oscillations: Noise and Signal. AIMS Neuroscience, AIMS Press, 2014, 1 (2), pp.169 - 182. 10.3934/Neuroscience.2014.2.169. inserm-01441461

HAL Id: inserm-01441461
https://www.hal.inserm.fr/inserm-01441461
Submitted on 19 Jan 2017

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Commentary

N-Methyl D-Aspartate Receptor Antagonists Amplify Network Baseline Gamma Frequency (30–80 Hz) Oscillations: Noise and Signal

Didier Pinault 1,2,3,*

1 INSERM U1114, Neuropsychopathologie cognitive et physiopathologie de la schizophrénie, Strasbourg, France.
2 Université de Strasbourg, Strasbourg, France.
3 FMTS, Fédération de Médecine Translationnelle de Strasbourg, faculté de médecine, Strasbourg, France.

*Correspondence: Didier Pinault, Email: pinault@unistra.fr; Tel: +33-36-885-3245.

Abbreviations: AMPAr, alpha-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor; DCM, Dynamical Causal Modeling; DSM, Dimensional Systems Model; EEG, electroencephalogram; GABA, gamma-aminobutyric acid; GFO, gamma frequency (30–80 Hz) oscillations; NMDAr, N-methyl D-aspartate type glutamate receptors

1. Introduction

In 1924, Hans Berger invented the cortical electroencephalogram (EEG). He discovered the alpha frequency (~10 Hz) rhythm, which is recorded particularly in the occipital cortex during the resting state, that is, during relaxed wakefulness and in the absence of sensory stimulation or conscious mental activity [1]. Once eyes are open, baseline alpha oscillations are reduced in the cerebral cortex. They are also reduced during drowsiness and sleep. The Berger’s waves would be generated by a thalamic pacemaker [2]. Berger was the first to suggest that brain rhythmic electric oscillations in the human EEG are associated with mental processes, including cognition, memory, arousal, and consciousness. Since then, a growing body of studies has been consolidating the notion that EEG oscillations, including gamma frequency (30–80 Hz) oscillations (GFO), are biomarkers of brain state and function. Brain field oscillations are versatile and directly linked to the structure of the neural networks and to the neurotransmitter systems.

When and how field or network GFO and N-methyl D-aspartate glutamate type receptors (NMDAr) contribute to normal and dysfunctional cognitive performances? This open question is
currently the object of intensive clinical, experimental and theoretical investigations and of passionate debates. In the previous issue, three reviews written by Moss and Moss [3], Cadonic and Albensi [4], and by Pinotsis and Friston [5] provide three appealing non-exclusive theoretical viewpoints. Moss and Moss [3] discuss the possible roles in health and disease of cortical columns through the notion of the “dimensional systems model”, paying attention to the generation of “signal and noise” in neural circuits. Cadonic and Albensi [4] introduced the basic physical model of “damped and forced harmonic oscillators”, which are under the constraints of inhibitory or driving “forces” that impede or amplify network oscillations. Pinotsis and Friston [5] show how GFO, neural field models and “Dynamical Causal Modeling” can be combined to understand the generation of relevant signal (visual perception) and noise in dynamic neural circuits and the connectivity between brain regions. This challenges lateral connections which, by generating a functional excitatory centre-inhibitory surround, play a crucial role in GFO-based information processing. Lateral inhibitions might be vulnerable during cognitive disorders. Interestingly, simulations of neural field models can yield predictions of recorded field GFO and on the anatomofunctional properties of the related cortical circuits.

Here I take this opportunity to discuss these different theoretical perspectives while integrating them in a basic-clinical translational framework in an attempt to understand neurophysiological and pathophysiological aspects regarding the relation between NMDAr-mediated activities and GFO in mental disorders and brain illnesses. I argue that spontaneously-occurring field GFO (or network gamma noise), which are usually mostly intracerebrally generated (e.g, from resident cognitive information), can—during neurological and neuropsychiatric diseases—increase in a manner such that they can become the source of abnormal activities (e.g, during hallucinations) and disturb function-related synchronized oscillations (or network gamma signal). The gamma signal-to-noise ratio is considered as a potential neurophysiological biomarker of the state and function of neural circuits.

2. Baseline and Function-Related Network Gamma Oscillations

Natural, spontaneously-occurring, synchronized and non-synchronized GFO are dominant in the desynchronized cortical EEG [6], a EEG state that can be recorded during conscious awareness in the awake state, executive functions, selective attention [7,8,9], Rapid Eye Movement sleep [10,11], hallucinations [12–16], in early psychosis [17,18], and in the process of meditation [19]. At rest, in the visual cortex, differences in peaks of GFO variations are associated with a \( \gamma \)-aminobutyric acid (GABA)-related inhibitory drive [5].

Large-scale, ephemeral synchronized field GFO emerge during the performance of cognitive tasks, that is, during global brain operations like attention, perception and memory [20,21,22]. They also arise during pain perception [23]. They are thought to play a key role in the temporal interaction and coordination between multiple cortical and subcortical brain regions during information integration (binding-by-synchronization) [24–29], the focused arousal, the resting wake state [30] and synaptic plasticity [31]. Field GFO can be recorded as local extracellular field potentials associated with irregular firing of single nerve cells [32]. Network GFO are multiple and operate in combination with theta frequency and other (slower and faster) brain rhythms [32,33,34]. Field GFO principally result from subthreshold synaptic and intrinsic membrane potential oscillations triggering action potentials at a precise instant during the oscillatory period. Their functions and mechanisms are still matter of debate.
3. From Vertical to Horizontal Network Gamma Oscillations

Function-related synchronized field GFO are usually recorded principally in adult small- and large-scale cortico-cortical networks. These “horizontal” cortical network GFO correspond to binding-by-synchronization of multiple cortical areas, which are also connected to subcortical structures including the thalamus. In human, ongoing and function-related synchronized GFO emerge during early childhood, and their spatiotemporal properties continue to mature until early adulthood, suggesting they are associated with synaptic and network plasticity involving myelination processes and the development of GABAergic neurotransmission [35].

In the rodent, the somatosensory vibrissae-related cortico-thalamo-cortical system is composed of topographically organized and interacting anatomofunctional modules, the barreloid-barrel circuits, each of them being already active at birth [36,37,38]. Field GFO start to play functional and structural roles early during the development of the neocortex. Remarkably, in the rodent, early “vertical” thalamically-generated GFO start to emerge in response to the ongoing activity of sensory inputs during the neurodevelopment of thalamocortical circuits, especially during the critical period for activity-dependent plasticity in thalamocortical synapses and before the appearance of intracortical GABAergic-dependent inhibition [39,38]. These synchronized GFO are very likely driven by the GABAergic thalamic reticular nucleus, the pacemaker of thalamic GFO [40,41]. These sensory-evoked, thalamically-generated early GFO appear when thalamocortical connections present enhanced plasticity (long-lasting potentiation of thalamocortical excitatory postsynaptic potentials) Highly localized spontaneous and sensory-related thalamocortical and corticothalamic GFO can be recorded at birth in the newborn rat barrel cortex [38]. These rhythmic events spread to adjacent ontogenetic columns at the end of the first postnatal week. Early GFO trigger repetitive synchronization of thalamic and cortical neurons during the neurodevelopment and maturation of the topographical organization of cortico-thalamo-cortical connections. The development of the column 6-layers architecture is driven by spontaneous and sensory-related thalamocortical activity [38]. In short, vertical field GFO start to play an anatomofunctional role early during the development of the topographic maps in the somatosensory cortex, a natural neurodevelopmental process that requires precise temporal binding-by-synchronization in thalamocortical networks. Then, vertical and horizontal GFO work together during adult global brain operations.

4. The Dimensional Systems Model and Memory

In the previous issue, Moss and Moss [3] argue that a cortical column is a basic unit having all the ingredients to compute ongoing information during global brain operations or functional integration. In their theory, the anatomofunctional integrity of the cortical column relies on the emergence of network GFO. The notion that the neocortex operates on a basic principle based on modular elements, that is, the cortical columns, took its roots in the pioneering neurophysiological discoveries of Mountcastle [42], who later proposed every column is made of “minicolumns” [43]. Since then, the cortical column becomes the unit of computation and a focus of interest to investigate the anatomofunctional properties of neuronal circuits [44]. Whether or not the cortical column has a function remains an open debate and the object of intensive investigations [44,45].

Moss and Moss [3] consider their theoretical cortical column, with its hundreds of minicolumns [each containing ~100–200 neurons], as an elementary unit involved in cognitive processes. Their theory, the so-called “DSM or Dimensional Systems Model”, suggests functional overlapping between columns and minicolumns during cortical processing, leading to a dynamic column
formation based on synchronized GFO. In their model, there is room for series and parallel “light-buzzer” circuits, thereby providing multiple patterns of connections, like electrical devices (electrical circuits with power supplies and switches). The DSM takes into consideration 5 systems: sensory inputs, arousal system, attention-memory system, cortical system (information processing) and motor system (output). Thus, their theoretical model provides a system definition of the simplest to the more complex memory, including multisensory and association memories. It also allows the implication of subcortical structures, in particular the thalamus and hippocampus.

In their DSM, Moss and Moss [3] highlight the importance of AMPA (α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid), NMDA and GABA receptors in the formation and consolidation of memory. Initial AMPAr-based activity would be strengthened by NMDAr-mediated synaptic potentiation; then a horizontal spreading would allow the consolidation of connections between the first activated minicolumns and the follower ones. The spreading would involve the pyramidal cell-parvalbumine positive GABAergic interneuron gamma-based feed-forward inhibition. The authors also highlight the importance of diverse types of interneurons (expressing parvalbumine, somatostatin and vasoactive intestinal polypeptide), which play a crucial GFO-based role in cortical information processing.

In their theory, a set of cortical columns that is coherently consolidated during a memory process would form the relevant “signal” while the other, overlapping and adjacent, columns that display irrelevant and distractive “noise” would be inhibited. Such a concept is nothing other than the principle of lateral inhibition, which consists in sharpening the receptive field by generating consolidated patterns of center-on (signal) surround-off (noise) connections. Moreover, local and distant lateral neuronal interactions play important roles in facilitating contrast augmentation during information processing in sensory and other systems. Furthermore, Pinotsis and Friston [5] emphasize the notion of intimate relationship between stimulus contrast, GFO and lateral inhibition in the visual cortex (excitatory-inhibitory balance).

5. Ketamine Amplifies Baseline Network Gamma Oscillations

The glutamatergic systems mediate most of the excitatory neuronal transmissions through the activation of ionotropic and metabotropic receptors. The ionotropic NMDAr play a key role in the synaptic plasticity, memory processes and in the modulation of field oscillations (see Cadonic and Albensi, the previous issue [4]). Ketamine, a non-competitive NMDAr antagonist, can safely be administered in humans under clinical monitoring. It has dose-dependent multiple properties, including positive and negative effects. For instance, a single subanesthetic administration can disturb cognitive and sensory-perceptual processes and induce schizophreniform psychosis in healthy subjects [46–49]; puzzlingly but of importance, ketamine can generate a durable antidepressant effect in patients refractory to conventional antidepressant therapies [50,51,52].

More specifically, brain scans recently revealed that a single subanesthetic administration of ketamine in healthy subjects at rest produces in the prefrontal cortex a state of hyperconnectivity, which resembles that recorded in people in the early stages of schizophrenia but not in patients with chronic (since several years) schizophrenia [53]. Also, using fMRI in healthy human subjects, it was demonstrated NMDAr antagonist ketamine increases global brain functional connectivity and reduces negative symptoms [54]. The acute ketamine effects are quick, transient and reversible. These findings (hyperconnectivity and hyperactivity) are consistent with preclinical studies demonstrating that, in rodents, non-competitive NMDAr antagonists increase the amount of field GFO in cortical and subcortical regions (see below). In healthy subjects, ketamine increases the
power of GFO during auditory-evoked network oscillations [55].

In rodents a single subanesthetic administration of ketamine (or other NMDAr antagonists like dizocilpine [MK-801] and phencyclidine) quickly and transiently induces abnormal behavior (hyperlocomotion, ataxy), memory deficits and abnormally persistent and generalized hypersynchronized (200%–400% increased power) ongoing GFO [56–61] (Figure 1B, top panel). The gamma frequency at maximal power is significantly increased by approximately 10 Hz on average [56]. Interestingly, using conductance and convolution models, Pinotsis and Friston [5] suggest that such a gamma frequency shift reflects an increase in the strength of inhibition. The amount of ongoing higher-frequency (> 80 Hz) oscillations is also increased following a single subanesthetic administration of ketamine [60,62,63].

In the ketamine (or MK-801)-treated rodent, the persistent generalized and hypersynchronized GFO are not dependent on muscle activity, locomotion-related brain state or conscious sensorimotor processing. Moreover, they are also recorded in anesthetized and immobilized rodents in almost all cortical and subcortical structures implicated in sensory, motor, limbic and associative/cognitive systems [60]. The ketamine-induced persistent generalized and hypersynchronized GFO are thought to represent an aberrant diffuse network noise, a potential electrophysiological correlate of a psychotic(-like) state (see below).

In addition, NMDAr antagonists transiently disrupt the expression, not the induction, of long-term potentiation in the thalamocortical system (Figure 1B, bottom panel; [63]), disorganize action potential firing in rat prefrontal cortex [64], increase the firing in fast spiking neurons and decrease that in regular spiking neurons [65]. These results suggest that the amount of ongoing GFO is inversely related to synaptic potentiation (assessed from the amplitude of the sensory-evoked potential) at least in the thalamocortical system [63]. They also suggest that the ketamine-induced state results in part from dysfunction of cortical GABAergic interneurons that would lead to hyperexcitation of projection glutamatergic neurons [65].

It may be worth precisioning that the acute, single low-dose (< 10 mg/kg) ketamine rat model models more hyperfrontality, which can be observed in first-episode schizophrenia [53,54,66], than the hypofrontality of patients diagnosed with the chronic disease schizophrenia. Therefore, the acute ketamine model may be appropriate to model the pathogenesis of acute psychotic states, a model translatable in humans [47,48,53,54,67]. The advantages and weaknesses and possible mechanisms of the acute ketamine model are still a matter for discussion [68,69,70].

6. Damped and Forced Harmonic Oscillators

Cadonic and Albensi [4] introduced the basic physical model of “damped and forced harmonic motion” referring to mechanical vibrations in real-world systems. The motion of the oscillator is under the constraint of inhibitory or driving “forces” that impede or amplify the motion of the oscillator. Such a model can be applied to neural oscillations although, as stressed by the authors, the activity of individual nerve cells is not representative of the corresponding field activity, which is the integration of collective activities from local and sparse neuronal populations. So, mathematical models, which approximate electrical properties (capacitance, conductances, voltage and current sources) of nerve cells, are necessary to describe how, in neurons, firing patterns are generated. Network systems can be described for instance with the Wilson-Cowan model, which considers at least two types, excitatory and inhibitory, of interconnect neurons.

As above-mentioned, both the duration and the amplitude of spontaneously-occurring EEG bursts of GFO significantly increase in the rat frontoparietal cortex following the administration of
ketamine at a subanesthetic dose [56]. So, from the mathematical viewpoint presented by Cadonic and Albensi [4], it is tempting to propose that natural, physiological ongoing GFO operate like damped harmonic oscillators, which would leave room for synaptic potentiation, learning and memory, whereas ketamine-induced persistently amplified GFO run like forced harmonic oscillators, which would brake the expression of synaptic potentiation. From this perspective, one may wonder what are the so-called inhibitory or driving forces that are responsible for the acute persistent amplification of network ongoing GFO that appear following the systemic administration of the NMDAr antagonist ketamine.

It is well known that, in health and disease, GFO interact with other neural oscillations, in particular with theta oscillations [71,72]. Such interactions are termed cross-frequency-couplings [73], which are of several types (power-to-power, phase-to-phase, phase-to-frequency and phase-to-power). The functional role of cross-frequency-coupling is not yet understood [71,74]. The ketamine-induced increase in ongoing GFO might in part be the result of privileged interactions with theta oscillations, dual oscillations forming a spatiotemporal code that would be implicated in processes underlying learning and memory.

Further investigations are necessary to understand the contribution of the possible inhibitory or driving forces that work from one rhythm to the other and vice versa. Indeed, there is a growing body of evidence suggesting that the NMDAr antagonist ketamine modulates not only GFO and higher frequency oscillations, as above-mentioned, but also lower frequency oscillations, including alpha, theta and delta oscillations [55,61,75,76]. However, this broad-spectrum effect depends on the injected dose, the experimental and recording conditions and on the anatomofunctional properties of the structures under investigation. For instance, in in vivo conditions, a single low-dose (< 10 mg/kg) ketamine administration alters more specifically GFO and higher frequency oscillations [56,59,60,69] while higher doses in addition affect slower rhythms [61,62,75,77–80]. Therefore, we must be prudent when comparing results and inferring mechanisms from studies using different doses of NMDAr antagonists and various and diverse animal and network models. This is fundamental for basic-clinical translational understanding.

7. NMDAr-Related Network Dysfunction Modulates the Gamma Signal-To-Noise Ratio

Interestingly, in an attempt to understand the functional role of NMDAr and minicolumns, Moss and Moss [3] introduced the concept of “disrupted column formation” as a neuronal substrate of mental disorders and brain illnesses, like schizophrenia and Alzheimer’s disease.

The notion of disrupted column formation comforts the universal concept of “Disconnection Syndrome” or “Cerebral Dis/Dysconnections”, which attempts to explain disorders of sensory-perception, thought, cognition, emotion and of sensorimotor integration that are observed in many complex brain diseases, including Alzheimer’s disease, autism, dementia, schizophrenia, bipolar and attention deficit hyperactivity disorders [81–85]. Nowadays, it is clear from the literature that many of these mental disorders, each arising from more than one etiology, share common pathophysiological mechanisms, which include at least three essential facets: 1) brain abnormal rhythms, in particular in GFO [86–91], 2) dysfunction of cortical and subcortical networks, including cortico-thalamo-cortical circuits [92–97] and 3) NMDAr hypofunction [98,99].

Here, I would like to further argue on the notion of “signal-to-noise ratio” pointed out in the previous issue in their terms by Moss and Moss [3] and by Pinotsis and Friston [5]. In any neuronal system, baseline field oscillations [recorded with EEG and local field potential electrodes] represent a dynamic “network noise”. The oscillation properties (frequency, period, amplitude, power, etc.)
depend on the physiological or pathological brain state and on the recording conditions. Under a given pathological condition, such a noise (background activity) can increase in a manner such that it can mask or interfere with function-related synchronized oscillations, thereby affecting the ratio signal power to noise power. Here, the notion of signal—more precisely “network signal” - is a function-related response (e.g., sensory-evoked potential—with its related wave components - that is time-locked to the stimulus) of the system under investigation challenged by the activation of an afferent pathway (e.g, sensory stimulus). In short, in any system, both the amount of the ongoing (background or baseline) activity and the amplitude (or power) of its global response to the activation of its inputs are indicators of its state and functionality (Figure 2). The possible noise-signal interplay(s) might in part explain some disparities between findings (e.g., increases and decreases in GFO in patients with schizophrenia).

Figure 1. In the thalamocortical system, ketamine increases the power of baseline GFO and decreases both the power of sensory-evoked GFO and the synaptic plasticity. (A): Experimental design showing the simplified three-neuron circuit involving the neocortex (CT, corticothalamic [from layer VI]), GABAergic thalamic reticular nucleus (TRN) and thalamus with its principal neurons that project to the cerebral cortex (TC, thalamocortical). Natural-like mechanical stimulus (sensory stim) of the vibrissae is provided by a piezo bender actuator. Baseline cortical activity and sensory-evoked potentials (SEP) are recorded simultaneously with the surface electrocorticogram (ECoG) electrode and intracortical (layer IV) lfp (local field potential) micro-electrode. (B): Ketamine transiently disrupts the expression of the sensory-induced long-term potentiation. Top: changes in the baseline GFO power; bottom: changes in the SEP amplitude before and after ketamine (keta) injection. Each point is an average of 15 values x 4 rats (± SEM). The insets in gray show traces of ongoing GFO (top) and of averaged (n = 12) SEP under the two conditions (cont, keta). (C): Time-frequency graph of the ECoG for each condition (90 SEP trials, stimulus given at 0 ms). (D): Quantitative and statistical analysis (t-test, P < 0.0001) shows that ketamine (keta) administration increases the power of baseline GFO and decreases the power of sensory-evoked GFO. The averaged power of the baseline GFO is measured during the 100 ms epoch before the sensory stimulation (from at least 45 trials, three rats). The power of the sensory-evoked GFO is directly measured from the averaged SEP (12 values from the post-stimulus 100 ms epoch from three rats per condition). Adapted from Kulikova et al., 2012.
Figure 2. Ketamine decreases the signal-to-noise ratio in a network model composed of interconnected glutamatergic and GABAergic neurons. (A): The simplified cortical network shows local anatomofunctional interactions between GABAergic parvalbumine positive interneurons that are interconnected, electrically and synaptically, and that innervate (recurrent and lateral inhibitions) glutamatergic (GLU) pyramidal cells. Both GLU and GABA neurons have operational NMDA and AMPA receptors. Natural ongoing GFO are locally generated through interactions in the GLU-GABA network, thereby inhibiting pyramidal neurons that display irregular firing with action potentials (in blue) phase-locked with the positive wave of ongoing GFO recorded in the extracellular local field potential (in black). The ongoing intrinsically-generated network activity generates a certain amount of gamma noise (ongoing γ) and intrinsically-generated signals (noise + i-signal). This ongoing activity characterizes the “normal or natural” network state (mind). When challenging the system by the activation of a sensory afferent pathway (stimulus), it produces a measurable sensory-evoked signal (e-signal), here the averaged sensory-evoked potential (SEP) of a given amplitude, which reflects synaptic potentiation. The averaging procedure attenuates or eliminates the ongoing noise as it is not time-locked to the stimulus. (B): Following a single administration of ketamine at a subanesthetic dose, NMDAr are less operational than AMPAr. The GABAergic interneurons, assumed as being more sensitive to the NMDAr antagonist, emit less action potentials leading to reduced inhibition of pyramidal neurons. These latter disinhibited GLU neurons, which are more numerous than the GABAergic neurons (~85% vs. 15%), spontaneously generate (locally and distantly), through their numerous axon collaterals, massive synchronized rhythmic activity at the gamma frequency. This “generalized” disinhibited pyramidal rhythmic activity is recorded as abnormally high amplitude (high power) ongoing GFO, which corresponds to a huge gamma noise and intrinsically-generated signal (NOISE + i-signal). Under such a pathological condition, the averaged SEP is recorded with an amplitude lower than that of the SEP recorded under the normal (control) condition, revealing an apparent decrease in synaptic potentiation. In short, ketamine alters both the state and the function of the GLU-GABA network, thereby affecting the network noise and accompanying intrinsically generated and sensory-evoked signals (i-signal and e-signal, respectively).
More precisely, in the rat thalamocortical system, ketamine simultaneously increases the power of spontaneously-occurring GFO (signature of a change in the state of the system) and decreases sensory-evoked GFO (signature of a disturbance of the functionality of the system) [56,60,63] (Figure 1C,D). Assuming that sensory-evoked GFO include a “true” sensory-related component, the ketamine-induced gamma noise amplification decreases the ability of the thalamocortical system to discriminate the sensory-evoked gamma signal drowned in the noise. In other words, the NMDAr antagonist ketamine decreases the gamma signal-to-noise ratio during sensory information processing (Figure 2). Such a ratio is considered as a suitable neurophysiological marker of neural networks to evaluate their function and dysfunction [61,100,101,102].

This abnormally excessive ongoing gamma noise is thought to affect global brain state and operation and to contribute to psychosis. Moreover, continuous and stereotyped GFO might be responsible for clinical positive symptoms [103]. Furthermore, ongoing abnormally hypersynchronized GFO have been recorded in patients experiencing sensory hallucinations [12–16]. Hypersynchronized GFO in cortico-thalamo-cortical systems are thought to play a key role during the appearance of hallucinations [12,14], arisng the question as to whether persistent amplification of ongoing GFO somehow could generate aberrant signals and conceal function-related GFO in the corresponding brain networks.

8. Conclusion

Healthy neural networks have the ability to discriminate, from ongoing intracerebrally generated background activities—under or not the influence of external world’s stimuli,—the appropriate signal(s) at the right time during cognitive and sensorimotor processes. During information processing, neuronal interactions play important roles in facilitating, via lateral GABAr-mediated inhibitions, contrast augmentation. As against, many mental disorders and brain pathologies have, in spite of their respective etiology, common pathophysiological characteristics, in particular dysfunction of brain networks leading them to exhibit abnormal GFO. Abnormally hypersynchronized ongoing GFO might be the source of distorted thoughts and hallucinations [14]. In the cortico-thalamo-cortical system, NMDAr antagonism dramatically amplifies baseline network GFO, impedes synaptic plasticity and disturbs function-related GFO [63]. The mechanisms underlying network dysfunction might in part involve hypofunction of NMDAr on GABAergic interneurons, which would lead to a deficit in GABAr-mediated inhibitions, a subsequent hyper-excitation of the postsynaptic projection glutamatergic neurons [99,104,105], and disruption of lateral inhibitions [95]. Testing theoretical and pathophysiological hypotheses is an appealing and effective basic-clinical translational approach to understand how, in health and disease, our brain at work combines its various and miscellaneous molecular, synaptic, cellular and architectural complexities.

Acknowledgement

This work is supported by the French Institute of Health and Medical Research (INSERM, Institut National de la Santé et de la Recherche Médicale) and by the Université de Strasbourg.

Conflict of Interest

The author reports no conflict of interest associated with this article.
References

1. Ben-Simon E, Podlipsky I, Arieli A, et al. (2008) Never resting brain simultaneous representation of two alpha related processes in humans. *PLoS One* 3: e3984.
2. de Munck JC, Goncalves SI, Huijboom L, et al. (2007) The hemodynamic response of the alpha rhythm an EEG/fMRI study. *Neuroimage* 35: 1142-1151.
3. Moss RA, Moss J. (2014) The role of dynamic columns in explaining gamma-band synchronization and NMDA receptors in cognitive functions. *AIMS Neurosci* 1: 65-88.
4. Cadonic C, Alhensi BC. (2014) Oscillations and NMDA receptors their interplay create memories. *AIMS Neurosci* 1: 52-64.
5. Pinotsis D, Friston K. (2014) Gamma oscillations and neural field DCMs can reveal cortical excitability and microstructure. *AIMS Neurosci* 1: 18-38.
6. Jasper HH. (1936) Cortical excitatory state and variability in human brain rhythms. *Science* 83: 259-260.
7. Sheer DE. (1975) Behavior and brain electrical activity. New York and London: Plenum Press.
8. Sheer DE. (1989) Sensory and cognitive 40-Hz event-related potentials behavioral correlates, brain function and clinical application Brain Dynamics. Berlin: Springer, pp 339-374.
9. Kulli J, Koch C. (1991) Does anesthesia cause loss of consciousness? *Trends Neurosci* 14: 6-10.
10. Ferri R, Cosentino FI, Elia M, et al. (2001) Relationship between Delta, Sigma, Beta, and Gamma EEG bands at REM sleep onset and REM sleep end. *Clin Neurophysiol* 112: 2046-2052.
11. Cantero JL, Atienza M, Madsen JR, et al. (2004) Gamma EEG dynamics in neocortex and hippocampus during human wakefulness and sleep. *Neuroimage* 22: 1271-1280.
12. Baldeweg T, Spence S, Hirsch SR, et al. (1998) Gamma-band electroencephalographic oscillations in a patient with somatic hallucinations. *Lancet* 352: 620-621.
13. Becker C, Gramann K, Muller HJ, et al. (2009) Electrophysiological correlates of flicker-induced color hallucinations. *Conscious Cogn* 18: 266-276.
14. Behrendt RP. (2003) Hallucinations synchronisation of thalamocortical gamma oscillations underconstrained by sensory input. *Conscious Cogn* 12: 413-451.
15. Ffytche DH. (2008) The hodology of hallucinations. *Cortex* 44: 1067-1083.
16. Spencer KM, Nestor PG, Perlmutter R, et al. (2004) Neural synchrony indexes disordered perception and cognition in schizophrenia. *Proc Natl Acad Sci USA* 101: 17288-17293.
17. Bartha R, Williamson PC, Drost DJ, et al. (1997) Measurement of glutamate and glutamine in the medial prefrontal cortex of never-treated schizophrenic patients and healthy controls by proton magnetic resonance spectroscopy. *Arch Gen Psychiatr* 54: 959-965.
18. Theberge J, Bartha R, Drost DJ, et al. (2002) Glutamate and glutamine measured with 4.0 T proton MRS in never-treated patients with schizophrenia and healthy volunteers. *Am J Psychiatr* 159: 1944-1946.
19. Lutz A, Greischar LL, Rawlings NB, et al. (2004) Long-term meditators self-induce high-amplitude gamma synchrony during mental practice. *Proc Natl Acad Sci USA* 101: 16369-16373.
20. Joliot M, Ribary U, Llinas R. (1994) Human oscillatory brain activity near 40 Hz coexists with cognitive temporal binding. *Proc Natl Acad Sci USA* 91: 11748-11751.
21. Tallon-Baudry C, Bertrand O. (1999) Oscillatory gamma activity in humans and its role in object representation. *Trends Cogn Sci* 3: 151-162.
22. Varela F, Lachaux JP, Rodriguez E, et al. (2001) The brainweb phase synchronization and large-scale integration. *Nat Rev Neurosci* 2: 229-239.
23 Zhang ZG, Hu L, Hung YS, et al. (2012) Gamma-band oscillations in the primary somatosensory cortex, a direct and obligatory correlate of subjective pain intensity. J Neurosci 32: 7429-7438.

24 Buzsaki G, Chrobak JJ. (1995) Temporal structure in spatially organized neuronal ensembles a role for interneuronal networks. Curr Opin Neurobiol 5: 504-510.

25 Buzsaki G. (2006) Rhythms of the brain. Oxford University Press.

26 Engel AK, Roelfsema PR, Fries P, Brecht M, Singer W. (1997) Role of the temporal domain for response selection and perceptual binding. Cereb Cortex 7: 571-582.

27 Fries P. (2009) Neuronal gamma-band synchronization as a fundamental process in cortical computation. Annu Rev Neurosci 32: 209-224.

28 Gray CM, Konig P, Engel AK, et al. (1989) Oscillatory responses in cat visual cortex exhibit inter-columnar synchronization which reflects global stimulus properties. Nature 338: 334-337.

29 Singer W. (1999) Time as coding space? Curr Opin Neurobiol 9: 189-194.

30 Mantini D, Perrucci MG, Del GC, et al. (2007) Electrophysiological signatures of resting state networks in the human brain. Proc Natl Acad Sci U S A 104: 13170-13175.

31 Buzsaki G, Draguhn A. (2004) Neuronal oscillations in cortical networks. Science 304: 1926-1929.

32 Buzsaki G, Wang XJ. (2012) Mechanisms of gamma oscillations. Annu Rev Neurosci 35: 203-225.

33 Steriade M. (2006) Grouping of brain rhythms in corticothalamic systems. Neurosci 137: 1087-1106.

34 Roux F, Uhlhaas PJ. (2014) Working memory and neural oscillations alpha-gamma versus theta-gamma codes for distinct WM information? Trends Cogn Sci 18: 16-25.

35 Uhlhaas PJ, Roux F, Singer W, et al. (2009) The development of neural synchrony reflects late maturation and restructuring of functional networks in humans. Proc Natl Acad Sci U S A 106: 9866-9871.

36 Woolsey TA, Van Der Loos H. (1970) The structural organization of layer IV in the somatosensory region. (SI) of mouse cerebral cortex. The description of a cortical field composed of discrete cytoarchitectonic units. Brain Res 17: 205-242.

37 Van Der Loos H. (1976) Barreloids in mouse somatosensory thalamus. Neurosci Lett 2: 1-6.

38 Yang JW, An S, Sun JJ, et al. (2013) Thalamic network oscillations synchronize ontogenetic columns in the newborn rat barrel cortex. Cereb Cortex 23: 1299-1316.

39 Minlebaev M, Colonnese M, Tsintsadze T, et al. (2011) Early gamma oscillations synchronize developing thalamus and cortex. Science 334: 226-229.

40 Pinault D, Deschenes M. (1992) Voltage-dependent 40-Hz oscillations in rat reticular thalamic neurons in vivo. Neurosci 51: 245-258.

41 Pinault D. (2004) The thalamic reticular nucleus structure, function and concept. Brain Res Rev 46: 1-31.

42 Mountcastle VB. (1957) Modality and topographic properties of single neurons of cat's somatic sensory cortex. J Neurophysiol 20: 408-434.

43 Mountcastle VB. (1997) The columnar organization of the neocortex. Brain 120. ( Pt 4): 701-722.

44 Feldmeyer D, Brecht M, Helmchen F, et al. (2013) Barrel cortex function. Prog Neurobiol 103: 3-27.

45 Horton JC, Adams DL. (2005) The cortical column a structure without a function. Philos Trans R Soc Lond B Biol Sci 360: 837-862.

46 Adler CM, Goldberg TE, Malhotra AK, et al. (1998) Effects of Ketamine on Thought Disorder, Working Memory, and Semantic Memory in Healthy Volunteers. Biological Psychiatr 43: 811-816.

47 Hetem LA, Danion JM, Diemunsch P, et al. (2000) Effect of a subanesthetic dose of ketamine on memory and conscious awareness in healthy volunteers. Psychopharmacology (Berl) 152: 283-288.
48. Krystal JH, Karper LP, Seibyl JP, et al. (1994) Subanesthetic effects of the noncompetitive NMDA antagonist, ketamine, in humans. Psychotomimetic, perceptual, cognitive, and neuroendocrine responses. Arch Gen Psychiatr 51: 199-214.

49. Newcomer JW, Farber NB, Jevtovic-Todorovic V, et al. (1999) Ketamine-induced NMDA receptor hypofunction as a model of memory impairment and psychosis. Neuropsychopharmacology 20: 106-118.

50. Fond G, Loundou A, Rabu C, et al. (2014) Ketamine administration in depressive disorders a systematic review and meta-analysis. Psychopharmacology. (Berl). In press.

51. McGirr A, Berlim MT, Bond DJ, et al. (2014) A systematic review and meta-analysis of randomized, double-blind, placebo-controlled trials of ketamine in the rapid treatment of major depressive episodes. Psychol Med 1-12.

52. Zarate CA, Jr., Singh JB, Carlson PJ, et al. (2006) A randomized trial of an N-methyl-D-aspartate antagonist in treatment-resistant major depression. Arch Gen Psychiatr 63: 856-864.

53. Anticevic A, Corlett PR, Cole MW, et al. (2014) NMDA Receptor Antagonist Effects on Prefrontal Cortical Connectivity Better Model Early Than Chronic Schizophrenia. Biol Psychiatr [Epub ahead of print].

54. Driesen NR, McCarthy G, Bhagwagar Z, et al. (2013) Relationship of resting brain hyperconnectivity and schizophrenia-like symptoms produced by the NMDA receptor antagonist ketamine in humans. Mol Psychiatr 18: 1199-1204.

55. Hong LE, Summerfelt A, Buchanan RW, et al. (2010) Gamma and delta neural oscillations and association with clinical symptoms under subanesthetic ketamine. Neuropsychopharmacology 35: 632-640.

56. Pinault D. (2008) N-methyl d-aspartate receptor antagonists ketamine and MK-801 induce wake-related aberrant gamma oscillations in the rat neocortex. Biol Psychiatr 63: 730-735.

57. Chrobak JJ, Hinman JR, Sabolek HR. (2008) Revealing past memories proactive interference and ketamine-induced memory deficits. J Neurosci 28: 4512-4520.

58. Kocsis B. (2012) Differential role of NR2A and NR2B subunits in N-methyl-D-aspartate receptor antagonist-induced aberrant cortical gamma oscillations. Biol Psychiatr 71: 987-995.

59. Ma J, Leung LS. (2007) The supramammillo-septal-hippocampal pathway mediates sensorimotor gating impairment and hyperlocomotion induced by MK-801 and ketamine in rats. Psychopharmacology (Berl) 191: 961-974.

60. Hakami T, Jones NC, Tolmacheva EA, et al. (2009) NMDA receptor hypofunction leads to generalized and persistent aberrant gamma oscillations independent of hyperlocomotion and the state of consciousness. PLoS One 4: e6755.

61. Ehrlichman RS, Gandal MJ, Maxwell CR, et al. (2009) N-methyl-d-aspartic acid receptor antagonist-induced frequency oscillations in mice recreate pattern of electrophysiological deficits in schizophrenia. Neuroscience 158: 705-712.

62. Hunt MJ, Raynaud B, Garcia R. (2006) Ketamine dose-dependently induces high-frequency oscillations in the nucleus accumbens in freely moving rats. Biol Psychiatr 60: 1206-1214.

63. Kulikova SP, Tolmacheva EA, Anderson P, Gaudias J, Adams BE, Zheng T, et al. (2012) Opposite effects of ketamine and deep brain stimulation on rat thalamocortical information processing. Eur J Neurosci 36: 3407-3419.

64. Molina LA, Skelin I, Gruber AJ. (2014) Acute NMDA receptor antagonism disrupts synchronization of action potential firing in rat prefrontal cortex. PLoS One 9: e85842.

65. Homayoun H, Moghaddam B. (2007) NMDA receptor hypofunction produces opposite effects on prefrontal cortex interneurons and pyramidal neurons. J Neurosci 27: 11496-11500.
66. Callicott JH, Bertolino A, Mattay VS, Langheim FJ, Duyn J, Coppola R, et al. (2000) Physiological
dysfunction of the dorsolateral prefrontal cortex in schizophrenia revisited. *Cereb Cortex* 10: 1078-1092.

67. Corlett PR, Honey GD, Fletcher PC. (2007) From prediction error to psychosis ketamine as a
pharmacological model of delusions. *J Psychopharmacol* 21: 238-252.

68. Adell A, Jimenez-Sanchez L, Lopez-Gil X, et al. (2012) Is the acute NMDA receptor hypofunction a
valid model of schizophrenia? *Schizophr Bull* 38: 9-14.

69. Frohlich J, Van Horn JD. (2014) Reviewing the ketamine model for schizophrenia. *J
Psychopharmacol* 28: 287-302.

70. Gunduz-Bruce H. (2009) The acute effects of NMDA antagonism from the rodent to the human brain. *Brain Res Rev* 60: 279-286.

71. Canolty RT, Knight RT. (2010) The functional role of cross-frequency coupling. *Trends Cogn Sci* 14: 506-515.

72. Kirihara K, Rissling AJ, Swerdlow NR, et al. (2012) Hierarchical organization of gamma and theta
oscillatory dynamics in schizophrenia. *Biol Psychiatr* 71: 873-880.

73. Jensen O, Colgin LL. (2007) Cross-frequency coupling between neuronal oscillations. *Trends Cogn
Sci* 11: 267-269.

74. Lisman JE, Jensen O. (2013) The theta-gamma neural code. *Neuron* 77: 1002-1016.

75. Palenicek T, Fujakova M, Brunovsky M, et al. (2011) Electroencephalographic spectral and coherence
analysis of ketamine in rats correlation with behavioral effects and pharmacokinetics. *Neuropsychobiology* 63: 202-218.

76. Tsuda N, Hayashi K, Hagihira S, et al. (2007) Ketamine, an NMDA-antagonist, increases the
oscillatory frequencies of alpha-peaks on the electroencephalographic power spectrum. *Acta Anaesthesiol Scand* 51: 472-481.

77. Caixeta FV, Cornelio AM, Scheffer-Teixeira R, et al. (2013) Ketamine alters oscillatory coupling in
the hippocampus. *Sci Rep* 3: 2348.

78. Hiyoshi T, Kambe D, Karasawa J, et al. (2014) Differential effects of NMDA receptor antagonists at
lower and higher doses on basal gamma band oscillation power in rat cortical electroencephalograms. *Neuropsycharmacology* 85: 384-396.

79. Nicolas MJ, Lopez-Azcarate J, Valencia M, et al. (2011) Ketamine-induced oscillations in the motor
circuit of the rat basal ganglia. *PLoS One* 6: e21814.

80. Buzsaki G. (1991) The thalamic clock emergent network properties. *Neurosci* 41: 351-364.

81. Friston KJ. (2002) Dysfunctional connectivity in schizophrenia. *World Psychiatry* 12: 66-71.

82. Melillo R, Lisman G. (2009) Autistic spectrum disorders as functional disconnection syndrome. *Rev
Neurosci* 20: 111-131.

83. de Haan W., Pijnenburg YA, Strijers RL, et al. (2009) Functional neural network analysis in
frontotemporal dementia and Alzheimer's disease using EEG and graph theory. *BMC Neurosci* 10: 101.

84. Bokde AL, Ewers M, Hampel H. (2009) Assessing neuronal networks understanding Alzheimer's
disease. *Prog Neurobiol* 89: 125-133.

85. Popescu BO, Toescu EC, Popescu LM, et al. (2009) Blood-brain barrier alterations in ageing and
dementia. *J Neurol Sci* 283: 99-106.

86. Herrmann CS, Demiralp T. (2005) Human EEG gamma oscillations in neuropsychiatric disorders. *Clin Neurophysiol* 116: 2719-2733.

87. van Deursen JA, Vuurman EF, Verhey FR, et al. (2008) Increased EEG gamma band activity in
Alzheimer's disease and mild cognitive impairment. *J Neural Transm* 115: 1301-1311.
88. Yordanova J, Banaschewski T, Kolev V, et al. (2001) Abnormal early stages of task stimulus processing in children with attention-deficit hyperactivity disorder--evidence from event-related gamma oscillations. *Clin Neurophysiol* 112: 1096-1108.

89. Spencer KM, Nestor PG, Niznikiewicz MA, et al. (2003) Abnormal neural synchrony in schizophrenia. *J Neurosci* 23: 7407-7411.

90. Uhlhaas PJ, Singer W. (2006) Neural synchrony in brain disorders relevance for cognitive dysfunctions and pathophysiology. *Neuron* 52: 155-168.

91. Whittington MA. (2008) Can brain rhythms inform on underlying pathology in schizophrenia? *Biol Psychiatr* 63: 728-729.

92. Cronenwett WJ, Csernansky J. (2010) Thalamic pathology in schizophrenia. *Curr Top Behav Neurosci* 4: 509-528.

93. Ferrarelli F, Peterson MJ, Sarasso S, et al. (2010) Thalamic dysfunction in schizophrenia suggested by whole-night deficits in slow and fast spindles. *Am J Psychiatr* 167: 1339-1348.

94. Lisman JE, Pi HJ, Zhang Y, et al. (2010) A thalamo-hippocampal-ventral tegmental area loop may produce the positive feedback that underlies the psychotic break in schizophrenia. *Biol Psychiatr* 68: 17-24.

95. Pinault D. (2011) Dysfunctional thalamus-related networks in schizophrenia. *Schizophr Bull* 37: 238-243.

96. Watis L, Chen SH, Chua HC, et al. (2008) Glutamatergic abnormalities of the thalamus in schizophrenia a systematic review. *J Neural Transm* 115: 493-511.

97. Zhang Y, Su TP, Liu B, et al. (2014) Disrupted thalamo-cortical connectivity in schizophrenia a morphometric correlation analysis. *Schizophr Res* 153: 129-135.

98. Javitt DC. (2007) Glutamate and schizophrenia phencyclidine, N-methyl-D-aspartate receptors, and dopamine-glutamate interactions. *Int Rev Neurobiol* 78: 69-108.

99. Moghaddam B. (2003) Bringing order to the glutamate chaos in schizophrenia. *Neuron* 40: 881-884.

100. Gandal MJ, Edgar JC, Klook K, et al. (2012) Gamma synchrony towards a translational biomarker for the treatment-resistant symptoms of schizophrenia. *Neuropharmacology* 62: 1504-1518.

101. Rolls ET, Loh M, Deco G, et al. (2008) Computational models of schizophrenia and dopamine modulation in the prefrontal cortex. *Nat Rev Neurosci* 9: 696-709.

102. Winterer G, Ziller M, Dorn H, et al. (2000) Schizophrenia reduced signal-to-noise ratio and impaired phase-locking during information processing. *Clin Neurophysiol* 111: 837-849.

103. Llinas RR, Ribary U, Jeantomond D, et al. (1999) Thalamocortical dysrhythmia A neurological and neuropsychiatric syndrome characterized by magnetoencephalography. *Proc Nail Acad Sci USA* 96: 15222-15227.

104. Gonzalez-Burgos G, Lewis DA. (2008) GABA neurons and the mechanisms of network oscillations implications for understanding cortical dysfunctions in schizophrenia. *Schizophr Bull* 34: 944-961.

105. Roopun AK, Cunningham MO, Racca C, et al. (2008) Region-specific changes in gamma and beta2 rhythms in NMDA receptor dysfunction models of schizophrenia. *Schizophr Bull* 34: 962-973.

© 2014, Pinault D, licensee AIMS Press. This is an open access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0)