In waking and sleeping states, thalamocortical system generates a variety of oscillations ranging from 0.1 Hz to hundreds of Hz. Most of them are present during NREM sleep, but slower activities prevail in this state of vigilance. Thalamocortical network is organized in a loop in which thalamocortical cells excite reticular thalamic and neocortical cells, reticular thalamic cells inhibit thalamocortical cells and corticothalamic cells excite thalamocortical and reticular thalamic cells. Despite stable anatomical connectivity, different types of oscillations preferentially originate either in neocortex or in thalamus.

During sleep stage 2, spindle oscillation (9–15 Hz) is a dominant type of activity. It is well accepted that spindles originate in the thalamus via interplay of firing of reticular thalamic and thalamocortical neurons, but neocortex controls spindle generation. Spindles can be divided on fast and slow. Several properties of slow spindles do not match known mechanisms of their thalamic origin. Slow oscillation (about 1 Hz) dominates slow-wave sleep stage. Each slow wave is composed of hyperpolarized or silent and depolarized and active state. Active states may be accompanied by spindles and higher frequency activities. Slow waves originate mainly in deep cortical layers from which they propagate to more superficial layers and they also propagate horizontally. Full expression of slow wave activities requires the presence of thalamus, although slow oscillation can be recorded in athalamic preparations.

Therefore, despite the fact of preferential origin of different sleep oscillations in either neocortex or in thalamus, only the full thalamocortical network can generate sleep activities with known properties. Support: CIHR and NSERC.

**Electropsychopharmacology: applying EEG and ERP to neurocognitive mechanisms that underlie pharmacological modulation of cognitive processes, or they may provide additional sensitivity to detect neurocognitive effects that are not readily observable in behavioral measures. This will be illustrated by means of pertinent examples. These include elucidating the mechanisms of stimulant action mediating deficient impulse control and the role of the cannabinoid system in human working memory, as well as drug effects on sensory gating and specific aspects of visual-spatial attention. Other examples concern the added sensitivity of EEG and ERP measures, relative to that of performance measures, in detecting effects of alcohol, and more generally in monitoring and predicting vigilance and the ability to detect external signals in the immediate future. Relations between brain signals and cognitive competences are revealed by either comparing different individuals, or moment-to-moment fluctuations within individuals, or differences in state (e.g., drug-induced) within individuals.

**EEG and ERP as key techniques for functional brain alterations studies**

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A3

Behavioral studies in rodents are basically based on inferring cognitive processes out of locomotor activity. In other words, we evaluate memory processes during Morris water maze, or Novel object recognition based on the time spent of the given subject in close proximity of an item, platform, or the time required to reach or leave an area of the cage/maze. By the mean of automatic scoring systems (e.g.: Ethovision, AniMaze, etc.) we are provided by objective measurements, which should be in any case interpreted by the researcher. There is always a lack of direct measurement of the cognitive processes. Integration of behavioural scoring with electrical activity evaluation of different areas involved in cognitive processing can be a useful tool, to provide scientists further parameter helpful in data interpretation. We will thus discuss the integration of EEG analysis in Alzheimer’s mild cognitive impairment and acoustic ERPs in AD and epilepsy.

**Targeting EEG network oscillations: translational opportunities for drug discovery science in psychiatric and neurodegenerative disorders**

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A4

Despite decades of research in psychiatric and neurodegenerative disorders, the attrition rate in clinical trials and late-stage drug discovery programs for the development of novel agents for disease interception has been unacceptably high. The major issue facing neuroscience drug discovery is that drugs that show good effectivity in preclinical models often fail to meet clinical trials endpoints. The
limitations of the traditional animal-based assays prompted a resurgence of interest in rethinking animal models and their predictability and translational validity in translational neuroscience. Better translation of a biomarker and endophenotype of the disease might rapidly provide information regarding the effects of drugs on the underlying disease biology, bridge the translational gap and potentially lower the rate of clinical trial attrition. An increasing number of experimental and clinical observations suggest that those chronic brain disorders arise from structural alterations in neuronal circuits, and therefore focus has been shifted towards investigation of electrophysiological correlates of the molecular pathology, with emphasis on neural oscillations and connectivity as promising candidate biomarkers of neuronal disorders. State-of-the-art examples of pharmacological studies modeling abnormal network oscillations and disturbed connectivity of several CNS disorders will be discussed.

A5
Source localization using LORETA software
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A5

The training course will be dedicated to the usage of LORETA software for localization of neuronal sources of EEG activity. After a brief introduction into the underlying physiology and theoretical assumptions of source localization techniques, an example of how to apply the software to EEG recordings will be shown, including limitations and caveats. Further focus will be on connectivity measures between intracortical areas as well as a short overview on statistical analysis implemented in the LORETA software.

A6
EEG Analysis methods beyond standard assessments
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A6

Data analysis methods for EEG have shown great progress over the last two decades, partially inspired or driven by methodological advancements in adjacent research methods such as MEG, intracranial recordings and functional MRI. In this educational session I will go over the analysis methods that can be considered part of the traditional repertoire for EEG assessments and will extend that by highlighting some recent methodological advancements for data processing. Among others, power spectral analysis techniques using multitapers, statistical approaches based on non-parametric hypothesis, cluster based inference, robust statistics and source reconstruction techniques based on spatial filtering will be explained. The focus will be on introducing these techniques in an intuitive manner and providing pointers to data analysis tools that implement them.

A7
Personalized medicine in Depression and ADHD: Introduction and EEG biomarkers for predicting treatment outcome
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A7

At present stimulant medication, antidepressants and behavior therapy are the most often applied and accepted treatments for ADHD and Depression. However, recent large-scale studies and meta-analyses have demonstrated limitations of these treatments including reduced long-term efficacy of stimulant medication, limited efficacy of antidepressant medications and overall limited efficacy of behavioral interventions on the group level. It hence becomes obvious there is a need for a re-conceptualization of psychiatric disorders along the lines of NIMH proposed Research Domain Criteria (RDoC) or referred to as biomarkers or personalized medicine. Personalized Medicine aims to prescribe the right treatment, for the right person at the right time as opposed to the currently employed ‘one-size-fits-all’ approach. This development relies on identification of subgroups using biomarkers. This presentation will summarize the current status of EEG based personalized medicine and present new results from large biomarker studies in depression and ADHD focused on resting-state EEG. Several results from the iSPOT study (international Study to Predict Optimized Treatment) in Depression and ADHD will be presented [1,2,3,4,5]. In iSPOT-D, 1008 depressed patients are randomized to Escitalopram, Sertraline or Venlafaxine and in iSPOT-A 336 ADHD patients are prescribed with methylphenidate and patients were assessed at baseline on resting-state EEG and other measures. Several promising biomarkers that can predict treatment response and remission using baseline biomarkers will be presented and the importance of gender differences will be discussed in more detail.

References

A8
Clinical applications of EEG in psychiatry
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A8

Starting with an outline of the evolving discipline of Clinical Psychiatric Electrophysiology a discussion of the value of diagnostic tests in general is given. The presentation then focuses on the well-established clinical indications of the standard EEG in the day to day practice of clinical psychiatry. Discussion will cover panic disorder, autistic spectrum disorders, and repeated aggression in some detail. Case vignettes are included to generate interactive discussion.

Keynotes
A9
The genesis of EEG phenomena: hot topics of the last decade
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A9
Since the turn of the century, the scope of EEG investigations became much broader, in particular due to the possibility of recording the full, physiologically relevant range of brain activities from the infra-slow to the high frequency spectral range, making use of wide dynamic direct-current (DC) coupled amplifiers, and of accurate recordings of high frequency oscillations up to hundreds of Hz. This has been denominated full-band or wide-band EEG. In this lecture, however, I focus on the high frequency EEG/MEG phenomena or High Frequency Oscillations (HFOs). These phenomena cover a number of activities that range from 60 – 80 Hz to approximately 500 Hz. Interest for these phenomena has gained momentum in the last decade. They appear in the healthy brain associated with sensory, motor and cognitive events, and also in pathological cases, particularly in epilepsy. Under the concept HFO, activities in the gamma band (30 – 70 Hz) occupy a prominent place. A variety of names are used to describe physiological EEG/MEG activity above 70 Hz such as high-gamma and the chi-band; in the context of epilepsy specific types of oscillations have been described, namely ripples (≈90 – 200 Hz) and fast ripples (≈200Hz), but the corresponding frequency boundaries are rather fuzzy [1]. Here I will discuss neuronal processes generating HFOs [1,2], a few relevant applications in cognitive neuroscience and in epilepsy [3], and practical questions regarding the possibilities and difficulties of reliably recording HFOs in human, and distinguishing the latter from artifacts [4].

**References**


**A10 Cellular basis of sleep slow oscillation: What is clear, what is not**

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The major type of activity generated by the thalamocortical system during slow-wave sleep is the slow oscillation, composed of slow waves repeated with a frequency of about 1 Hz. Each slow wave is comprised of hyperpolarized silent state, often called down state, and depolarized active state, often called up state. During wake and REM sleep cortical neurons remain in active state. What is likely known on slow wave generation: (a) Slow oscillation is essentially cortical in origin, but it is modulated by thalamic activities. (b) Cortical slow waves in adults start more in frontal areas and propagate to other cortical areas, but multiple slow waves recorded throughout cortical mantle remain local. (c) In ferrets, mice and cats slow waves start mainly in layer 5, but in epileptic patients they originate around layer 3. (d) Silent states of slow oscillation are essentially periods of disfacilitation, but GABAergic activities can be detected in a subset of neurons prior to the onset of silent states. (e) Active states are dominated by excitatory and inhibitory synaptic activities; in anesthetized animals, these activities are balanced, in sleeping animals inhibition largely dominates active states. What we don’t know: (a) What triggers the onset of active state? Three hypotheses are present: (1) Stochastic summation of spike-independent minis occasionally leading to the first spike that engages the whole network. (2) Intrinsic activity of layer 5 pyramidal cells (h-like current). (3) Self-organized onset of activity in groups of neurons. (b) What terminates active states? Following hypotheses are proposed: (1) Intrinsic neuronal firing frequency accommodation that decreases the overall excitatory drive. (2) Use-dependent synaptic depression. (3) Activity of Na+- and Ca2+-dependent K+ current. (4) Extracortical signaling. All these mechanisms are present during wake and REM sleep, but why active states are not terminated in these states of vigilance remains unclear.

**A11 GABAergic modulation of neuronal oscillations in animals and humans and its consequences for working memory**

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A11

**Background**

Networks in the brain must rely on powerful mechanism for routing and prioritizing information processing. In a larger set of attention and memory studies we have investigated the notion that alpha oscillations (9 – 12 Hz) are inhibitory and serve to route the information flow: ‘gating by inhibition’ [1]. The alpha band activity is under top-down control by areas in the dorsal attention network. As such the alpha band activity – previously believed to reflect a state of rest - serves an important role for shaping the functional architecture of the working brain. Gamma band activity (50 – 100 Hz) reflects feed-forward processing and is modulated by the alpha oscillations. In animals it has been demonstrated that GABAergic interneurons play an important role for synchronizing neural populations [2]; however, it remains unknown if these mechanistic principles generalize to human oscillations.

**Methods**

To investigate how GABAergic modulated affects gamma oscillations, we recorded ongoing brain activity using magnetoencephalography (MEG) in human subjects participating in a double-blind pharmacological study receiving placebo and lorazepam. Lorazepam is a benzodiazepine upregulating GABAergic conductance. This was done in participants while they performed a visuospatial working memory (WM) task.

**Results**

The key finding was that occipital gamma power associated with WM recognition increased with lorazepam dosage [3]. In addition, the frequency of the gamma activity decreased with dosage. This is consistent with models derived from the rat hippocampus. With respect to oscillations in the alpha band, we observed a parametrical decrease with drug dosage that also predicted a performance decrease. This is consistent with alpha oscillations reflecting functional inhibition.

**Conclusion**

We conclude that GABAergic interneurons are implicated in the generation of gamma and alpha oscillations in humans. As we will discuss these findings allow us to link neuronal dynamics to behavior in humans by embracing established animal models.

**References**


**A12 Refining brain oscillatory targets for intervention in ADHD**

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A12

**Background**

Brain oscillatory patterns are increasingly used as biological markers of psychiatric disease, developmental course, and treatment response. In
order to maximize effectiveness, advanced approaches should enable a mechanistic understanding of how brain rhythms carry out the cognitive and emotional processes that, when disordered, may lead to mental disorders. This will aid in the identification of interventions that show evidence of mechanism and target engagement and will increase our understanding of how efficacious interventions achieve their effect within clinical populations.

In this presentation, important considerations in the process of identifying and refining potential brain oscillatory targets that may be useful for treatment monitoring and response will be presented. For example, approaches to characterizing the significant variability and heterogeneity that exists within ADHD and typically developing populations are discussed. In addition, the need for refined measurements and signal processing techniques that increase the signal-to-noise ratio are described. Finally, issues such as targeting cognitive dysfunction and/or modeling developmental changes are considered.

We then describe how this approach has been implemented to identify and validate brain targets (biomarkers) in clinical trials for children with attention-deficit/hyperactivity disorder (ADHD). In addition, these data will be used to illustrate potential applications for neuromodulation approaches in ADHD as well as other neurodevelopmental disorders such as autism spectrum disorder (ASD) and Tourette’s Syndrome.

Relevance and Implications for future research

The data presented suggest that EEG-based biomarkers may be useful indices of developmental course of disorder, behavioral and cognitive functioning, and prediction of treatment response. Although the clinical utility of EEG measures is promising, more research is needed before these findings can be implemented in clinical practice.

A13

Elucidating mechanisms of sleep-wake regulation in humans with pharma-genetic tools

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A13

Epidemiological studies demonstrate that sleep-wake disorders are highly prevalent in society and rank third in the prevalence of all brain diseases. The normal alternation between sleep and wakefulness is tightly regulated, and prolonged EEG recordings show that specific sleep and wake states reflect highly complex behaviors. Little is currently known about the molecular underpinnings of physiological sleep-wake regulation and functions. To foster our knowledge of the pathophysiology of sleep-wake disorders and their possible rational treatment, a molecular understanding of sleep-wake regulatory processes is indispensable. Accumulating evidence suggests that important aspects of sleep-wake regulation in animals and humans are genetically controlled and, thus, have a molecular basis. Consistent with this view, the combination of neurophysiologic, genetic and pharmacologic tools revealed specific roles for adenosine, dopamine and glutamate receptors and metabolic pathways in sleep-wake regulation. These studies also showed that functional allelic variation in candidate genes can profoundly affect functional aspects of sleep and wakefulness, even in healthy humans and under physiological conditions, as well as modulate individual responses to hypnotic and wake-promoting agents. These insights may provide a rationale for personalized sleep-wake pharmacotherapy (Holst et al., Annu Rev Pharmacol Toxicol, 2016). In the future, together with novel ‘omics’-studies of sleep in health and disease, they may pave the way for the discovery of new evidence-based treatments of sleep-wake pathologies such as insomnia and the pharmacological enhancement of sleep-associated brain functions such as neuronal plasticity.

Research supported by the Swiss National Science Foundation (grant # 320030_135414 & 320030_163439) and the Clinical Research Priority Program “Sleep and Health” of the University of Zürich.

Symposia

A14

The dos and don’ts for electrophysiological connectivity analysis

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A14

In recent years it has been increasingly recognized that insight into the dynamics of interareal interactions is crucial for our understanding of normal and pathological brain function. Methodological developments and open source availability of advanced analysis tools have enabled the wider neuroscientific community to estimate a wide range of connectivity metrics from non-invasively obtained electrophysiological signals. Next to deciding on an appropriate analysis strategy, researchers are faced with the challenge to correctly interpret their findings. Volume conduction and electromagnetic field spread cause neuronal signals to be picked up by multiple channels at once, causing spurious estimates of connectivity. Comparison across experimental groups and conditions may be confounded by differences in univariate signal properties such as signal-to-noise ratio. I will illustrate some of these interpretational pitfalls and provide some recommendations that may need to be taken into account to improve the validity of the interpretation of EEG/MEG connectivity studies.

A15

MEG as a routine diagnostic tool in memory clinic patients

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A15

Background

Electro-encephalography (EEG) has been used as a routine diagnostic tool for patients of the Alzheimer Center Amsterdam since 2001 [1]. Recently, EEG has partly been replaced by magneto-encephalography (MEG), because it may be more sensitive for pathology in disease specific regions. We investigated its discriminative value between Alzheimer’s disease (AD) and subjective cognitive decline (SCD) using a machine learning approach.

Methods

MEG was recorded in an unselected proportion of memory clinic patients as part of a routine workup. MEG data were co-registered with a head-size matched template-MRI and source-reconstructed by projection onto 90 AAL-regions using beamforming [2, 3]. Clinical reports were made using visual and spectral analyses, blinded for clinical information. Diagnoses were made in a weekly multidisciplinary meeting using full clinical information and additional investigations, such as MRI and neuropsychological examination. The first 20 AD and 20 SCD patients were further analysed. Directed connectivity (directed phase transfer entropy [dPTE]) and minimum spanning tree (MST) based network measures (8–13 Hz band) were calculated per region [3–5], where the imbalance in information flow between regions was used to construct the MST. Combinations of MEG measures at eight AD-specific regions (left and right hippocampus, parahippocampal gyrus, precuneus, cuneus) were entered into random forest models to classify between patient groups.

Results

From April 2015 to July 2016, 101 patients received an MEG. Diagnoses were AD (n = 26); SCD (n = 24); psychiatric disorder (n = 18); mild cognitive impairment (n = 10); fronto-temporal dementia (n = 7); Lewy body dementia (n = 5); vascular dementia (n = 1); and other/
postponed diagnosis \( (n = 10) \). One patient’s MEG diagnostic report could not be made because of movement artefacts. Her MRI and neuropsychological examination could also not be completed and she was diagnosed with severe AD based on clinical information and cerebrospinal fluid biomarkers. In the distinction between AD (age 64.8 ± 7.9, 50 % female) and SCD (age 61.4 ± 21.8, 55 % female), a random forest model with relative theta power of the eight AD regions yielded an accuracy of 0.810. Addition of dPTE for these regions increased the accuracy to 0.843. When network measures (leaf fraction, diameter, tree hierarchy) were added to the model with theta power and dPTE an accuracy of 0.812 was found.

Conclusion
Routine diagnostic MEG is feasible in a memory clinic screening and has a high accuracy in the discrimination between AD and SCD using theta power in AD-specific regions. Directed connectivity has modest additional diagnostic value whereas network measures did not add to the diagnostic accuracy.

References

Identification of the epileptogenic zone using MEG network analysis
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A16

Introduction
Epilepsy is increasingly seen as a brain network disorder [1–3]. Patients with epilepsy have been shown to have different networks compared to healthy controls, deviating from the optimal configuration and with abnormal network hubs [4–8]. A potent treatment for pharmacoresistant epilepsy is epilepsy surgery. The goal of epilepsy surgery is to remove or disconnect the epileptogenic zone, which renders the patient seizure-free [9]. A hypothesis about the location of the epileptogenic zone can be based on techniques such as electroencephalography (EEG) and magnetoencephalography (MEG). However, establishing a hypothesis is challenging and not always successful as currently one third of the patients continue to experience seizures after surgery [10]. New methods are therefore needed to generate more accurate hypotheses about the location of the epileptogenic zone such that more patients become seizure-free. Our aim was to develop such a new method based on network theory. We hypothesized that the epileptogenic zone coincides or connects with hubs and information senders in the network.

Methods
We analyzed eyes-closed resting-state MEG recordings of 22 patients with pharmacoresistant epilepsy. The time series in source space (virtual electrodes) were reconstructed using beamforming for 90 regions of the AAL atlas [11]. We estimated functional connectivity between those regions using phase lag index (PLI) [12] in the broad-band (0.5–48Hz). We used 20 epochs of 3.28 s each without artefacts or epileptiform activity. We generated the minimum spanning tree based on the PLI and calculated the betweenness centrality (an indicator of hubs) for each region. Furthermore, we assessed effective connectivity (an indicator of information senders) using the directed phase transfer entropy (dPTE) [11] for different frequency bands.

Results
ROIs with high broadband betweenness centrality (hubs) coincided with the resection cavity (or resection lobe) in 8/14 (9/14) seizure-free patients and in 0/8 (0/8) patients with remaining seizures (73 % (77 %) accuracy). For the effective connectivity, high dPTE values coincided with the resection cavity (or resection lobe) in 8/14 (10/14) seizure-free patients and only in 2/8 (2/8) patients with remaining seizures (64 % (73 %) accuracy) in the delta band (0.5–4Hz).

Implications
Hub regions and strong senders are markers of the epileptogenic zone. These results are a first step towards a localization method that can be applied to MEG recordings even in the absence of epileptiform activity, yielding an improvement in localization and finally surgery outcome.

References

EEG biomarker integration for better decision making in clinical trials
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A17

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Brain disorders are a huge burden on the health care system, key issues being inaccurate diagnosis and insufficient treatment options. Hence, there is an urgent need for biomarkers that monitor disease status or therapeutic response. Current biomarkers lack the desired accuracy, because of the large variability in healthy subjects and the often subtle disease-related changes. In EEG, however, pathophysiology is often expressed in multiple ways. Here we show that an integrative approach in which any biomarker that carries complementary information about a disease or therapeutic intervention can result in an accurate diagnostic index for better decision making in clinical trials.

Recently, we showed that EEG biomarker integration improves the prediction of conversion from mild cognitive impairment to Alzheimer’s disease (AD) compared with a single-biomarker based prediction [1]. The integrative biomarker index can be used for stratification of patients at recruitment in clinical studies and for documenting and quantifying effects of intervention. Here, we provide additional proof-of-concept that EEG-based prediction can be improved with the integrative biomarker approach in clinical trials where a drug is tested in a scopolamine challenge model in healthy subjects. Scopolamine is the most extensively studied and used model for cognitive impairment and resembles the changes seen in AD patients [2]. It is used in drug development to demonstrate the reversal of the temporary scopolamine-induced cognitive deficits by a cognition enhancing compound. For this purpose, we have developed an integrative EEG biomarker index (mAChR index) that is optimally sensitive to the CNS effects of scopolamine, to objectively determine whether reversal of scopolamine effects by a cholinergic compound is successful. The mAChR index yielded higher classification performance than any individual EEG biomarker with accuracy, sensitivity, specificity and precision of 90%. This significantly outperforms the single-best EEG biomarker (relative delta power). Validation on an independent dataset indicated the robustness of the index. To examine the validity of scopolamine as a cognitive impairment model, we applied this integrative index on healthy elderly controls and Alzheimer’s patients and observed that this index indeed differentiates patients from controls.

We address this by using novel features of the Neurophysiological Biomarker Toolbox (http://www.nbtwiki.net/), which employ data mining algorithms to combine the information from multiple biomarkers. Our results demonstrate that integrating information from multiple EEG biomarkers better captures the unique phenotype of an individual patient and is a promising approach to enhance accuracy and reduce the multiple-comparisons problem when using EEG in clinical trials.

References

A18 Continuous EEG and deep learning for outcome prediction in postanoxic coma
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A18

Introduction
Reliable outcome prediction in approximately 50% of patients with a postanoxic encephalopathy is possible with visual interpretation of continuous EEG recorded within 24 h after cardiac arrest [1-6]. To assist in the visual assessment, we developed the Cerebral Recovery Index [7] and, more recently, a random forest classifier, showing similar performance for outcome prediction as visual assessment of the EEG [8]. Deep Learning may advance the prognostic value of EEG significantly, in part as it does not depend on ‘hand-made’ features [3, 9].

Methods
We used data from the EEG database of the Medisch Spectrum Twente and Rijnstate hospitals with recordings from patients treated in the Intensive Care Unit with a postanoxic encephalopathy after a cardiac arrest. EEGs were recorded with twenty-one silver/silver chloride cup electrodes placed on the scalp according to the international 10–20 system using a Neurocenter EEG recording system (Clinical Science Systems, Voorschoten, The Netherlands) or a Nihon Kohden system (VCM Medical, the Netherlands). Neurological outcome (Cerebral performance category scores) was dichotomized as good (no or mild neurological impairment) or poor (severe neurological impairment, vegetative state or death) at 6 months after cardiac arrest.

We implemented a convolutional neural network (CNN) in python with TensorFlow on a CentOS system with the NVIDIA GTX-1080 as GPU. The input layer had dimensions 128x19 to process the raw 19-channel EEG. EEGs were analyzed using non-overlapping 2 s epochs using 5 min segments at each hour after cardiac arrest. For each patient in the validation set, we calculated the percentage of 2 s epochs within the 5 min segment that is predicted as poor neurological outcome. Using ROC curves the threshold at which poor outcome could be predicted with 100% specificity was determined.

Results
After training with 131 EEGs, evaluation in an independent set with 33 patients showed that poor outcome could reliably be predicted in 67% of the patients, without false positives (specificity 100%) at t = 12 h after cardiac arrest; poor outcome prediction at a later instance (t = 24) was not possible with a specificity of 100%.

Discussion and Conclusion
We show feasibility of CNN to process EEG in patients with a postanoxic coma for prognostication. Pilot results show high predictive value for poor neurological outcome. As temporal evolution of EEG patterns in these patients is significant [2], recurrent neural nets may outperform convolutional networks. To understand the discriminating features, we currently explore methods for interpretation and visualization of networks.

References
A19

Prospects and challenges of Alzheimer’s classification using resting-state EEG rhythms
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A19

Background and aim
In the European FP7 “DECIDE” project (www.eu-decide.eu), a computing-grid infrastructure was developed to compute electroencephalographic (EEG) and other biomarkers for diagnosis and instrumental assessment of patients with Alzheimer’s disease (AD). In the framework of that project, previous evidence showed a 75.5% best accuracy in the classification of 120 Alzheimer’s disease (AD) patients with dementia and 100 matched normal elderly (Nold) subjects based on cortical source current density and linear lagged connectivity estimated by eLORETA freeware from resting state eyes-closed electroencephalographic (rsEEG) rhythms (Babiloni et al., 2016). Specifically, that accuracy was reached using the ratio theta/alpha1 rhythms (Babiloni et al., 2016). It also showed an accuracy of 72% using the most 4 discriminative rsEEG markers of source current density and linear lagged connectivity estimated by eLORETA freeware from healthy volunteers at 3 timepoints: baseline, 3, and 9 h following 1 of 4 doses of basmisanil (2x15, 2x60, 3x130, 3x1250 mg). A separate EEG study in 12 volunteers measured at baseline, midazolam (5 mg), and 14 days of basmisanil treatment (240 mg, bid).

Results
Basmisanil (ROS186582, RG1662) bound to cloned human GABA-A alpha5 subunit-containing receptors shows target and neuronal circuit engagement in man
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A20

Background
Inhibitory GABAergic signaling plays a key role in brain function. Drugs that enhance GABA-A receptor function (e.g. Benzodiazepines) are widely used to treat conditions such as anxiety, insomnia, and epilepsy. In contrast, no inhibitors of GABA-A receptors exist for clinical use. Preclinical animal studies suggest that releasing inhibition by selectively inhibiting GABA-A alpha5 subunit-containing receptors may be beneficial in conditions of impaired cognition such Down syndrome, Schizophrenia, and Alzheimer’s disease, and may also promote functional recovery after ischemic stroke, importantly without the side effects associated to non-selective inhibitors. Here we characterize basmisanil, a novel selective negative allosteric modulator of GABA-A alpha5 receptors, in terms of in vitro pharmacology as well as receptor occupancy and EEG signature in healthy volunteers.

Methods
Radioligand binding (3H-flumazenil) and voltage-clamp electrophysiology experiments were conducted in vitro on GABA-A receptors expressed in HEK293 cells and Xenopus oocytes to demonstrate binding affinity and functional selectivity for the GABA-A alpha5 vs. alpha1/2/3 subunit-containing receptors. A receptor occupancy study with the GABAA a5 PET tracer [11C]Ro15-4513 was conducted in 10 healthy volunteers at 3 timepoints: baseline, 3, and 9 h following 1 of 4 doses of basmisanil (2x15, 2x60, 3x130, 3x1250 mg). A separate EEG study in 12 volunteers measured at baseline, midazolam (5 mg), and 14 days of basmisanil treatment (240 mg, bid).

Results
Basmisanil is a highly selective GABA-A alpha5 negative allosteric modulator that reaches the desired target with good safety and tolerability, and modulates neuronal activity in humans. These data suggest basmisanil as a promising candidate drug for further clinical
testing in conditions which may benefit from a reduction in excessive GABA-mediated tonic inhibition, such as cognitive impairment and stroke recovery.

Competing interests
All authors are or were employees of F. Hoffmann-La Roche Ltd.

A21
EEG cross-frequency coupling associated with attentional performance: an RDoC approach to attention
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A21

The quality of attentional performance plays a crucial role in goal-directed behavior in daily life activities, cognitive task performance, and in multiple psychiatric illnesses. The Research Domain Criteria (RDoC) approach put forward by the National Institute of Mental Health aims to investigate cognitive constructs while abandoning the conventional diagnostic system of psychiatric illnesses. The current study used an RDoC approach to investigate functions underlying attentional performance.

One of the previously postulated physiologic mechanisms that could explain variance in attentional performance is the quality of interplay between neuronal networks1,2. Various attempts to visualize this interplay have been made using different approaches. In our current study, we aimed to validate the approach of functional Independent Component Analysis (fICA) based on electroencephalograms (EEGs) for this purpose. This method yields components that reflect EEG cross-frequency coupling patterns between networks (details about the method can be found elsewhere3).

We first applied fICA to combined Eyes Open resting state EEG and EEG during an n-back task data in a large sample of healthy adults (n = 1397), yielding 32 components. Secondly, we obtained individual component loadings for every subject for the two conditions as well as a difference loading score (Loadingtask-LoadingEO) per network. Thirdly, we operationalized attentional performance by differentiating between attenders (n = 704) versus non-attenders, (n = 320) on the n-back task and found a significant difference between groups for the difference loading score for component 10. We proposed that component 10 reflects the anti-correlated interaction of an attention network and a resting state network. This finding was cross-validated in an adolescent Attention-Deficit/Hyperactivity Disorder (ADHD) population (n = 80), clinically suffering from attentional problems. As expected, the difference loading scores in this group were similar to the pattern observed in non-attenders. Furthermore, it was accompanied by a lower overall loading on component 10 in both conditions.

The current findings seem to validate fICA as a method to visualize neuronal networks and their interactions. Combining this method with objective behavioral measures may contribute to the understanding of brain mechanisms involved in attention and attentional problems such as observed in multiple psychiatric illnesses.

References

A22
Attention for inhibition in ADHD: new insights with ERP source imaging
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A22

Background
Deficits in response inhibition figure prominently in models of ADHD and have been documented in cognitive [1], ERP [2] and fMRI studies [3]. Parallel to these developments, some authors have criticized the inhibition model of ADHD and associated methodology, suggesting that attentional factors confound former results [4]. In a previous fMRI study [5] we aimed to control for attentional confounds during a stop-signal task (SST). Despite this modified SST, we found evidence for reduced activation in key-areas of the inhibition network, such as the right inferior frontal gyrus (rIFG), supplementary motor area (SMA) and anterior cingulate cortex (ACC). However, according to Barkley [6], inhibition problems precede other cognitive dysfunctions, such as attentional deficits. In order to investigate this hypothesis at the brain level, both high spatial and temporal resolution are needed, which have not yet been fully integrated in one imaging technique. In the current study [7], we addressed this issue by localizing ERP components associated with response inhibition in children with ADHD.

Methods
Dense array ERPs (128 electrodes) were obtained for 46 children with ADHD and 51 controls during the SST. Early and late components were compared between groups. N2 and P3 components were localized with LAURA distributed linear inverse solution for each participant, and statistically compared between groups (Bonferroni-corrected based on the number of electrodes, with p = 0.05/128 = 0.0004).

Results
A success-related N1 modulation was only apparent in the ADHD group. N2 and P3 amplitudes were reduced in ADHD. During the successful inhibition N2, the ADHD group showed reduced activation in rIFG, SMA, and right temporoparietal junction (rTPJ), and during failed inhibition in the rIFG. During the successful inhibition P3, reduced activation was found in ACC and SMA.

Conclusions
Source localization of N2 revealed not only a typical inhibition network (rIFG and SMA) that was affected, but also a major hub of the ventral attention system, the rTPJ. The ventral attention system supports attentional reorienting to salient and behaviourally relevant external stimuli. The fact that this ventral attention network is implicated in the same 50 ms time window (240-290 ms after stop stimulus) as the inhibition network creates a challenge to Barkley’s theory of ADHD.

Competing interests
The author(s) declare no potential conflicts of interest

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The findings do not yet allow for clear recommendations as to which participants will benefit from neurofeedback beyond practical considerations. Several predictive EEG markers have been proposed but await replication.

**A23 Neurofeedback and pharmacological treatments in ADHD - evidence and EEG-markers**

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A23

**Scientific background/Introduction**

Neurofeedback is a promising treatment for ADHD despite recent evidence cautioning that probably blinded ratings draw a less positive picture.

**Theoretical framework/Hypothesis/Purpose of the work**

The current presentation will focus on how recent metaanalytic evidence for neurofeedback in ADHD, address specifically studies on the add-on use and interactions with pharmacological treatment, and on the predictive EEG-based markers for neurofeedback.

**Used Methods and Materials**

Review of evidence from recent metaanalysis during the last 8 years [1–3] and selected individual papers on the relation to pharmacological treatment, EEG markers [4, 5] and mechanism of learning self-regulation.

**Findings**

The clinically relevant efficacy of neurofeedback is reduced in probably blinded ratings and compared to active or sham control conditions. This may reflect considerable unspecific effects, compromised neurofeedback quality, or lack of learning self-regulation. How neurofeedback depends on previous or concurrent pharmacological treatment is unclear. Some head to head studies report comparable or additive effects of neurofeedback and medication, but not that neurofeedback works best as an add-on or second stage treatment (beyond practical considerations). Several predictive EEG markers have been proposed but await replication.

**Discussion of Relevance and Implications for future Research**

The findings do not yet allow for clear recommendations as to which patients profit most from neurofeedback alone or in combination pharmacological intervention. Studies on stepped care approaches and personalized approaches – what works for whom – are urgently needed.

**References**


**A24 Background**

Attention-deficit hyperactivity disorder (ADHD) is characterized by an inappropriate pattern of inattentiveness. Increasing evidence demonstrates that the modulation of alpha oscillations plays an important role in the allocation of attention. A failure to modulate alpha activity might therefore reflect ADHD. The first study presented here aimed to investigate alpha modulation in children with ADHD during attentional performance. The second study aimed to replicate and extend previous findings with respect to electroencephalographic (EEG) biomarkers that have shown promise in predicting treatment outcome to stimulant medication in ADHD.

**Methods**

For the first study [1], posterior alpha activity (8–12 Hz) was measured in 30 healthy children and 30 children with ADHD aged 7–10 years, using EEG while they performed a visuospatial covert attention task. We focused the analyses on healthy boys (N = 9) and boys with ADHD (N = 17). For the second study [2], data from the international Study to Predict Optimized Treatment Response in ADHD (SPOT-A), 336 children and adolescents with ADHD were included and prescribed methylphenidate, and 158 healthy children were included. Treatment response was established after six weeks using the clinician rated ADHD-Rating Scale IV (ADHD-RS-IV). Responders to treatment were defined as >25 % improvement. The EEG Theta/Beta ratio (TBR) and alpha peak frequency (APF) were investigated as predictors for treatment outcome.

**Results**

In the first study, alpha activity in typically developing boys was similar to previous results of healthy adults: it decreased in the hemisphere contralateral to the attended hemifield, whereas it relatively increased in the other hemisphere. However, in boys with ADHD this hemispheric lateralization in the alpha band was not obvious (group contrast, p = .018). In the second study, male-adolescent non-responders exhibited a low frontal APF (ES = 0.83), whereas no differences in TBR were found between responders and non-responders. 62 % of the ADHD group was classified as a responder. Responders were more often males (63 % versus 51 %, p = .031), but did not differ from non-responders in age, medication dosage, and baseline severity of ADHD.

**Conclusions**

The first study demonstrated that the ability to modulate alpha oscillations in visual regions with the allocation of spatial attention was clearly present in healthy boys, but not in boys with ADHD. The second study demonstrated that male adolescent non-responders to methylphenidate display a lower frequency at which frontal alpha oscillations is peaking. The typical maturational changes in EEG emerging in adolescents observed in ADHD responders and controls, are absent in non-responders.

**Trial registration**

Clinical trial registration information; www.clinicaltrials.gov; NCT01932398 & NCT00863499

**Competing interests**

MA reports research grants, options/shares from Brain Resource Ltd. (Sydney, Australia) and neuroCare group and he is also a co-inventor on 4 patent applications (A61B5/0402; US2007/009323, A1; WO2001/013951 A1) related to EEG, neuromodulation and psychophysiology, but does not own these nor receives any proceeds related to these patents.
Brain arousal regulation: a predictive biomarker in psychiatry

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Arousal fundamentally impacts normal and abnormal behavior, and recently research on disturbed arousal regulation in mental disorders has attracted increasing interest. Accordingly, arousal has been implemented as a basic dimension of mental disorders in the Research Domain Criteria (RDoC) project of the National Institute of Mental Health (NIMH). The talk introduces the arousal regulation model of affective disorders and ADHD, which suggests hyper-arousal as a core pathogenetic factor in uni- and bipolar depression, and, in contrast, hypo-arousal in mania and ADHD. The model explains different clinical phenomena, as manic behavior is in parts interpreted as an autoregulatory attempt to stabilize brain arousal by creating a stimulating environment, whereas the withdrawal and sensation avoidance as well as insomnia symptoms in depression is seen as reflecting the underlying chronic hyperarousal. Many depressed patients experience themselves as subjectively fatigued and in need of rest, extended bed-times and inactivity in most cases do not result in the desired recovery but in the contrary depressive symptoms and sleep problems tend to increase. As inferable from the model, interventions that decrease arousal (e.g. antidepressants) or increase sleep need (e.g. sleep deprivation, sleep restriction) are efficient, whereas the inefficacy of stimulants has often been shown in depression. The arousal model contributes to delineating more homogenous subgroups within affective disorders and predicts response to treatment based on the respective brain arousal disturbance. Electroencephalography under resting conditions is most suitable for the assessment of brain arousal regulation, as different arousal states (also called EEG-vigilance stages) can be differentiated during the transition from high alertness to drowsiness until sleep onset according to specific EEG characteristics. The second part of the talk will introduce a computer-algorithm (Vigilance Algorithm Leipzig, VIGALL2.1), allowing semi-automatic classification of EEG-vigilance stages during resting-EEG recordings. The time sequence of these EEG-vigilance stages indicates the individual arousal regulation of the recorded subject. The final part of the talk will outline results from current studies applying VIGALL 2.1 and investigating hypotheses derived from the arousal regulation model with regards to the usage of brain arousal regulation as a diagnostic and/or predictive biomarker in psychiatric research.

Competing interests
The author(s) declare(s) that they have no competing interests.

Electrophysiological markers in the prediction of various treatment approaches in major depression and obsessive compulsive disorder

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A27

Despite large research efforts - also in electrophysiology - within the last decades, up to now there are no clinically accepted or used biomarkers that help to diagnose affective or anxiety disorders: It seems that the discriminative abilities of clinicians concerning diagnostic decisions are fair enough. However, what is even more important than a correct diagnosis is choosing the best treatment. Here, the decisions of the clinicians are not supported by any evidence. Especially in disorders where with various treatment approaches - such as major depressive disorder (MDD) and obsessive compulsive disorder (OCD) - but a still very high percentage of non-responders to first line therapies, biomarkers could contribute to an individualized medicine with faster responses and less trial and error approaches. Electrophysiological biomarkers provide a direct window to brain function combined with cost-effective settings and broad availability. Therefore, data on electroencephalogram (EEG) based algorithms derived from different studies will be presented that help to discriminate patients with better response to different types of treatment. Within the first part, the focus will be on predictors for treatment outcome following therapy with different types of antidepressants. Further, EEG-based biomarkers with discriminative power concerning outcome of electro-convulsive therapy (ECT) and will be shown. The second part is dedicated to treatment of OCD with cognitive-behavioral therapy, selective-serotonin inhibitors (SSRIs) or a combination of both. EEG-based biomarkers will be presented that might support the right choice of treatment.

The presentation of data will be followed by a brief outlook to what will has to come to transfer research knowledge into every day routine.
Besides the need for large, prospective multicenter studies, new analytical approaches will be presented that could help to establish an individualized medicine in neuropsychiatric disorders one day.

Competing interests
Authors report no competing interests.

A28
The sgACC in depression: getting at the heart of it
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A28

Major depressive disorder (MDD) is a chronic, mental disease with a remitting and relapsing course. Antidepressant medication is the most common treatment for MDD, however, the precise working mechanism underlying these treatments remains unclear. Recent neuromodulation treatments demonstrate that direct stimulation of the dorsolateral prefrontal cortex (DLPFC) and subgenual anterior cingulate (sgACC) relate to clinical improvement, suggesting connectivity alterations in this network to mediate antidepressant response, which might be similar for pharmacological treatments as well. This will be the focus of part 1. The international Study to Predict Optimized Treatment in Depression (iSPOT-D) is an international multicenter study that collected EEG data for 1008 MDD patients. We investigated whether connectivity in alpha and theta frequencies within this network changed during antidepressant treatment between: patients and controls, and responders and non-responders. Women exhibited higher alpha and theta connectivity compared to males, both pre- and post-treatment. Decreased alpha connectivity after treatment was found only for male responders, while non-responders and females exhibited no changes in alpha connectivity. Furthermore, it could be useful to a priori stratify by gender for future MDD studies [1].

Part two focusses on functional connectivity assumptions between the DLPFC, sgACC and the vagal nerve. Preliminary results will be presented regarding a method to localize the DLPFC: neuro-cardiac-guided rTMS (NCG-rTMS). The efficacy of rTMS in the treatment of MDD has been well established in recent years, however with various methods of locating the DLPFC. It has been proposed that the efficacy of rTMS in MDD is more related to stimulating the area that is functionally connected to the sgACC rather then to specific cortical areas. Therefore, we set-out to develop and test a new method that employs the functional role of the sgACC to establish in real time if the correct cortical area is targeted. Several studies have shown that the sgACC is involved in parasympathetic regulation such as heart rate (HR) and respiratory sinus arrhythmia (RSA), whereas HRV distinguishes healthy subjects from depressed patients. Our data suggest that prefrontal REM sleep-deprived cordance may predict response to antidepressant treatment in depressed patients, whereas HRV distinguishes healthy subjects from depressed patients.

Results
First, fourteen responders had significantly higher prefrontal theta cordance as compared to nineteen non-responders after the first week of antidepressant medication. Second, HRV in REM sleep was decreased in depressive patients at week four as compared to controls (high effect size; Cohen’s d > 1). Third, HRV showed negative correlation with REM density in healthy subjects and patients at week four.

Conclusions
Our data suggest that prefrontal REM sleep-deprived cordance may predict response to antidepressant treatment in depressed patients, whereas HRV distinguishes healthy subjects from depressed patients.

A30
Applying integrated EEG-behavioural analyses in genetic mouse models for Autism Spectrum Disorder; the identification of translational neuronal biomarkers
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A30

Autism Spectrum Disorder (ASD) is a highly heterogeneous neurodevelopmental disorder that is clinically defined by impaired social interaction, as well as repetitive and restricted behaviours. More than 200 ASD risk genes have been identified. Understanding the functional impact of these genetic variants will contribute to unravel clinical heterogeneity and to improve treatment efficacy. To provide clinically relevant neuronal biomarkers for specific biological subgroups, we initiated integrated EEG-behavioural analyses in selected genetic mouse models for ASD. So far, EEG data from ASD patients have shown changes in the power of brain oscillations in several frequency bands as well as in response to sensory stimuli. Although these results tell us that ASD patients have alterations at the level of brain circuits, and/or during processing of specific tasks, these biomarkers are currently not used for diagnosis of the disease.

To increase the selectively and sensitivity of neurophysiological biomarkers we analyzed the EEG of genetically modified mice during specific aspects of a behavioral task. To this end we coupled the systems used for EEG recordings and the software that tracks the location of the animal, allowing us to select parts of EEG measurements based on animal behaviour. We used Protocadherin 9 (Pcdh9) mutant mice as a model for the social interaction deficits seen in ASD patients (Bruining et al., 2015). These mice show deficits in sensorimotor development and long-term social discrimination capacity, while long-term fear conditioning is normal. Preliminary EEG results showed a decrease in gamma band oscillations in mutant mice during social interaction with both familiar and novel intruder mice. Furthermore, mutant mice showed impaired sensory information processing during an auditory mismatch negativity task. Currently, we are expanding these findings and investigate both EEG
Anesthesia, an opportunity to measure a pharmaco-EEG par excellence

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A31

The IPEG is an association for researchers involved in electrophysiological brain research and pharmacology, and the contribution of pharmaco-EEG research to the field of neuroscience is gaining in importance. In the past few years the functions of brain circuits, i.e. functional neuroanatomical resting-state networks, have come to be on the verge of being understood. This progress is the result of close collaboration between many disciplines: neuroanatomy, psychology, physics and pharmacology, to name just a few, which are making a joint effort to understand the functioning of the brain.

In view of these developments, the EEG measured during anesthesia might hold keys to disentangle (or to the contrary perhaps to unify), behavioral, pharmacological and neurophysiological signatures of various states of behavior, especially of the difficult to quantify states of consciousness. This is because anesthesia is a drug-induced state in which patients do not have any sensation, they are unconscious.

Moreover, during the whole period of anesthesia, the anesthesiologist meticulously monitors the state of wakefulness, so this procedure complies perfectly with the IPEG recommendation, which advises to measure EEG activity under vigilance-controlled conditions [4].

To induce a state of anesthesia, a variety of drugs can be used, all with quite different molecular targets. One of the still unanswered questions is: are different drugs inducing different states of anesthesia, or is anesthesia a well-described state that might be induced by modifying different stations in a hypothesized anesthesia circuit?

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In this oral I will mini-review the literature to point out characteristic EEG and connectivity changes induced by various types of anesthetics, propofol, isoflurane and ketamine included [e.g.1,3]. I will illustrate the findings in the literature with our own data of both rats and humans [2,3]. Further research questions will be proposed and discussed with the audience, in the hope to boost interest and research in our IPEG society in the EEG under anesthesia, the pharmaco-EEG par excellence.

Competing interests
None.

References

Alzheimer’s disease (AD) is a disconnection syndrome manifested by the disruption of white matter integrity, loss of synapses and functional connectivity (FC) across different cortical and subcortical regions. Early in AD progression, tau pathology can be found in the brainstem locus coeruleus (LC) prior to its amyloid-induced exacerbation and clinical symptoms. Accordingly, the pathological process of AD is characterized by the cell-cell spread of tau pathology from the LC into the medial temporal lobe, which triggers pathological changes causing functional disconnectivity. Network dynamics have become a leading model to assess both the anatomical relationships (structural networks) and the coupling of dynamic neurophysiology (functional networks) linking separate brain regions.

The present study used a tau seeding model in which preformed synthetic tau fibrils (K18) were unilaterally injected into the LC of transgenic mice expressing mutant human P301L tau, equipped with multichannel electrodes in frontal cortical and CA1–CA3 hippocampal areas. This approach allows us to 1) quantify longitudinal coherence and FC using phase-amplitude theta-gamma coupling (PAC); 2) identify the directionality of connectivity, using lagged and extended partial direct coherence (PDC); 3) measure pre-attentive auditory P50 potentials; 4) investigate sleep-wake organization; and 5) quantify phospho(p)-tau pathology in regions of interest using immunohistochemistry (AT100 antibody).

At the functional level, a decrease in spectral power at a range of frequencies in the hippocampal regions ipsilateral to the injection site is found at 2 weeks post-K18 injection, while an increase in power in contralateral hippocampal regions is hypothesized to be indicative of early compensatory mechanisms. Inter-hippocampal coherence is reduced in slow frequency oscillations and FC is significantly impaired as evidenced by: decreased intra- and inter-hemispheric hippocampal directionality of theta frequency oscillations; and reduced intra- and inter-hemispheric functional PAC strength. At the structural level, abnormal pTau aggregation is regionally specific, with AT100-positive tau detected in the pons, medulla, thalamus and cerebellum.

Ongoing assessment of pre-attentive auditory information processing, sleep-wake alterations and changes in the activity of GABAergic interneurons, which play a critical role in theta-gamma interactions, will allow further investigation into this aforementioned network dysfunction. Electrophysiological abnormalities in the hippocampus and cortex following injection of K18 into the LC convincingly support the relevance of tau pathology early in the LC. These functional alterations offer a reliable in vivo assay to test AD therapeutic agents for early intervention of tau pathology and possible prevention of the impairments in synaptic plasticity and neuronal network connectivity as seen in AD.
A33
Public-private initiative to align EEG biomarkers of Alzheimer’s disease in human and mouse models for early stages of drug discovery: the achievements of IMI PharmaCog project

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A33

Background and aim
In the European FP7 IMI “PharmaCog” project (Grant Agreement n° 115009, www.pharmacog.org), we evaluated whether cortical electroencephalographic (EEG) rhythms in quiet wakefulness reflected prodromal Alzheimer’s disease (AD) in amnestic mild cognitive impairment (aMCI) and had a back-translational value in transgenic mouse models of Alzheimer’s disease (AD).

Methods
The research data (including human biological samples) were sourced ethically and used in line with international ethical standards. EEG rhythms were recorded in 127 aMCI subjects. Cortical sources of EEG rhythms were estimated by eLORETA package (http://www.uzh.ch/keyinst/loreta.htm). Back translation of the EEG markers was tested on on-going EEG rhythms in wild type and transgenic mouse models of AD developing an accumulation of Aβ1-42 in the brain (i.e. one mutation in PDAPP and two mutations in TASTPM).

Results
(1) Compared with the aMCI sub-group showing “negativity” to Aβ1-42/phospho tau in the cerebrospinal fluid, the aMCI sub-group showing “positivity” (profound AD) exhibited an abnormal delta (<4Hz) source activity in widespread cortical regions while a posterior source activity in low-frequency alpha rhythms (8–10.5 Hz) pointed to a progressive abnormality across disease progression in 2 years; (2) On-going EEG rhythms in the same frequency range were abnormal in the transgenic PDAPP and TASTPM mice when compared to the control wild-type animals. Furthermore, these EEG rhythms were modulated by an Aβ1 – 42 lowering agent (monoclonal antibody 3D6) administered for 4 weeks in TASTPM mice. No effect was observed in wild-type mice.

Conclusions
The results of the PharmaCog project suggest that markers of on-going cortical EEG rhythms < 12 Hz may reflect profound AD processes in aMCI subjects and can be back-translated to transgenic mouse models of AD. These results encouraged the use of EEG biomarkers for an early evaluation of new AD modifying drugs in transgenic mouse models of AD.

A34
Neurophysiological biomarkers in first episode psychosis

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A34

A biomarker is an objective biological measure that can be used for health risk prediction, or for screening, diagnosis, or tracking disease progression. In psychosis research, a key recent development has been the application of a staging framework. The understanding that there is a progressive course to psychosis, including a potential prodromal phase of increasing disability before the emergence of frank psychosis, has spurred the search for screening biomarkers to indicate those truly prodromal clinical high risk for psychosis individuals that will convert to psychosis (approximately 30 % in 3 years), crucial for early prophylactic treatment. Practically, true prodromal cases are rare and studies tracking conversion long and costly. To be useful for screening any disease presence biomarker must obligatorily be reduced at first psychosis. Our group has been testing various auditory-based neurophysiological tasks in first episode psychosis individuals to develop screening biomarkers for an incipient psychotic break. Here we describe several candidate biomarkers. Passive listening, simple mismatch negativity (MMN) to a rare deviant tone is not abnormal in first episode psychosis (Study 1: 29 first episode schizophrenia and 40 controls; Study 2: 35 first episode schizophrenia and 35 controls). Complex MMN tasks that depend on extraction of patterns in auditory sequences show more promise. In a series of passive tasks where the number of tones in groups were occasionally changed, we saw significant reductions of complex MMN to a rare extra tone (19 first episode schizophrenia and 19 controls). We also observed abnormalities of a slow potential that appeared to indicate the formation of each group as an acoustic object. Having participants actively count stimuli in each group revealed that a missing tone (e.g. a group of 3 instead of a group of 4) elicited an emitted P300, and that the posterior P300b component was abnormal in first episode schizophrenia (20 first episode schizophrenia and 32 controls). Using a single tone auditory evoked potential task, attention to stimuli was manipulated by having participants either press a button to every 7 tones or watch a silent video (10 first episode schizophrenia and 10 matched controls). Healthy participants modulated their N100 with attention, but first episode psychosis individuals did not. These data suggest that several neurophysiological measures may be suitable as biomarkers for the presence of psychosis. Future work will deploy these tasks in clinical high risk individuals to track whether they show promise as screening biomarkers for an incipient psychotic break.

A35
Perception of sleep in patients with insomnia related to generalized anxiety disorder, patients with apnea and in healthy controls

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A35

Background
Although sleep research has been trying to elucidate the relations between objective and subjective sleep and awakening quality for five decades, findings have often been controversial. This might be due to the lack of large data sets, to the differences between different sleep disorders and to the inter-individual differences in sleep perception per se.

Objectives
The aim of the present study was to investigate relations between objective and subjective sleep variables in a large number of healthy subjects as well as in 2 clinically relevant patient groups, i.e. nonorganic insomnia in generalized anxiety disorder (insomnia/GAD) and sleep apnea.
Material and methods
One hundred and seventy-seven healthy subjects (94 females, 83 males, aged 20 – 95 years), 61 insomnia GAD patients (32 females, 29 males, aged 21 – 66 years) and 51 apnea patients (7 females, 44 males, aged 29 – 73 years) underwent two polysomnographic nights analyzed by the Somnolyzer [1] and completed the self-rating scale for sleep and awakening quality [2].

Results
Patients with insomnia/GAD underestimated their sleep efficiency in both nights (objective sleep efficiency index (SEI) 77 % and 84 % versus subjective SEI 57 % and 64 % for adaptation and baseline night, respectively, p < 0.001 Wilcoxon test). Apnea patients showed no differences between subjective and objective SEI (objective 80 % and 87 % versus subjective 79 % and 86 %). Healthy controls – specifically males and subjects older than 60 years – overestimated their sleep efficiency in the adaptation night (objective 80 % and 86 % versus subjective 84 % and 87 %, p < 0.001 for night 1). Correlation analysis between objective and subjective SEI on change values from adaptation to baseline night revealed highly significant correlations for all three groups (r = .77 for GAD, r = .57 for apnea and r = .51 in healthy controls). Interestingly, the regression lines go through the origin in all three groups, i.e. no change in objective SEI is perceived as no change in subjective SEI.

Conclusions
Relations between subjective and objective sleep efficiency are influenced by age, gender and the type of sleep disorder. In correlation analyses, the problem of inter-individual judgements of sleep perception can be reduced by using change values between adaptation and baseline nights rather than raw values. The variety of correlations requires a parallel evaluation of subjective and objective variables as they are not interchangeable.

Competing interests
Peter Anderer, Georg Gruber, Silvia Parapatics and Georg Dörfner are employees of The Siesta Group Schlafanalyse GmbH. Gerda M Saletu-Zyhlarz and Bernd Saletu have no competing interests.

References

A36
Catecholaminergic regulation of learning rate in a dynamic environment
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A36

Adaptive behavior in a changing world requires flexibly adapting one’s rate of learning to the rate of environmental change. Recent studies have examined the computational mechanisms by which various environmental factors determine the impact of new outcomes on existing beliefs (i.e., the “learning rate”). However, the brain mechanisms, and in particular the neuropeptudors, involved in this process are still largely unknown. The brain-wide neurophysiological effects of the catecholamines norepinephrine and dopamine on stimulus-evoked cortical responses suggest that the catecholamine systems are well positioned to regulate learning about environmental change, but more direct evidence for a role of this system is scant. Here, we report evidence from a study employing pharmacology, scalp electrophysiology and computational modeling (N = 32) that suggests an important role for catecholamines in learning-rate regulation. We found that the P3 component of the EEG—an electrophysiological index of outcome-evoked phasic catecholamine release in the cortex-predicted learning rate, and formally mediated the effect of prediction-error magnitude on learning rate. P3 amplitude also mediated the effects of two computational variables—capturing the unexpectedness of an outcome and the uncertainty of a preexisting belief on learning rate. Furthermore, a pharmacological manipulation of catecholamine activity affected learning rate following unanticipated task changes, in a way that depended on participants’ baseline learning rate. Our findings provide converging evidence for a causal role of the human catecholamine systems in learning-rate regulation as a function of environmental change.

A37
Dopamine and the cortical representation of reward
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A37

The way humans behave is greatly affected by the principle expected utility, the combination of subjective value (SV) of the outcome of an act (is it rewarding?) and subjective probability (SP) of that outcome. In an initial study (n = 42) we examined the electro-cortical representations of the anticipation of SV and SP during a cued Go/NoGo experiment. During this task cue letters signaled upcoming target letters to which participants had to respond. The probability of target letter appearance after the cue letter and the amount of money that could be won for correct and fast responses were orthogonally manipulated across four task blocks. Results show that reward availability affected a prefrontal reward P200 and a centro-parietal P300 ERP. Moreover, a fronto-central ERP was affected by both reward and probability manipulations. These results suggest that reward and probability are partially separately processed in the cortex. Furthermore, reward and probability information are integrated around 300 ms after presentation of the cue and possibly processed via a shared underlying cortical mechanism that may act to reduce uncertainty or to prepare for action. In a follow-up study we investigated the role of dopamine (and noradrenaline) in either of these processes by employing a within-subjects haloperidol/clonidine/placebo cross-over design (n = 24) with the same cued Go/NoGo paradigm.

Trial registration
The Netherlands National Trial Register (NTR) (CC = 4493).

A38
Optimizing the earliest memory stages: a role for acetylcholine and serotonin?
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A38

Acetylcholine and serotonin both play an important role in encoding and consolidation of memories. However, it has also been suggested that these two neuromodulators take part, and might even interact, in processes initiated before conscious encoding even takes place. Sensory memory and novelty detection, two processes related to reduction of surprise, are part of those early stages. They can be measured with the mismatch negativity (MMN) and P3a components of the event-related potential, respectively. In a series of experiments, we examined whether cholinergic and serotonergic manipulations affect MMN and P3a components during a novelty oddball task. In this task, frequent standard stimuli were interspersed with infrequent deviant and infrequent novel stimuli at a pace of one stimulus presentation per second. Biperiden, a cholinergic agonist, and rivastigmine, a cholinesterase inhibitor, did not affect MMN amplitude. Acute tryptophan depletion, a method to reduce serotonin in the brain, and citalopram, a selective serotonin re-uptake inhibitor, were
also unable to affect the MMN. No significant interactions between treatments were found related to the MMN. Cholinergic treatments did, however, affect the P3a amplitude: P3a was decreased after Biperiden intake and increased after Rivastigmine. The serotonergic manipulations did not affect P3a amplitude, neither were interactions found between treatments. Our results thus show that, although the cholinergic and serotonergic systems do not seem to play a role in sensory memory, acetylcholine’s role in novelty detection, and thus in handling surprise, is evident.

A39
Processing of the mismatch negativity under the LSD state
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A39
Background
Lysergic acid diethylamide (LSD) is a classic psychedelic drug and serotonin 2A receptor agonist. A common feature of the LSD state is its capability to provide an experience of modulating the salience of external events. The Mismatch Negativity (MMN) is an event-related potential/field (ERP/ERF), that indexes expectation disruption (or ‘surprise’) mechanisms, which have been shown to be modulated in patients with disorders of consciousness, as well as in schizophrenia and following ketamine administration. In this study the MMN paradigm was used to assess expectation and surprise mechanisms under LSD and placebo conditions in healthy participants.

Methods
A balanced order, within subject design was used for the study. 20 Healthy volunteers underwent MEG recordings following intravenous administration of LSD (75 mcg IV) and placebo at least 2 weeks apart. Participants were presented with auditory stimuli consisting of oddball and standard tones while resting inside the MEG scanner. Following preprocessing and averaging, the resulting event-related fields (ERF) were converted into scalp-map images and smoothed for statistical analysis corresponding to four conditions: auditory stimuli of standard tones under LSD (1) and placebo (2) and deviant tones under LSD (3) and placebo (4). The ERFs were entered into a within-subject analysis of variance with 2 main factors: ‘drug’ (LSD and placebo) and ‘expectation disruption’ (standard and deviant).

Results
An interaction effect between ‘expectation disruption’ and ‘drug’ factors was found in a right lateralized cluster in the scalp. Post hoc analyses within this ROI, reveal significant differences in the processing of standard tones between placebo and LSD conditions as well as deviant tones. Within the placebo condition the difference between standards and deviants was significant, while it wasn’t following LSD administration.

Conclusions
Results indicate a reduction of activity related to the processing of novel stimuli, while showing that the surprise response was increased under the LSD condition in large areas of the scalp for familiar stimuli. These findings may inform how salience mechanisms may be disrupted under LSD and is consistent with reports of “increased novelty” to familiar stimuli in the LSD state. Mechanisms underlying this modulation may be accounted by modulation of prediction error in the psychedelic state.

This research received financial and intellectual support from the Beckley Foundation and was conducted as part of a wider Beckley-Imperial research programme.

A40
The effects of psilocybin on human EEG, comparison with animal models
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A40
Objectives
Psilocin, the active metabolite of psilocybin, is a classical psychedelic tryptamine acting as an agonist at serotonin 5-HT2A/C and 5-HT1A receptors. It has been used extensively to model psychosis in humans as well as in animals, and during the last decade given more attention as a potential antidepressant and anxiolytic drug. Recent studies in healthy volunteers have shown that psilocybin leads to a global desynchronization of brain activity and disconnection of the main brain networks. The purpose of the current study is to compare the translational effects of psilocybin/psilocin on EEG activity and connectivity between healthy volunteers and rats.

Methods
For the human study, a 19 channel EEG of a standard 10/20 system was recorded with BioS-DAQ9 amplifier (M8). Subjects (M10/F10) were administered 0.26 mg/kg of psilocybin or placebo orally in a double-blinded, crossover manner. EEG was recorded before and at 50, 90, 180 and 240 min after drug administration. In the animal study, psilocin was administered to male Wistar rats (n = 10) subcutaneously in dose of 4 mg/kg. Multichannel EEG with 12 cortical (6 homolateral pairs) electrodes at frontal, parietal and temporal regions were recorded for 10 min before and 90 min after dosing using BrainScope EADS 221 amplifier (M8). EEG in animals was co-recorded with behavioural activity and epochs of inactivity were then subjected for further processing. Data was pre-processed using Brainvision and WaveFinder software followed by further power spectral and coherence analyses using Neuroguide software. Source localization of EEG activity was analysed by LORETA, 3D brain mapping in animals was performed using an in-house developed Matlab tool.

Results
Psilocybin induced an absolute as well as relative alpha power decrease in occipital regions, while beta and gamma power increases in fronto-temporal areas in humans. The effects were most robust 50 min after drug administration. Source localization by LORETA confirmed the localization of EEG changes. However, it is of note that the gamma power cannot be distinguished from artificial motor activations. EEG coherence was mainly decreased in theta, alpha and beta bands, with some increases observed in beta and gamma bands. In rats there was a global power decrease in absolute power however relative power showed partly similar profile to humans with theta power decrease and beta and gamma power increase. Furthermore, there was a global decrease of coherence in rats following drug administration.

Conclusions
The effects of psilocin/psilocybin resulted in similar direction of EEG changes in both humans and rats with disconnection (decreased coherence) being the most stable phenomenon observed, indicating good translational validity.

This study was supported by the projects IGA MHCR NT/13897, ED2.1.00/03.0078, LO1611/NPU I, MH CZ - DR (NIMH-CZ, 00023752) and PRVOUK P34.

A41
Salvinorin-A induces a unique pattern of neurophysiological effects in humans characterized by alpha suppression and widespread increases in cortical delta activity
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A41
Objectives
Salvinorin-A induces a unique pattern of neurophysiological effects in humans characterized by alpha suppression and widespread increases in cortical delta activity.

Methods
Psilocybin was administered to male Wistar rats (n = 10) subcutaneously in dose of 4 mg/kg. Multichannel EEG with 12 cortical (6 homolateral pairs) electrodes at frontal, parietal and temporal regions were recorded for 10 min before and 90 min after dosing using BrainScope EADS 221 amplifier (M8). EEG in animals was co-recorded with behavioural activity and epochs of inactivity were then subjected for further processing. Data was pre-processed using Brainvision and WaveFinder software followed by further power spectral and coherence analyses using Neuroguide software. Source localization of EEG activity was analysed by LORETA, 3D brain mapping in animals was performed using an in-house developed Matlab tool.

Results
Psilocybin induced an absolute as well as relative alpha power decrease in occipital regions, while beta and gamma power increases in fronto-temporal areas in humans. The effects were most robust 50 min after drug administration. Source localization by LORETA confirmed the localization of EEG changes. However, it is of note that the gamma power cannot be distinguished from artificial motor activations. EEG coherence was mainly decreased in theta, alpha and beta bands, with some increases observed in beta and gamma bands. In rats there was a global power decrease in absolute power however relative power showed partly similar profile to humans with theta power decrease and beta and gamma power increase. Furthermore, there was a global decrease of coherence in rats following drug administration.

Conclusions
The effects of psilocin/psilocybin resulted in similar direction of EEG changes in both humans and rats with disconnection (decreased coherence) being the most stable phenomenon observed, indicating good translational validity.

This study was supported by the projects IGA MHCR NT/13897, ED2.1.00/03.0078, LO1611/NPU I, MH CZ - DR (NIMH-CZ, 00023752) and PRVOUK P34.
Background
Salvinorin-A (SA) is a potent perception-modifying drug found in the leaves of the plant Salvia divinorum. Unlike 5-HT2A agonists such as DMT, LSD and psilocybin, SA is a selective kappa-opioid receptor (KOR) agonist. Its pattern of effects in humans also shows important differences with that of the classical psychedelics. While subjects usually experience intense visual and auditory phenomena, SA completely blocks external sensory perception, and leads to a characteristic total loss of contact with external reality. Here we investigated the neurophysiological correlates of SA effects in humans.

Methods
We measured spontaneous brain oscillations (EEG) in 24 healthy volunteers, before and after the administration of 1 mg vaporized SA. We recorded the EEG from 19 scalp leads and we calculated drug-induced energy changes in ten frequency bands between 1.3 and 40 Hz. Additionally, we computed the changes in the intracerebral current density distribution associated with the voltage values recorded at the scalp.

Results
SA administration led to rapid and significant changes in brain oscillations that coincided with maximum drug levels in plasma. SA suppressed the alpha rhythm (7.5-13 Hz) and markedly increased slow delta activity (1.3-3.5 Hz). Less prominent effects included increases in the theta (3.5-7.5 Hz) and low gamma (35-40 Hz) bands. Alpha decreases were localized over parieto-occipital regions, including the posterior cingulate cortex and visual areas. Delta increases were observed over most of the brain, with the maximum located over auditory and visual cortex in the left temporal lobe. Theta increases were found over left temporal and frontal areas. Finally, gamma increases were restricted to visual areas in the occipital cortex.

Conclusion
These results show a unique pattern of neurophysiological effects for SA in humans. While it shares with serotonergic psychedelics the alpha-suppressing action, its main neurophysiological signature is an atypical enhancement of slow delta activity. These differences may explain the marked differences in subjective effects between SA and 5-HT2A agonists.

This research was supported by Grant P112/02758 from the Spanish government.

Results
Ketamine induced immediate (10 min and 30 min) decrease of parieto-occipital sources of alpha-1 and alpha-2 activities and an increase of gamma-sources in all subjects. Responders to medication were characterized by excess of mediofrontal delta and theta sources in comparison to non-responders. Moreover, only the responders showed significant changes that persisted 24 h after infusion, while no significant changes were observed in non-responders. Among the clinical variables we have found a significant correlation between the BPRS score during ketamine infusion and MADRS score at day 7, and the intensity of psychotomimetic symptoms during infusion seems to be the strongest clinical predictor of antidepressive effect of ketamine. Regarding the QEEG parameters, the patients with better responses showed higher pre-treatment theta activity in mediofrontal areas and in the rostral anterior cingulate. Better response to ketamine was also connected with higher pre-treatment lagged phase synchronization (i.e. higher connectivity) between anterior cingulate and mediofrontal cortex at theta and alpha-1 frequency bands.

Our results suggest that an acute increase of mediofrontal cortical sources of theta and delta activities after ketamine infusion could be potential biomarkers to differentiate responders and non-responders to ketamine. Higher pre-treatment theta activity in mediofrontal areas together with higher lagged phase synchronization between anterior cingulate and mediofrontal cortex at theta and alpha-1 frequency bands could serve as predictors of treatment response to ketamine. Moreover, the antidepressive effect of ketamine seems to be undoubtedly connected with patient’s psychotomimetic experience. Supported by grants AZV MZCR 15-33250A and by the project PRVOUK P34.

Oral presentations

A42

QEEG signatures predicting antidepressant response to ketamine
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A42

Objective
Treatment-resistant depression (TRD) is a disabling disorder that negatively impact patient’s morbidity and mortality and constitutes a major challenge for current psychopharmacology. The discovery of rapid-acting antidepressive action of ketamine has motivated studies aiming to reveal the molecular mechanism of this effect and to enable the clinical application of similarly rapid-acting antidepressants. In our two studies, the time-course of effects of ketamine was assessed in treatment-resistant depressive patients by QEEG to elucidate changes associated with treatment effect and to assess potential predictors of treatment response.

Methods
The pool analysis was completed from data of two double-blind, cross-over, placebo-controlled studies, assessing the effect of single infusion of ketamine (0.54 mg/kg within 30 min) in altogether 50 inpatients with major depressive disorder. EEG data were analysed during the infusion (10 min and 30 min) and 24 h after ketamine administration using exact low-resolution electromagnetic tomography (eLORETA). Response to treatment was defined as a ≥50 % reduction of MADRS score.

Results
In our two studies, the time-course of effects of ketamine was assessed in treatment-resistant depressive patients by QEEG to elucidate changes associated with treatment effect and to assess potential predictors of treatment response. The pool analysis was completed from data of two double-blind, cross-over, placebo-controlled studies, assessing the effect of single infusion of ketamine (0.54 mg/kg within 30 min) in altogether 50 inpatients with major depressive disorder. EEG data were analysed during the infusion (10 min and 30 min) and 24 h after ketamine administration using exact low-resolution electromagnetic tomography (eLORETA). Response to treatment was defined as a ≥50 % reduction of MADRS score.

Oral presentations

A43

Nonlinearity of the visual system assessed by cross-frequency phase coupling
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A43

Background
Processing of visual input by the brain is a highly nonlinear operation, involving complex interactions among neuronal networks. Nonlinear visual system activity includes harmonic interactions, thought to reflect resonance of neural processing, whereas intermodulation, being the contribution of multiple input frequencies to one output frequency, relates to functional integration. Using a sum-of-sinusoid signal as visual input [2], it is possible to elicit a richer class of nonlinear responses than the classic pulse train stimulus, thereby providing a more complete description of nonlinearity. Here, we will use nonlinear EEG analyses to quantify higher-order nonlinearities in visual processing.

Methods
Ten healthy participants were subjected to bi-sinusoidal light stimulation of 13 and 23 Hz for 320 1 s-epochs, while scalp EEG (8 electrodes) was recorded at the occipital, parietal and frontal lobes. The frequencies of light stimulus were chosen to guarantee no overlap of their harmonic and intermodulation frequencies for different orders of nonlinearity. Nonlinear interactions and time delay from stimulus to cortex were analyzed in the frequency domain using novel phase synchronization measures [3] and amplitude spectrum.

Results
Higher harmonic and intermodulation interactions were detected between visual input and cortical responses. First to fourth order
phase coupling interactions were enhanced in the visual cortex com-pared to parietal and frontal responses. Spectral amplitude differ-ences were less pronounced between cortical regions. Time delay estimation showed a delay between light stimulus and visual cortex of 116 ± 21 ms, significantly higher than the delay between stimulus and frontal or parietal lobes.

Discussion
This study demonstrates the potential of using sum-of-sinusoid light stimulation and quantitative nonlinear EEG analysis to identify higher-order nonlinear dynamics of visual processing. We foresee that application of the described frequency interaction analyses can further our insight in the nonlinear dynamics of visual processing not only in healthy subjects, but also with respect to the pathophysi-ology of neurological diseases with visual manifestations that relate to cortical hyperexcitability, like migraine and epilepsy.

References

A44
Whole-brain time-frequency analysis of event-related potentials for the assessment of pharmacodynamic effects in the human brain
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A44

We are developing Whole Brain Time-Frequency (WBT) analysis as a new physiological biomarker for clinical trials of pharmacodynamics of novel drugs. WBT analysis expands the power of event-related potential (ERP) assessment by using wavelets to measure both evoked (phase-locked) and induced (non-phase-locked) activity. Unlike traditional ERP measures, which are indexed by specific electrodes and peak latencies, WBT analysis measures integrated change in brain responses across time-frequency and space to infer whether a drug has a significant effect. WBT analysis also uses permutation tests and multiple comparison corrections to identify important within-subject changes between conditions and rule out differences arising from recording noise, artifacts or random variability.

The specific aim of this study was to assess the sensitivity and specificity of WBT analysis to drug effects that are typically measured with ERP amplitudes and latencies. We simulated effects of dose-related changes in N1-P2-P3 ERP components and 40-Hz induced gamma bursts at 24 electrodes. Simulations included a range of amplitude effects, latency effects and signal-to-noise ratios, serving to define the sensitivity and specificity of WBT analysis to ERP differences. The simulations allowed us to optimize parameters for WBT analysis, including choice of analyzing wavelets, energy normalization, baseline correction, measures of evoked and induced activity, and method of testing significant differences. We found that WBT analysis reliably detects small differences in evoked activity (on the order of 10%) in realistic noise and background EEG conditions. We found similar detectability of small differences in induced 40-Hz gamma bursts. It is the goal of the further studies to investigate the clinical relevance of these observed differences using WBT analysis, and to relate the evoked and induced components ERP differences to mechanisms of drug action. Currently we are applying WBT analysis to data from three Phase 1 clinical trials of novel compounds for schizophrenia in both healthy controls and schizophrenia patients.

A45
Disregulation of hyperpolarization-activated inward current current (Ih) affects thalamocortical oscillations: the role of the auxiliary subunit TRIP8b on HCN channel function in thalamic and cortical neurons
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A45

Background
The family of hyperpolarization-activated cyclic nucleotide-gated cation (HCN) channels consisting four different isoforms (HCN1-4) have a major role in controlling neuronal excitability and generation of rhythmic oscillatory activity in individual neurons and neuronal networks [1, 2]. These channels activate in response to hyperpolarizing potentials negative to −50 to −60 mV and depolarize the resting membrane po-tential. HCN channels are regulated by small molecules like cyclic nucleo-tides and different accessory proteins. TRIP8b is a brain-specific accessory subunit of HCN channels which controls the gating, surface expression and trafficking of different HCN channels subunits in many regions of brain [3–5]. The role of this protein for Ih characteristics in thalamic and cortical neurons and the functional consequences of TRIP8b dysregulation for thalamocortical oscillations however is not yet fully understood. The present study aimed at providing a better understand-ing of the functional role of TRIP8b in the thalamocortical system and shedding some light on possible dysfunctional aspects by combin-ing in vitro and in vivo electrophysiological approaches.

In this study, Ih was measured in whole cell patch clamp recordings from thalamocortical (TC) neurons of different thalamic nuclei, as well as pyramidal neurons in layer V and VI of the somatosensory cortex of TRIP8b-deficient (TRIP8b−/−) and control (C57Bl/6 J) mice (p15 – p90). Effects of TRIP8b-dependent dysregulation of Ih on thalamocor-tical oscillations was monitored by local field potential (LFP) record-ings from the ventral-posterior medial complex of the thalamus (VPM) and somatosensory cortex (p 90 – p120), regions which are known to be involved in generation of normal and also pathological thalamocortical oscillations.

Results
Characterization of Ih in the thalamocortical system in the absence of the auxiliary subunit TRIP8b showed a significant decrease in ih density and changes in intrinsic properties and firing patterns of TC and cortical pyramidal neurons. These changes were accompanied by an increase in cAMP sensitivity in TC neurons. Dysregulation of Ih in the thalamocortical system of TRIP8b−/− mice was associated with altered thalamocortical oscillations revealing a significant increase in slow osci-lillations in the delta frequency range (0.5–4 Hz) during episodes of active-wakefulness.

Conclusion
The results of our study point to the importance of TRIP8b, as a brain-specific auxiliary subunit of HCN channels, in regulation of cell and network oscillations. It was demonstrated here that the presence of TRIP8b is necessary for modulation of thalamocortical delta oscilla-tions due to its direct effect on HCN channels protein expression in the thalamocortical system.

References
Cognitive processes are based on the coordination of interactions of populations of neurons that are distributed within and across different specialized brain areas. Accumulating evidence suggests that neuronal oscillations play a pivotal role in driving brain communication. This communication is affected in many neurodegenerative diseases. Accordingly, understanding network interactions during cognitive activity is crucial for a better comprehension of neurodegenerative consequences for cognitive functioning as well as for assessing the efficacy of novel pharmacological treatments. The purpose of the present study was to evaluate putative EEG-based biomarkers during the trial-unique delayed nonmatching-to-location (TUNL) task in rats. The task assesses memory for location across different delays and spatial separations in a computer-automated touchscreen set-up. Once EEG-instrumented rats reached performance criteria (80% accuracy in an 8 s delay for two consecutive days), brain activity in the CA1 region of the hippocampus, medial prefrontal cortex, and retrosplenial cortex was monitored during two consecutive TUNL sessions using an 8 s and a 16 s delay. Time-frequency-based analysis of EEG readouts was used to investigate neuronal connectivity during the different delays comparing correct vs. incorrect trials. In particular, cross-frequency coupling (CFC, when the phase of a low frequency oscillation drives the amplitude of the coupled higher frequency oscillation) was analyzed as this has been suggested a possible mechanism facilitating working memory. It was hypothesized that functional connectivity during the delay will be reduced during incorrect trials compared to correct ones for both delays and separations. Behavioral results confirmed that accuracy was higher in the larger separation for the 8 s delay (M = 64.29%, 95% CI [81.03, 87.55]) compared with smaller separation for the same delay (M = 74.99%, 95% CI [69.89, 80.09]). Furthermore, the 16 s appeared to be more challenging as accuracy was reduced for both delays (M = 63.05%, 95% CI [55.95, 70.15]) and small separation (M = 60.41%, 95% CI [55.57, 65.25]) compared with the 8 s delay. Interestingly, results from the CFC during the delay support our hypothesis as 8 Hz frequency modulation of 90 Hz amplitude in the medial prefrontal cortex showed a more rapid decrease in CFC during the delay for the incorrect trials in both delays and separations compared to the correct trials. Overall, results identified the critical role of neuronal oscillations and connectivity for working memory in the TUNL task. This study reinforces the strength of combining multiple approaches to further understand cognitive processes and assessment of pharmacological treatments.

A47

EEG functional connectivity of Brodmann area 24 in obsessive-compulsive disorder
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A47

Background
A growing body of evidence have challenged the traditional orbitofrontostraial hypothesis and suggested that dysregulation of widespread brain networks may underlie the OCD disorder. In last decades, the increasing attention of neuroscience research has been paid to large-scale network organization of the brain. Three prominent networks were identified: “Central Executive Network”, “Default Mode Network” and “Salience Network”, responsible for synchronization of anticorrelated activity of DMN and CEN. Several studies have stressed the role of the dorsal anterior cingulate cortex, a core structure within the salience network, in OCD pathophysiology. Our two previous studies also revealed abnormal EEG activity in this structure (Brodmann area, BA 24), however little is known about its EEG functional connectivity in OCD. Based on our previous findings, we tested functional connectivity between EEG sources in BA 24 and rest of the brain in the group of OCD patients and in healthy controls.

Methods
96 in-patients diagnosed with OCD and 95 healthy controls matched for age and sex were included in the study. All subjects were right-handed. 27 OCD patients were drug-free and 69 were medicated with SSRIs. All subjects underwent 19-channel resting-state EEG examination. Functional connectivity was analysed in LORETA-KEY software. We assessed connectivity between centroid of BA 24 and centroids of all the other Brodmann areas as defined in the LORETA-KEY software. Lagged nonlinear connectivity was computed in eight frequency bands: delta (1.5 - 6 Hz), theta (6.5 - 8 Hz), alpha 1 (8.5 - 10 Hz), alpha 2 (10.5 - 12 Hz), beta 1 (12.5 - 18 Hz), beta 2 (18.5 - 21 Hz), beta 3 (21.5 - 30 Hz) and gamma (30.5 - 44.0 Hz). Groups were compared using t-statistics and permutation testing to correct for multiple comparisons.

Results
Drug-free and SSRIs medicated patients did not differ from each other in functional connectivity and therefore they were further tested as a unitary group. Compared with controls, OCD patients had higher lagged nonlinear functional connectivity between BA 24 and BA 5 and BA 7 in the beta 3 as well as in the gamma frequency band (p < 0.05). In the gamma band the results were significant only for the left BA 5 and 7, however connectivities in the right hemisphere were close to threshold.

Conclusion
We hypothesize that an aberrant synchronization between default mode and central executive network related to the aberrant activity of the salience network may underlie symptoms of OCD. This work was supported by the projects ED2.100/03.0078, LO1611 and MH CZ - DRO 00023752.

A48

Cortical network reorganization in mild and prodromal Alzheimer disease: graph theory approach on resting state EEG recordings
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A48

Cortical network reorganization in mild and prodromal Alzheimer disease: graph theory approach on resting state EEG recordings
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A48
Background
Alzheimer’s disease (AD) has a long preclinical period in the absence of overt symptoms in which the process progresses until it crosses a threshold to clinically recognizable dysfunction [1]. Graph theoretical analysis of cerebral networks has been implemented in AD challenging the classical concept of neurological disorders being either ‘local’ or ‘global’, and have pointed to the overload and failure of hubs as a possible final common pathway in neurological disorders. Previous EEG studies on the comparison between AD and control subjects reported divergent results [2]. An intermediate trend was found in subjects with mild cognitive impairment (MCI) with respect to AD and control subjects [3]. The aim of the study was to assess by means of graph theory analysis if AD patients with mild dementia could show a different cortical organization from age matched control subjects and if these possible differences could be already present at the stage of MCI.

Results
The main finding of the present study is that network reorganization is evident in AD since the prodromal stage (AD-MCI). Specifically, AD-MCI and AD showed a lower number of links among nodes than control group (p = 0.0007). Both within and outward links among nodes and brain areas with a high level of functional connectivity (so-called hubs) were found to be reduced in both AD-MCI and AD patients. Hubs in the parietal areas (P3, P4, and Pz) showed lower number of links in AD-MCI and AD than control group. Temporal nodes showed lower lower clustering coefficient and local efficiency in patients than control group. Significant differences between AD-MCI and AD were found in the right occipital node. Indeed, the clustering coefficient and the local efficiency was reduced in AD compared with AD-MCI in O2 (p < 0.05).

Conclusions
Our results suggest that brain network functional alterations mainly involved the temporal nodes in prodromal stage of AD, whereas brain dynamic changed in the posterior areas with disease progression to overt AD dementia.

The functional disconnection between temporal and parieto-occipital areas could be related to medial temporal lobe atrophy which is a characteristic neuropathological change in the early stage of AD [4].

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A49 Sleep disturbances in obsessive-compulsive disorder: association with response to repetitive transcranial magnetic stimulation (rTMS)
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A49

A50 Glutamatergic deficit and negative symptoms: new evidence from the ketamine model of schizophrenia
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Targeting the N-methyl-D-aspartate-receptor (NMDAR) is a major approach for treating negative symptoms of schizophrenia. The ketamine model of schizophrenia has the advantage of comprehensively producing schizophrenia-like symptoms such as positive, cognitive and negative symptoms in healthy volunteers. The amplitude of the Mismatch Negativity (MMN), a neurophysiological parameter related to infrequent stimuli, is known to be significantly reduced in schizophrenic patients but also in healthy controls receiving ketamine [1,2]. Accordingly, it was the aim of the present study to investigate whether changes of MMN during ketamine administration are related to the emergence of negative symptoms in healthy subjects. Therefore, we examined the impact of ketamine on MMN amplitudes and its sources (sources localization approach: low resolution electromagnetic tomography (LORETA)) by means of 64-channel electroencephalography (EEG) recording during performance of an auditory MMN paradigm and assessed the psychopathological status using the Altered State of Consciousness (SD-ASC) Rating Scale and the Positive and Negative Syndrome Scale (PANSS). Twenty-four male, healthy volunteers were measured with pharmacological EEG using a single-blind, randomized, placebo-controlled crossover design. We identified significant changes of the MMN response, to both duration and frequency deviants, under ketamine condition as well as a significant increase in all PANSS scores. Reductions of MMN amplitudes were significantly correlated with more pronounced negative symptoms, assessed by the PANSS. Accordingly, the MMN might represent a biomarker for negative symptoms in schizophrenia related to an insufficient NMDAR system and could be used to identify schizophrenia patients with negative symptoms due to NMDAR dysfunction and thus to determine a maximal benefit of drugs modulating neurotransmission at the NMDAR.

Competing interests
The authors declare that they have no competing interests.

References

A51 Isolated epileptiform discharges in psychiatry: outcomes in an integrative practice
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A51

Background
The search for biomarkers that can inform medication decisions in neuropsychiatric disorders is a goal of the Research Domain Criteria project under the National Institute of Mental Health. Isolated epileptiform discharges (IEDs) may be such a biomarker. IEDs have been linked to increased psychopathology that traverses many diagnoses [1]. It has been suggested that IEDs may represent an epiphenomenon with an etiology of unappreciated significance [2]. The literature suggests that anticonvulsants should be considered when IEDs are identified [3, 4]; however, outcome studies have yet to be published. This study investigates the predictive value of IEDs as a biomarker for the use of anticonvulsants on a large cohort of patients.

Method
We reviewed refractory cases from a large multidisciplinary practice whose EEG readings contained IEDs and were subsequently medicated with anticonvulsants by the clinic’s psychiatrist. The psychiatrists followed up progress notes were assessed to determine the impact of adding anticonvulsants. Ratings were based on clinical presentation and reported in three categories: Improved, unchanged, and more severe. There were two exclusion criteria: a prior diagnosis of seizure disorder and a history of prior treatment with anticonvulsants. Of the 735 patients in our database, 325 (44.22 %) were identified with IEDs. The final sample was comprised of 76 refractory cases. The study included 61 males (80.26 %) and 15 females (19.74 %) ages 5 to 52.

Results
Of the 76 cases treated with anticonvulsants, the vast majority were found to be improved in follow-up progress notes: Improved 65 (85.53 %), Unchanged 6 (7.89 %), and More Severe 5 (6.58 %).

Conclusions
IEDs predict positive treatment outcome to anticonvulsant medication and may not only represent a biomarker for medication selection but also a step towards an evidence-based diagnosis. This review serves as the first large outcome study in which patients with IEDs were treated with anticonvulsants. Our findings suggest that EEG screening should be utilized in all refractory cases regardless of age, gender, or diagnosis. When IEDs are identified, anticonvulsants should be considered as a treatment option.

Consent to publish
This study does not contain details relating to individual participants.

Competing interests
The authors declare that they have no competing interests.

References

A52 Do cannabinoid antagonists affect cognition? SLV326 induces changes in theta and gamma bands in active rats
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A52

Cannabinoid CB1 antagonists have been investigated for possible treatment of e.g. obesity-related disorders. However, clinical application was halted due to their symptoms of anxiety and depression. In addition to these adverse effects, we have shown earlier that chronic treatment with the CB1 antagonist rimonabant may induce convulsive seizures which were EEG-confirmed. However, due to the wide distribution of CB1 receptors throughout the CNS, it is highly unlikely that chronic blocking of the CB1 receptor is only manifested in seizures. CB1 agonists have been described to alter the EEG frequency spectrum. No such data are available for CB antagonists.

In a regulatory repeat-dose toxicity study “muscle spasms” were observed in Wistar rats, daily dosed with the CB1 receptor antagonist SLV326 during 5 months. In selected SLV326-treated and control
animals, EEG and behavior were monitored for 24 h. Subsequently, random segments of the interictal EEG were selected, totaling 20 min per animal. These segments were assigned to subsets of ‘active state’ or ‘passive state’, based on Passive Infrared (PIR) motion detection. Spectral information was calculated using a Fast Fourier Transformation analysis. 25% of SLV326 treated animals showed, EEG-confirmed, spontaneously occurring generalized convulsive seizures, whereas all controls were seizure-free. The behavioral signs of the seizures were typical for a limbic origin. The frequency spectrum of the interictal EEG of the treated rats showed a lower theta peak frequency, as well as lower gamma power compared to the controls. These frequency changes were state-dependent: they were only found during high locomotor activity. However, the treatment did not affect the amount of locomotor activity itself. Apart from confirming our previous finding that long-term blockade of the endogenous cannabinoid system can provoke limbic seizures in otherwise healthy rats, this study shows that SLV326 alters the frequency spectrum of the EEG, but only when rats are highly active. It is therefore likely that the EEG effects caused by SLV326 are linked to higher order behavior that might be present during locomotion. Theta rhythm is shown to be a marker of complex behavior, and gamma rhythm is typically associated with cognitive functions. Therefore, these observations suggest that CB antagonists might have effects on complex behavior and cognition.

Results
Our connectivity analysis found that during eyes-closed, nicotine decreased feed-back connectivity (from precentral gyrus to precentral, angular gyrus, cuneus and superior occipital gyrus) at 10.5-12.4 Hz (α2). During eyes-open, no significant results were found at any frequency range.

Conclusions
We interpreted the results by help of previous anesthesiology literature about an anti-correlated relationship between feed-back and feed-forward connectivity. Such relationship emerged by pharmacologically-induced sedation during eyes-closed condition. Our results suggest that nicotine potentially increases the level of vigilance. Such nicotine-effect is particularly prominent during the eyes-closed condition.

Competing interests
Nothing to declare.

References

Poster presentations
A55
Modulation of the NMDA receptor function enhances hippocampal network oscillations, connectivity and synaptic LTP in vivo: A case study with a Glycine Transporter-1 Inhibitor
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Hypofunction of N-methyl-d-aspartate receptors (NMDARs) has been associated with deficits in synaptic plasticity and cognitive decline as found in neuropsychiatric and neurodegenerative disorders such as Alzheimer’s disease. Glycine and D-serine are endogenous ligands of the NMDAR and therapeutic approaches that enhance NMDAR activity through increases in glycine and/or D-serine levels are expected to enhance synaptic strength and to potentially improve have impact on cognition processes. The present in-vivo study investigated whether positive modulation of brain glycine levels, through modulation by the glycine transporter 1 (GlyT1) inhibitor SSR504734, affects network connectivity and long-term potentiation (LTP) at the hippocampus. For in-vivo network oscillations and connectivity, multichannel EEG recordings were performed in conscious Sprague-Dawley rats from frontal cortical, hippocampal CA1 and CA3 and dentate gyrus (DG) structures after subcutaneous administration of vehicle or SSR504734 (2.5, 10 and 40 mg/kg). For hippocampal synaptic plasticity, rats were anesthetized with urethane and recording and stimulating electrodes were inserted at the DG and at the medial perforant pathway (MPP), respectively. Population spike (PS) amplitudes and excitatory post-synaptic potential (EPSP) slope were measured before and 2 h after high-frequency stimulation (HFS). SSR504734 (at 40 mg/kg) elicited robust EEG slow theta oscillations (4–6.5 Hz) at the DG, CA1 and CA3 and in addition slow gamma oscillations (30–50Hz) in the frontal areas, next to network coherence changes between frontal and CA1 recording sites, which were dissociated from motor behavior. SSR504734 (at 40 mg/kg) enhanced LTP of the PS amplitude after HFS of the MPP synapse, whereas the potentiation of EPSP slope was short-lived. The present data support the hypothesis on a facilitating role of the NMDARs glycine binding site on network oscillations and synaptic efficacy at the medial perforant path of the DG. Future studies will evaluate novel approaches targeting D-Serine modulatory sites, for
example by inhibition of the enzyme d-amino acid oxidase (DAO), which slows the break-down of D-serine, or by its transporter, the alanine-serine-cysteine-1 (Asc-1), the abnormal glia-transmission of which has been linked to synaptic failure in Alzheimer’s disease.

A56
Withdrawn

A57
Neurofeedback training as a treatment for dyslexia
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A57

Dyslexia is amongst the commonest of neurobiological disorders, affecting about 20% of children in Norway. Its heterogeneity makes it difficult to establish a single treatment which is suitable for most of the affected. According to the phonological theory of dyslexia, the disorder is caused by a deficit in the representation, storage and recall of speech sounds. Different brain areas have been linked to the phonological deficit by means of different brain imaging techniques, among other quantitative electroencephalography (qEEG).

The aim of this study was to improve reading ability in children with dyslexia by means of individualized neurofeedback training. The study was conducted as a pre-post intervention single-subject design with 5 participants, aged from 13 to 14 years. The intervention consisted of 25 sessions of neurofeedback training, 15 beta/theta frontocentral sessions and 10 individualized sessions, mostly towards the language areas. The effect of the intervention was measured by means of qEEG and the LOGOS (a Norwegian dyslexia assessment battery). The results showed improvement in reading abilities and phonological skills amongst all participants. Furthermore, qEEG analysis showed increased alpha activity in several brain areas, and normalization of theta and beta activity in comparison to a normative database. An increase of alpha activity may possibly indicate changes in alpha coherence which can be an indication of improved attentional processes. This may explain the improvements in reading and phonological skills. The analysis also confirms the heterogeneity of dyslexia, and the complicity of several brain areas that are involved in dyslexia.

This study is limited by the small number of participants, and the restriction in time (the number of training sessions offered). However, the improvement in reading and phonological skills in this study suggests that neurofeedback training may be an effective and relevant intervention for adolescents with dyslexia. But, further research in this area with larger samples and a larger number of training sessions is required. The study was approved by the regional ethics committee.

Competing interests
There are no competing interests.

A58
Mismatch negativity (MMN) in serotonergic model of psychosis induced by psilocybin
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A58

Background
The auditory MMN (mismatch negativity) is considered to be an index of automatic context-dependent information processing and auditory sensory memory. MMN deficit is a characteristic endophenotype of schizophrenia. A 5-HT2A agonist psilocybin induces acute transient psychotic symptoms and is extensively used as a putative pharmacological model of schizophrenia. Our aim was to investigate the effect of psilocybin on this pre-attentive cognitive functions.

Methods
A double-blind, placebo controlled study design was used. 20 healthy adult volunteers were administered a dose of psilocybin (0.26 mg/kg) and placebo per os in 2 separate sessions. Auditory MMN was recorded in sound and electrically shielded room, 120 min after ingestion of psilocybin/placebo. Participants were lying down with their eyes closed in a comfortable setting with two sitters who were present during whole experiment.

MMN

A single deviant paradigm with 1350 standard (1000Hz, 75 dB SPL, 100 ms duration) and 75 deviant in frequency (1200Hz, 75 dB SPL, 100 ms) tones were presented binaurally in regular order when every 20th was deviant tone. Data was acquired with a standard 32-channel digital EEG amplifier BrainScope (unimed, Prague) with 21 Ag/AgCl scalp electrodes placed according to the 10/20 system and sampled at 1000 Hz.

Results
Mismatch negativity was calculated by subtracting the average of frequently occurring stimuli from the average of deviants. There were no significant differences in latency, absolute amplitude and area under the curve of MMN during psilocybin intoxication compared to placebo. Furthermore, there were no correlations between subjective effects induced by psilocybin (HRS and ASCS) and MMN.

Conclusion
Our results correspond with previous findings [1]. Psilocybin does not affect processing at the level of pre-attentive cognition and the auditory sensory memory.

This effect is probably due to different underlying receptor mechanism as the generation of MMN is strongly dependent on NMDAR dysfunction. Another reason for negative results could be inappropriate timing of recording or insufficient single-deviant paradigm.

This work is supported by Ministry of Health of the Czech Republic, grant nr. 15-29900A. All rights reserved.

References

A59
Comparison between SMR and Upper Alpha Neurofeedback trainings as a non-pharmacological treatment of ADHD and sleep disorders in children and adolescents
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Background
About 5% of school-aged children may have an Attention Deficit–Hyperactivity Disorder (ADHD), a neurodevelopmental disorder often associated with other comorbid conditions including sleep disorders. ADHD became a public health concern. Psychostimulants are the first line pharmacological treatments for ADHD. However, parents are often reluctant to medicate their children and, additionally, a proportion of patients stop their treatment because of side effects. Non-pharmacological treatments are also available. Recently, improvements of cognitive functioning and hyperactivity level of patients with ADHD have been reported after Neurofeedback trainings with a relative Upper Alpha Power enhancement paradigm. Sensorimotor rhythm (SMR) Neurofeedback has been also proposed to improve
ADHD symptoms. The aim of this study is to compare the benefits of Upper Alpha and SMR trainings on ADHD symptoms and concomitant improvement of sleep.

Methods
In this controlled and randomized study, 60 French medication-free children and adolescents with ADHD aged from 8 to 15 years old participated in 30 neurofeedback sessions. They will be assigned to either in the SMR or the Upper Alpha training group. EEG, ADQI rating scales, cognitive assessment, and actigraphic records will be performed at pre-, mid- and post-training times, and 6 months after the end of protocol.

Results
The main expected outcome is the clinical reduction of at least 30% of ADHD symptoms, and we anticipated the superiority of Upper Alpha training over SMR in reducing hyperactivity levels. Improvement of sleep quality is a secondary outcome.

Conclusion
To date, no comparison between SMR and Upper Alpha Neurofeedback trainings with a significant number of sessions and enough patients in each group has been conducted. We hope to gain valuable insights into specific effects of both trainings on ADHD symptoms and sleep without any medication. This study would foster the development of research on Neurofeedback and its clinical applications, which are under-investigated in France.

Trial Registration
NCTD RCB 2016-A00655-46

Keywords
ADHD, Neurofeedback, SMR-Upper Alpha Training, EEG, non-pharmacological treatment, sleep disorders

A60
Effect of Tai-Chi and Cyclic Meditation on hemodynamic responses of the prefrontal cortex
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Keywords
Tai-Chi-Chuan (TCC); Yoga; Cyclic Meditation (CM); Walking; Prefrontal Cortex (PFC); Heart Rate Variability (HRV)

References

A61
Withdrawn

A62
EnkephaloVision: fast dynamic EEG analysis in combination with eye tracking for efficacy testing of plant-derived and low triturated homeopathic drugs.
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A62

Efficacy testing of plant-derived and homeopathic drugs still remains a challenge in pharmacology. In the past several neurophysiological techniques have been successfully applied. However, interpretation of spectral power values with respect to different brain regions is contradictory. Conventional quantitative EEG analysis uses averaged data from epochs of 2 or 4 s. Analysis of shorter epoch length of 364 ms has been achieved by definition of specific frequency ranges [1]. Surprisingly, focal spectral power within these short periods reached tremendous values (up to 9000 %) at single brain regions when compared to the average of other assessed regions (global median) [2]. In order to learn more about these short periods of high electric activity, EEG analysis was combined with eye tracking. The eye tracking software served to present different cognitive and emotional audio-visual challenges in series. Synchronization of the gaze overlay video from the eye tracking with screen capture of the online quantitative EEG analysis was achieved by starting the recording with a gong. The combined technology has been published [3]. Synchronized scenes were evaluated before and after intake of the preparations. In the presence of cognition activating drugs (i.e. Zembrid®) more flashing of delta (1.375 - 4.125 Hz) and theta spectral power (4.125 - 6.875 Hz) was observed in frontal brain in comparison to placebo during performance- and psychometric testing. In the presence of calming drugs (i.e. plant-derived drug Pasconflair®) more flashing of alpha1 spectral power (6.875 - 9.625 Hz) was recognized in comparison
to placebo. The same increase of spectral alpha1 power was detected after intake of 6 homeopathic Calmvalera Hevert tablets at a time. Since about 3 pictures per second are difficult to follow, slow motion videos will be presented. Finally, averaged data were fed into a discriminant analysis. Comparison of several plant-derived and homeopathic drugs with each other revealed for example projection of data from 3 calming drugs in close vicinity to each other.

References

A63 Quantitative EEG assessment of students with ADHD undergoing neurofeedback training
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A63

Several studies have demonstrated abnormalities in quantitative electroencephalogram (qEEG) in children and adults with ADHD. Based on these findings, neurofeedback training (NFT) has emerged as a new treatment option for ADHD. In this preliminary study, qEEG was used to assess the efficacy of NFT training for college students with ADHD. Participants received computer attention training intervention using NFT two times a week over a period of four months. A group of college students with ADHD who did not undergo NFT training was used as a control group. Brain activity was measured using qEEG prior to, midway through, and post NFT training. ADHD behavioral symptoms were also assessed pre- and post- training using the Conners’ Adult ADHD Rating Scale (CAARS-SL) and the IVA-2. Changes in qEEG were detected following NFT. Significant changes in resting-state brain activity were observed in the experimental group. Participants who underwent training demonstrated significant decreases in absolute power across a wide spectrum of frequency bands (delta, theta, alpha and beta) as well as a relative decrease in alpha activity and increase in delta and beta activity. In order to identify if anomalous patterns of brain activity were related to symptoms of ADHD, neuroelectric measures were compared to behavioral measures. Changes in neural activity in the experimental group correlate with improvements in ADHD symptoms. Although further research is warranted to determine the exact impact of NFT on the neural correlates of ADHD, these preliminary findings suggest that it might be a promising cognitive training treatment for students with ADHD.

Keywords
qEEG, neurofeedback, ADHD, ADD

A64 Improving drug discovery using brain oscillations as biomarkers for movement disorders
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A64

Movement disorders represent a group of neurological syndromes characterised by an alteration of voluntary and/or automatic movements. Here, we focused on Parkinson’s disease (PD) and essential tremor (ET), which are the most common forms of movement disorders. Motor symptoms of Parkinson’s disease result from a dysfunction of cortico-basal ganglia circuits mainly due to dopaminergic neurons death in the substantia nigra pars compacta. A hypersynchronization of beta frequency oscillatory activity in these circuits has been described in both patients and animal models of the disease. Essential tremor (ET) is characterized by the symptom of action tremor (which intensifies when the affected muscles are used). ET typically involves a tremor of the arms, hands or fingers. The classically-used animal model of ET is generated by the administration of the beta-carbolin harmaline in mice. Harmaline induces action tremors lasting several hours and the classical read-out is the recording of behavioural tremor frequency that occurs between 8 to 10Hz. The aim of this poster was to provide two examples of the use of brain oscillations in preclinical drug development for movement disorders. Here, 1) we assessed the use of aberrant cortical oscillations in the unilateral 6-OHDA injected rat as translational biomarkers for drug development in PD, and 2) studied the effect of harmaline-induced tremor on cortical and cerebellar oscillatory activities. The sensitivity of these functional biomarkers was challenged with the reference drugs for each pathology. In the 6-OHDA rat model of PD, we found a prominent beta band (~30Hz) in the motor cortex, which was inexcised in control Sprague-Dawley rats. Acute injection of the dopaminergic receptor agonist L-DOPA (6 and 20 mg/kg) induced body rotations along with a significant reduction of the beta band. This treatment also induced a prominent 80-100Hz gamma increase. By contrast, the D2/D3/D4 agonist ropinirole at 0.2, 0.4, and 0.8 mg/kg also decreased the beta band but caused only a slight gamma band increase.

We found that administration of harmaline (10-20-30 mg/kg) in male C57BL/6 J mice dose-dependently increased the cortical and cerebellar power in a wide 15-60Hz frequency range, along with action tremors. Pre-treatment with 20 mg/kg propranolol, one of the first-line medications used in ET patients attenuated the tremors and decreased the 35-60Hz range.

In this study, we identified aberrant EEG oscillations in two rodent models of movement disorders. These oscillations and their pharmacological modulation may represent predictive biomarkers for the identification, selection and validation of new therapeutics in movement disorders.

A65 Telemetric electroencephalography (EEG) and in vivo microdialysis to study dopaminergic hyperactivity in freely moving rats
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A65

Dopamine is a key regulator of cognition, mood, reward and movement in the human and rodent brain. The dopamine homeostasis is tightly controlled by the dopamine transporter (DAT), which is a target for addictive drugs (such as cocaine and amphetamine), and therapeutic antidepressants.

The present study was designed to investigate the effects of cocaine in freely moving rats using telemetric electroencephalography (EEG) to monitor a hyperdopaminergic state. Additionally, in vivo microdialysis in the nucleus accumbens shell was carried out to measure extracellular cocaine levels and dopamine itself by LC-MS-MS and HPLC coupled to electrochemical detection. Behaviour was assessed by an automated motor activity system using light beam interruptions. Cocaine (3, 10 and 15 mg/kg, i.p.) dose dependently induced an increase in motor activity, which reached its maximum level after 20 min and lasted for 90 min. In addition, cocaine appeared to affect the EEG power spectrum, increasing gamma frequency band power up to 90 min after administration, whilst causing a decrease of power in delta, theta, alpha, and beta frequencies. Maximum cocaine levels measured from the dialsates appeared 30 min after dosing.
(300 nmol/l) and extracellular dopamine levels showed a peak concentration at 30 min and then returned to basal levels 120 min later. In conclusion, these results indicate that cocaine induces an increase in dopaminergic transmission in the nucleus accumbens shell, and as expected, produces hyperactivity. The effect observed on the EEG frequency bands and in vivo microdialysis could serve as a physiological biomarker of target engagement studies and to set up a PK-PD relationship in drug discovery research.

A66
Effects of clozapine on auditory steady-state response in schizophrenia
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A66
Auditory steady-state responses (ASSR) provide a non-invasive technique to assess neural synchrony at a particular frequency. Attenuated phase-locking (PLI) of ASSRs in gamma frequency range is observed in schizophrenia and in animal models for psychosis [1]. State-sensitivity of 40 Hz ASSRs has been shown for schizophrenia, where PLIs increased with eyes closure in patients [2]. The effect of clozapine, which is prescribed in cases of treatment-resistant schizophrenia, on ASSRs in humans is not clear. The aim of this study was to evaluate the effect of clozapine use on phase-locking of 40Hz ASSR and state-sensitivity in schizophrenia patients. 48 male patients with schizophrenia (according to ICD-10 criteria) were recruited from the in-patients of Republican Vilnius Psychiatric Hospital. Patients were divided into two groups: (1) resistant to standard antipsychotic medication and treated with clozapine (Cloz, n = 23); and (2) responsive to standard antipsychotic treatment (NCloz, n = 25). ASSRs to click stimuli at 40Hz were recorded using 9 channels in eyes open and eyes closed conditions, with 60 stimuli presented binurally per condition. After conventional cleaning procedures, epochs of 700 ms were created starting at 100 ms prior to the stimulus onset and lasting for 600 ms post-stimulus. ASSRs were analyzed from Cz location, 38-42Hz window was calculated for 100 ms bins and subjected to RM-ANOVA with time bin and task as within-subjects factors and group as a between-subjects factor. Significant interaction of condition (eyes open vs eyes closed) and group (Cloz vs NCloz) factors (p = 0.038) was observed. This suggests that in Cloz group subjects tended to have lower PLIs in open eyes (p = 0.08), which increased with eyes closure (p < 0.001). In NCloz group, PLIs did not change with eyes closure (p > 0.05). Our data propose that state-sensitivity of 40 Hz ASSRs vary depending on the treatment in patients with schizophrenia, subject receiving clozapine showing response increase with eyes closure in contrast to those on standard antipsychotic treatment.

Competing interests
The author declare that they have no competing interests.

References

A67
Neuro-Cardiac-Guided TMS (NCG TMS): a new and cost-effective method for accurately localizing the DLPCA in the treatment of depression
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A67
Background
The efficacy of rTMS in the treatment of major depressive disorder (MDD) has been well established in recent years. Most studies to date have employed the ‘5-cm’ rule for targeting stimulation of the Dorsolateral Prefrontal Cortex (DLPFC). New variations and improvements of this targeting technique include a ‘6-cm’ rule, the Beam-F3 method, and neuronavigated rTMS. Furthermore, it has been proposed that the efficacy of rTMS in MDD is more related to stimulating the area that is functionally connected to the subgenual anterior cingulate cortex (sgACC) rather than to specific cortical areas (Fox et al., 2012). Therefore, we set out to develop and test a new method that employs knowledge about the functional role of the sgACC to establish in real time if the right cortical area is targeted.

Method
Several studies have shown that areas in the ventromedial prefrontal cortex are involved in parasympathetic regulation such as heart rate and respiration, and that neurostimulation of these areas led to heart rate decreases (Makovac et al., 2016), most likely through connectivity with the nucleus vagus. Therefore, based on the notion that rTMS aims to transsynaptically stimulate the sgACC, we used electrocardiogram (ECG) R-peak triggered single pulse TMS to various frontal locations to establish the location that most consistently resulted in a lengthening of the R-R latency (reflective of a heart rate deceleration). This method of Neuro-Cardiac-Guided TMS or NCG TMS thus could be the equivalent of what the Motor Threshold is for the motor system, but then for the DLPFC with heart rate as an output.

Results
First preliminary results using a burst of 10 Hz TMS stimulation demonstrated that in a subject with a relatively large head circumference, no response was found at the ‘5 cm’ site (corresponding to FC4 in this subject), whereas the F4 location did result in a consistent heart rate deceleration. More data are currently being collected using a single pulse R-peak triggered approach and data will be presented. This method is pending for patent. Dutch Patent office: P100241NL0

Conclusions
In the treatment of MDD, Neuro-Cardiac-Guided TMS has the potential to become the equivalent of the ‘motor threshold’ for the DLPFC, and thereby would be a cost-effective and easy to use method for localizing the right stimulation target in the treatment of MDD, and also serve as a real-time control of adequate coil contact in patients undergoing rTMS treatment.

A68
Electroencephalogram connectivity in frontal networks to predict outcome of electroconvulsive therapy in major depressive disorder
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A68
Background
Major depressive disorder (MDD) is a common and potentially lethal disorder affecting up to 14 % of all persons worldwide. However, 1/3 to 2/3 of patients are non-responders to first line therapy [1]. Even the electroconvulsive therapy (ECT) as the option of choice in therapy-resistant MDD still shows a high proportion of non-responders [2]. Due to the invasive nature of the ECT it would be desirable to know which subjects are likely to respond. In case of a predicted non-response to
ECT, e.g. by means of electrophysiological electroencephalogram (EEG) parameters, other therapies of MDD (e.g. augmentation, polypharmacy etc.) could be chosen.

Methods
In this study, we retrospectively analysed two minute resting state EEG from patients with MDD who underwent ECT (4–12 sessions with 3/week) between 2005–2015 at the University Hospital of Zurich. Following several lines of evidence, we hypothesized altered non-linear connectivity in frontal networks including subgenual-, dorsolateral- and medio-prefrontal cortices being predictive for treatment outcome. Symptom severity and response/emission rates were assessed using the Global Clinical Impression (GCI) rating scale. Source estimates and connectivity measures were mapped using Low Resolution Brain Tomography (LORETA).

Results
Responders in comparison to non-responders showed a significant stronger non-linear connectivity in the frontal network within the EEG delta, alpha 1 and beta 1 frequency bands, while connectivity was weaker in theta, alpha 2, beta 2 and gamma frequency bands. Additionally, there were several non-significant correlations (from $r = .15$ to $r = .20$) between symptom change and source estimates with e.g. a low midline theta-activity being associated with response to ECT.

Conclusions
Pre-treatment EEG-connectivity in frontal networks seems to have a predictive value for the efficacy of ECT treatment. Prospective trials following several lines of evidence, we hypothesized altered non-linear connectivity in frontal networks including subgenual-, dorsolateral- and medio-prefrontal cortices being predictive for treatment outcome. Symptom severity and response/emission rates were assessed using the Global Clinical Impression (GCI) rating scale. Source estimates and connectivity measures were mapped using Low Resolution Brain Tomography (LORETA).
Methods
Randomized Clinical Trial (RCT), conform to a randomized, balanced placebo-controlled design with two arms: in condition 1, patients (n = 25) will receive 6 weeks of additional clonidine treatment to their current medication, in condition 2, patients (n = 25) will receive 6 weeks of additional placebo treatment to their current medication. In addition, 25 age and gender matched healthy subjects will function as controls. Primary outcome is change in symptom severity, expressed as a change in total score on the Positive and Negative Symptom Scale (PANSS) from baseline to end of the 6-week treatment. Secondary outcomes are changes in cognitive functioning (measured through the Brief Assessment of Cognition in Schizophrenia; BACS and Cambridge Neuropsychological Test Automated Battery; CANTAB), change in GAF (global assessment of functioning) scores and the measurement of various psychophysiological parameters of basic information processing, such as P50 suppression, prepulse inhibition of the startle reflex (PPI) and mismatch negativity (MMN).

Results
In line with our pilot-study it is expected that early information processing will improve. We predict that this will lead to an improvement in cognitive functioning after six weeks, which expectantly leads to lower symptom severity and a better quality of life.

Trial Registration
EudraCT Number: 2014-003008-53

Conclusion
Our results find that compared to the healthy population, a large number of patients with ASD have IEDs despite never having a seizure. The findings support the use of EEG in children, adolescents, and young adults with ASD, regardless of gender or age. This is particularly true for those who have failed prior medication attempts with stimulants, antidepressants, and/or antipsychotics. Utilizing the EEG for refractory cases in a psychiatric practice allows for more individualized and precise medication selection.

Consent to publish
This study does not contain details relating to individual participants.

Competing interests
The authors declare that they have no competing interests.

References

A71
Isolated epileptiform discharges: an electroencephalographic abnormality underlying medication failure in autism spectrum disorder
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Background
Autism Spectrum Disorder (ASD) often presents a treatment challenge due to the variety of symptoms that make each case unique. Medication prescribed to manage ASD associated symptoms such as anxiety, depression, attention issues, and behavioral problems often fail to alleviate symptoms and can produce undesirable side effects. The question is, why are the stimulants, selective serotonin reuptake inhibitors, and antipsychotics prescribed to alleviate these issues [1] effective in some patients but fail in others? The answer could be related to the increased prevalence of electroencephalographic abnormalities in psychiatric patients [2]. The presence of isolated epileptiform discharges (IEDs) may account for the treatment failure of these medications, especially antipsychotics, because these drugs lower seizure threshold, thus resulting in increased epileptiform activity. Electroencephalography (EEG) can be used to document the presence of IEDs that would otherwise go undetected. The purpose of the study was to reveal the prevalence of IEDs in the ASD patient population and to demonstrate the usefulness of the EEG for providing data to psychiatrists, neurologists, and developmental pediatricians to improve medication selection and outcomes for patients with ASD.

Method
The data was obtained from an Institution Review Board approved data archive from a multidisciplinary practice that treats a wide variety of refractory and neuroatypical patients. The study is comprised of 140 non-epileptic children, adolescents, and adults diagnosed with ASD, ages 4 to 25. A board certified electroencephalographer interpreted the EEGs in order to identify abnormalities.

Results
Of the 140 patients with ASD, 36.4 percent were found to have IEDs after an EEG screening. Chi-square analysis found no significant difference between genders among the three age groups. The findings indicate a high prevalence of IEDs among individuals with ASD.

A72
Neuromodulation using maintenance TDCS optimized by qEEG leads to full recovery from myalgic encephalopathy/chronic fatigue syndrome: a case report
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A72

Background
A 61-year-old man with progressive myalgic encephalopathy/chronic fatigue syndrome was referred for neuromodulation. His condition, likely virally induced decades ago, was characterized by recurring periods of extreme fatigue, lasting months at a time. Severe fatigue had become unrelenting over the prior two years, impairing many dimensions of his life. Multiple immunological and neurological workups were negative and fibromyalgia had been ruled out. Patient failed many medically advised approaches, including antidepressants, acupuncture and a gluten-free diet.

Methods
Genetic analysis suggested he would respond to dopaminergic agents and to neuromodulation [1]. Trials of both amphetamines and methylphenidate ultimately failed but modafinil 200 mg did provide partial relief. Distance from the office precluded daily treatment with repetitive transcranial magnetic stimulation (rTMS). Transcranial direct current stimulation (tDMS) was chosen as a safe alternative feasible treatment [2], allowing cumulative, ongoing treatment to target ongoing inflammation. We used neurophysiological state markers of qEEG. Patient was trained with tDCS in the office and then treatment was self-applied at home daily with anode on left dorsal-lateral prefrontal cortex (LDLPFC), cathode on right (RDLPFC), 2 mA/min, 20 min, 40 mA total dose, using 1.5” diameter electrode pads. After four weeks, maintenance tDCS sessions were increased to twice daily (6 AM and 12 Noon) and modafinil was lowered to 100 mg.

Results
Follow-up qEEG testing was done one year after the initial qEEG when patient was in full recovery. Comparison of pre-treatment and post-treatment qEEG findings show minor improvement in excessive hypercoherent frontal alpha, a substantial 50 % drop in excess left temporal alpha, and a normalization at the very low end of the qEEG spectrum (less than 1 Hz). The patient noted: “this treatment has given me sustained relief from a chronic fatigue condition from which I’ve suffered throughout my adult life.”
Conclusions
Maintenance treatment with daily tDCS and modafinil likely exerted synergistic effects on the brain and immune system. The clinical recovery with notable improved sleep, energy, and ability to tolerate exercise are most likely to be reflected in slow wave oscillations changes. This case supports the need to look more closely at glial as well as neuronal impact, perhaps expanding qEEG to include slow wave markers. Clinicians are eager to have qEEG personalized biomarkers to optimize adjuvantive tDCS stimulation in chronic psychiatric and neurological conditions, so often neuroinflammatory in nature [3].

Keywords
tDCS, myalgic encephalopathy, chronic fatigue syndrome, neuroinflammation, qEEG biomarker, personalized medicine

Consent to publish
Informed consent was obtained and the subject’s rights were protected.

Competing interests
No competing interests.

References

A73
Changes of CNS- and ANS arousal levels following successful antidepressant treatment with ketamine: a case series
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A73

Introduction
Ketamine has been established as an alternative in the treatment of therapy-resistant major depressive disorder (MDD). Although response rates are reportedly high with up to 60-70 %, until now no biomarkers that could predict treatment response exist. As a first step, this case series aimed at identifying electrophysiological markers of arousal that reflect alterations of ongoing neuronal activity after treatment with ketamine.

Methods
Two patients (one 65-year-old female, one 78-year-old male) with therapy-resistant depression (> two treatment -approaches with SSRIs, SNRIs or TCAs) were treated with ketamine infusion four times respectively six times during three weeks. Resting state electroencephalogram (EEG) and electrocardiogram (ECG) were recorded at baseline and after treatment with four/six time ketamine infusion. Central nervous system (CNS) arousal was assessed using Vigilance Algorithm Leipzig (VIGALL). Autonomous nervous system (ANS) function was quantified using heart rate and heart rate variability measures (HRV). Changes of depressive symptoms were assessed using Hamilton Depression Rating Scale (HDRS).

Results
Both patients showed a marked decrease of depressive symptoms with a drop from 28 HDRS to 9 HDRS after four ketamine infusions and from 20 HDRS to 6 HDRS after six infusions respectively. In parallel, both patients showed a decrease of CNS arousal levels as assessed by VIGALL with increased amounts of low vigilance stages and decreased EEG-alpha peak frequencies after therapy in comparison to baseline EEG recording. Further, both patients revealed a lowered ANS arousal level as assessed by a reduction of heart rate >24 h after the last ketamine infusion in comparison to pretreatment condition.

Discussion
Following the arousal framework in MDD with a suggested high EEG-vigilance level in depression, the found decrease of CNS-arousal could be interpreted as a consequence of the anesthetic, i.e. vigilance decreasing effect of ketamine. In contrast, the decrease of heart rate remains elusive in the light of an initial increase of sympathetic function following infusion of ketamine. However, decrease of CNS- and ANS arousal level could lead to less pronounced MDD related behavioral aspects such as withdrawal and sleep disturbances. The predictive value of the EEG in ketamine treatment should be in the focus of further prospective randomized studies.

Consent to publish
Written informed consent has been obtained by all patients prior to publication.

Competing interests
The authors report no competing interests.

A74
ECoG spectral analysis of the Interaction between caffeine and nicotine
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A74

Caffeine and nicotine are the most consumed psychostimulants worldwide. Although the electrophysiological effects of each drug alone were studied extensively, the literature on the effects of their combined treatments on brain electrical activity is scarce. The present study aims to investigate the effects of the intraperitoneal injection of caffeine followed by the subcutaneous injection of nicotine after 1 h on electrical activity recorded from the cortex of rats (ECoG). It was found that the successive injection of caffeine and nicotine resulted in a significant increase in the power of delta frequency band but a significant decrease in the power of theta, beta-1 and beta-2. It was suggested that the caffeine and nicotine interaction could have an adverse effect by altering the cortical electrical activity that may indicate impair in memory encoding.

A75
Dishabilitation of central nervous system to tonic pain following chiropractic care - a standardized low resolution brain electromagnetic tomography (sLORETA) based study
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A75

It has been demonstrated that after chiropractic spinal manipulation neural plastic changes occur in different areas of the brain. Different methods have been utilized to assess these changes, but the majority of the measurements to find the involved brain areas have been indirect. The objective of this study was to determine the changes in brain
activity during tonic pain after single session of chiropractic care in a sub-clinical pain population by using source localization of the EEG. 

Fifteen healthy volunteers (10 males, 32.1 ± 7.2 years) participated in two experimental sessions on separate days; chiropractic or control (sham) session in random order. The EEG was recorded continuously using a 61-channel system before and after either intervention during 72 s of cold pressor test at 2 °C (left hand). The pain and unpleasantness ratings were obtained on two separate numeric scales (range: 0 = no unpleasantness/pain to 10 = maximum unpleasantness/pain). The EEG was divided into 9 epochs (8 s each), which were separated into four frequency bands: delta (1–4 Hz), theta (4–8 Hz), alpha (8–12 Hz) and beta (12–32 Hz). Subsequently, standardized low resolution brain electromagnetic tomography (sLORETA) was done on these frequency bands. 

In the control experiment, the brain activity decreased in all frequency bands (all p < 0.05). The decrease in activity in frequency bands (all p < 0.05). In the control experiment, the brain activity decreased in all frequency bands. 

The decrease in brain activity in the control arm reflects central habituation which occurs due to repetitive painful stimulation. The lack of this phenomenon in the chiropractic arm could imply that the chiropractic care normalizes the central nervous system leading to central dishabituation.

References


A76 Animal 3D brain-mapping 

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A76

Introduction

To this date there are no standardized mapping methods that display animal cortical EEG on the brain surface. Therefore, this study describes a 3D imaging method to be used for EEG mapping on the surface of the rat brain. The aim of our study was to develop a software module and a standard for statistical brain mapping. Animal EEG data recorded during behavioral activity and inactivity served as a subject for analysis and brain mapping.

Methods

In this study we measured electrical activity of the rat brain. For imaging purpose, we used 3D brain model from atlas [1] and adjusted it for our own module. We confirmed the validity of the 3D brain model by comparing the dimensions of normalized brain scans of 9 rats of the Wistar strain typically used in our laboratories. We have created a MATLAB module for brain mapping with the use of the rat brain model and a possibility to place any number of electrodes on the surface of the rat brain. The spline interpolation was used for imaging activity on surface of the brain. The spline interpolation was used to compare the two example behavioral conditions. 

Results

The module was effectively used to display EEG activity on the 3D surface and to display the statistical group differences in the sample of the animal data between behavioral activity and inactivity. The module can also compare data from individual measurement with a group mean.

Conclusions

This study describes computation of splines interpolation curves that are important for the brain mapping in rats. This approach will be used for effective comparisons of brain activity of rats under various conditions and with variable number and placement of cortical electrodes.

This study was supported by the project Nr. LO1611 from the MEYS under the NPU I program, by project “National Institute of Mental Health (NIMH–CZ) (grant number ED2.1.00/03.0078 from the European Regional Development Fund), by Czech Technical University research program SGS (SGS15/229/CHK4/3 7/17), MH CZ–drq (National Institute of Mental Health – NIMH) project nr.: 00023752 and PRVOUK34. I declare no conflict of interests.

A77 Effects of γ-aminobutyric acid-modulating drugs on resting state brain oscillations and executive function in healthy volunteers: a pilot study

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A77

Background

Recent findings have suggested a relationship between abnormal γ-aminobutyric acid (GABA) function, disordered neuronal oscillations, and impaired executive function in schizophrenia. Additionally, there has been an increasing amount of interest in the therapeutic potential of these drugs in the treatment of this disorder. However, the neural oscillations that underlie the effects of GABA-modulating drugs on cognitive functioning require further work.

Objective

In an attempt to begin identifying which receptor subtypes may alter the neural oscillations underlying executive function via selective agonist actions, the study examined the effects of: a) a benzodiazepine drug with broad spectrum agonist actions at all GABA receptors containing the α1, α2, α3, and α5 subunits, and the γ subunit (in addition to the obligatory β subunit) and b) a drug with agonist actions at GABAA receptors. The objective of this pilot study was to examine the effects of single doses of these GABA enhancing drugs on resting state brain oscillations and executive function in healthy volunteers stratified by executive function performance.

Method

30 participants were assessed in a randomized, double-blind, placebo-controlled design. Three minutes of eyes closed resting state brain oscillations were measured from 8 electrode sites in response to an acute administration of lorazepam (Ativan®; 1.0 mg), a GABAA receptor positive allosteric modulator, and baclofen (Lioresal®; 10 mg), a GABAB receptor agonist. Executive function was assessed using the Groton Maze Learning Task (GMLT) of the CogState Schizophrenia Battery. 

Results

Spectral analysis revealed overall reductions in alpha and theta oscillations with the lorazepam treatment. Follow-up analyses indicated that these reductions were in the better performing participants. Correlational analyses revealed that greater lorazepam-induced reductions in alpha and theta oscillations were associated with greater
lorazepam-induced cognitive impairment. Reduced theta at placebo was also associated with worse performance. Additionally, smaller theta activity at placebo was associated with greater lorazepam-induced cognitive impairment.

**Conclusion**

The results suggest that GABA_A-modulated alpha and theta oscillations are involved in the neural underpinnings of executive processing.

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**A78**

**EEG machine learning for enhanced monitoring of Alzheimer’s disease and cholinergic modulation**

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Scopolamine is a muscarinic acetylcholine receptor antagonist (mACHR) that induces cognitive impairments resembling those observed in Alzheimer’s disease (AD) and schizophrenia. It is used in drug development to demonstrate the reversal of the temporary scopolamine-induced cognitive deficits by a cognition enhancing compound. However, there is an urgent need for biomarkers that monitor therapeutic response; current biomarkers lack the desired accuracy, because of the large variability in healthy subjects and the often subtle disease-related changes. In EEG, pathophysiology is often expressed in multiple ways. Here we show that an integrative approach in which any biomarker that carries complementary information about a disease or therapeutic intervention can result in an accurate diagnostic index for better decision making in clinical trials.

Recently, we showed that EEG biomarker integration improves the prediction of conversion from mild cognitive impairment to Alzheimer’s disease (AD) compared with a single-biomarker based prediction [1]. The integrative biomarker index can be used for stratification of patients at recruitment in clinical studies and for documenting and quantifying effects of intervention. Here, we provide additional proof-of-concept that EEG-based prediction can be improved with the integrative biomarker approach in clinical trials where a drug is tested in a scopolamine challenge model in healthy subjects. For this purpose, we have developed an integrative EEG biomarker index (mACHR index) that is optimally sensitive to the CNS effects of scopolamine, to objectively determine whether reversal of scopolamine effects by a cholinergic compound is successful. The mACHR index yielded higher classification performance than any individual EEG biomarker with accuracy, sensitivity, specificity and precision ranging from 88–92%. This significantly outperforms the single-best EEG biomarker (relative delta power). Validation on an independent dataset indicated the robustness of the index. To support the validity of scopolamine as a model for AD pathophysiology, we show that the mACHR index discriminates healthy elderly from patients with AD.

We address this by using novel features of the Neurophysiological Biomarker Toolbox (http://www.nbtwiki.net/), which employ data mining algorithms to combine the information from multiple biomarkers. Our results demonstrate that integrating information from multiple EEG biomarkers can enhance the accuracy of identifying disease or drug intervention, which should be of interest to a wide range of clinical trials.

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**A79**

**Midfrontal theta dynamics reflect the ability to overcome motivational biases in decision making**

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A79

Our motivations influence our actions in predictable ways. The promise of a reward promotes behavioural activation, while the threat of a punishment context promotes inhibition. However, these motivational biases can at times be at odds with our goals. At such times, we need to be able to suppress them, which has been suggested to be implemented by the midfrontal cortex. We developed a novel paradigm and computational models of behavior to disentangle the impact of such motivational response biases, from the impact of learning from reward and punishment outcomes. Participants (N = 34) completed this task while recording surface EEG. As expected, cue valence strongly biased action. Midfrontal theta-band oscillatory activity was increased in those trials, where the motivational response bias conflicted with the required response, particularly when subjects successfully suppressed the motivational bias. We will present further analyses to dissociate the role of midfrontal cortex in learning from reward and punishment outcomes. This work will allow us to characterize how motivations drive biases in both choice and learning, and how we may learn to suppress these when they are at odds with our instrumental goals. This work has relevant implications for a range of psychiatric disorders associated with a maladaptive reliance on impulsive, motivation-driven responding including addiction, impulse control and ADHD.

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**A80**

**Neurophysiological substrates of memory processes: assessment of glutamatergic and cholinergic modulation of sharp wave ripples in rats**

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A80

Sharp wave ripples (SPW-Rs) represent the most synchronous population patterns observed in the mammalian brain and are considered a cognitive biomarker for episodic memory and planning. SPW-Rs occur during several off-line states of the brain including non-REM sleep; are modulated by many neurotransmitter systems; and affect both cortical and subcortical structures by their excitatory output. Selective disruption of SPW-Rs impairs memory formation and pathological SPW-Rs have been observed in rodent models for neurodegenerative diseases. Quantification of these synchronous population patterns associated with memory processes is instrumental for a better comprehension of neurodegenerative diseases as well as for assessing the efficacy of novel pharmacological treatments. The purpose of this study was twofold: first, to develop and val-

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idate a novel computer-automated touchscreen-based spatial search task assessing either working memory or memory consolidation in Long-Evans rats; second, to quantify SPW-Rs’ activity in this spatial search task during working memory or memory consolidation combined with pharmacological glutamatergic and cholinergic modulation. For the working memory component of the task, rats had to find a hidden location on the touchscreen with either a short (2 s) or long (10 s) delay between 10 consecutive trials with each delay having 4 different locations presented within one session. During these delays, hippocampal SPW-Rs from the CA1 stratum pyramidale cell layer were measured following each completed trial, using implanted 4-shank silicon electrodes. Here, SPW-Rs were measured when the rat was moving at speeds of less than 4 cm/s, by use of video monitoring to ensure events analyzed were associated with quiescent periods only. Results indicate that Scopolamine 0.1 mg/kg but not 0.05 mg/kg decreased performance for the long but not for the short delay. For the memory consolidation component of the task, rats received 1-day or 4-day acquisition session/s of a single hidden location with variable encoding strength using few (10) or many (>40) trials per session. Memory consolidation of the location was measured 24 h after acquisition by the use of a probe trial. SPW-Rs were measured when the rats were asleep both before and after the acquisition session. A differential effect on memory consolidation was addressed using pharmacological manipulation of glutamatergic and cholinergic systems. This study reinforces the strength of combining neurophysiological and cognitive behavioral assessment to further understand memory processes and effects of pharmacological treatments thereon.

A81
Glutamatergic deficit and negative symptoms: new evidence from the ketamine model of schizophrenia
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1): A81

Targeting the N-methyl-D-aspartate-receptor (NMDAR) is a major approach for treating negative symptoms of schizophrenia. The ketamine model of schizophrenia has the advantage of comprehensively producing schizophrenia like symptoms such as positive, cognitive and negative symptoms in healthy volunteers. The amplitude of theMismatch Negativity (MMN), a neurophysiological parameter related to infrequent stimuli, is known to be significantly reduced in schizophrenic patients but also in healthy controls receiving ketamine [1,2]. Accordingly, it was the aim of the present study to investigate whether changes of MMN during ketamine administration are related to the emergence of negative symptoms in healthy subjects. Therefore, we examined the impact of ketamine on MMN amplitudes and its sources (sources localization approach: low resolution electromagnetic tomography (LORETA)) by means of 64-channel electroencephalography (EEG) recording during performance of an auditory MMN paradigm and assessed the psychopathological status using the Altered State of Consciousness (SD-ASC) Rating Scale and the Positive and Negative Syndrome Scale (PANSS). Twenty-four male, healthy volunteers were measured versus with pharmacological EEG using a single-blind, randomized, placebo-controlled crossover design. We identified significant changes of the MMN response, to both duration and frequency deviants, under ketamine condition as well as a significant increase in all PANSS scores. Reductions of MMN amplitudes were significantly correlated with more pronounced negative symptoms, assessed by the PANSS.

Accordingly, the MMN might represent a biomarker for negative symptoms in schizophrenia related to an insufficient NMDAR system and could be used to identify schizophrenia patients with negative symptoms due to NMDAR dysfunction and thus to determine a maximal benefit of drugs modulating neurotransmission at the NMDAR.

Competing interests: The authors declare that they have no competing interests.

References

A82
Modulation of the serotonin system in an animal model of psilocin-induced psychosis – time course of quantitative eeg changes
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1): A82

Introduction The serotonergic hallucinogen psilocybin and its active metabolite psilocin nowadays receive a lot of attention in the scientific community as a research tool for modeling psychosis. First experiments assessing brain activity after psilocybin administration in humans using PET and fMRI found contradictory results [1,2]. More recently, a study directly measuring neuronal activity using MEG confirmed massive inhibition of brain activity [3]. The aim of our animal study was to assess psilocin-induced changes in quantitative EEG (QEEG) in rats in order to explore the role of different serotonergic receptors in psilocin action.

Methods The substances used were: psilocin (4 mg/kg s.c.), 5HT1A antagonist WAY 100635 maleate (1 mg/kg s.c.), 5HT2A antagonist MDL-100907 tartrate (0.5 mg/kg s.c.), 5HT2C antagonist SB-242084 (1 mg/kg s.c.), haloperidol 0.1 mg/kg s.c. and clozapine 5 mg/kg i.p. For EEG experiments, rats were stereotactically implanted with 12 active electrodes onto the surface of the cortex under isoflurane anesthesia. EEG was recorded in freely moving rats after one-week recovery from surgery. EEG power spectra (local synchronization) and coherence (long projections) were subsequently analyzed comparing the drugs’ effect in time (20–30, 50–60 and 80–90 min post administration) to the baseline record. To avoid moving artifacts and effects of behavior on EEG, only EEG traces corresponding to behavioral inactivity were included in the analysis.

Results Psilocin generally decreased both EEG absolute spectral power and EEG coherences. The changes in spectral power induced by psilocin were normalized by all substances used, mainly in the lower frequency bands. However, only 5HT1A and 5HT2A antagonists partially normalized the psilocin-induced decrease of EEG coherences. The specific QEEG pattern of each substance and the temporal dynamics of QEEG changes will be presented.
Conclusions
Psilocin-induced changes in QEEG in rats are very similar to our recent human data with psilocybin and are in accordance with the concept of psychosis as a disconnection syndrome. All the specific 5HT antagonists and both antipsychotic drugs specifically affected the EEG spectral power by psilocin. Surprisingly, only SHT1A and SHT2A antagonists were able to partially reverse psilocin-induced disconnection. These results indicate that SHT1A and SHT2A receptors might be involved in the increase of entropic brain activity during psychedelic state as well as acute psychosis. This study was supported by the grant IGA MZCR NT/13897, by Charles University research program PRVOUK P34, by project “National Institute of Mental Health (NIMH–CZ)”, grant number ED2.1.00/03.0078, and by the European Regional Development Fund. I declare no conflict of interests.

References

A84
P300 in pharmacological models of psychosis
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A84

Background
P300 (P3) is an index of focal attention processes and memory updating. Impaired cognition is one of hallmark features of psychotic disorders. Both psilocybin (5-HT2A agonist) and cannabinoid induces acute transient psychotic symptoms and have previously been used as putative models for psychosis. In order to investigate the extent of cognitive disruption during psilocybin and cannabis intoxication, information processing was evaluated by means of both sensory event related potentials (P2, N2) and cognitive potential P3.

Methods
Data from two separate studies are presented. 1) In a placebo-controlled design, 20 healthy adults were administered a dose of psilocybin per os (0,26 mg/kg) and placebo during 2 separate sessions. 2) In an ecologically valid model of cannabis intoxication, 34 recreational users, 32 chronic users and 30 healthy age- and gender-matched cannabis non-users were recruited. ERPs were recorded in a sound-attenuated room with each participant lying down with their eyes closed in a comfortable setting with two sitters (male and female) being present for the whole time. An oddball paradigm with 120 frequent and 30 target tones presented binaurally in a pseudo-random order was used. Data were acquired with a standard 32-channel digital EEG amplifier BrainScope (unimeds, Prague) with 20 active scalp electrodes and oculogram according to the 10/20 system.

Results
Psilocybin: A repeated-measures ANOVA on latencies and amplitudes of P2, N2 and P3 revealed a main effect of 5-HT2c agonists have been suggested to exhibit antidepressant-like profile that fits to the sleep changes observed in our study. Further, 5-HT2C agonists have been reported to inhibit theta oscillation, desynchronizing the EEG and leading to shifts to lower frequencies [2,3]. Yet, despite the inhibition of theta oscillation and desynchronization of the EEG by CP-809,101, the ratio between delta/theta revealed no changes underlying the wake-promoting effects of CP-809,101.

Disclosures
All authors are employees of AbbVie. The design, study conduct, and financial support for this research was provided by AbbVie. AbbVie participated in the interpretation of data, review, and approval of the publication.

References

A83
Novel methods assessing electrophysiological alterations by 5-HT2C receptor agonist CP-809,101 in sleep EEG and power spectral activity
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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A83

The serotonergic 5-HT2C receptor is a key contributor to a variety of medical conditions including psychiatric and neurological diseases. The development of therapeutic approaches at this receptor, with both, agonists and antagonists continues to be in focus [1]. Using a novel wireless EEG device (Neural Activity Tracker-1) and a novel in-house developed statistical algorithm we investigated electrophysiological changes in sleep structure and EEG power spectral distribution caused by the highly selective 5-HT2C receptor agonist CP-809,101.

In two independent studies, male Fischer rats with chronically implanted suprachrival EEG-electrodes were treated with 10 mg/kg of CP-809,101. In the 1st study, sleep structure changes in terms of total sleep time, percent of time spent in different vigilance states, the number of rapid eye movement (REM) episodes, and latency to first REM episode were analyzed. Treatment with CP-809,101 led to attenuation of time spent in mild, deep, and REM sleep. It increased time spent in wake state and latency to first sleep and first REM episode. The 2nd study investigated power spectral distribution changes. A refined statistical method of baseline-adjusted power spectral changes revealed an attenuation of delta and theta band by CP-809,101 in comparison to vehicle recordings while maintaining the delta/theta ratio. Our results clearly demonstrate that acute treatment with CP-809,101 changes both sleep architecture and power spectral parameters in Fischer rats.
In major depressive disorder (MDD) research, frontal alpha asymmetry (FAA) has frequently been reported as a potential discriminator between depressed and healthy individuals, although contradicting studies and non-significant results have been published [1, 2]. Locating an MDD biomarker could benefit many people, as MDD is predicted by the WHO to become the second most debilitating disease by 2020. The aim of the current meta-analysis is to clarify the relationships between MDD and FAA further, through analyzing new research from the last decade and put it in perspective by comparing current and past findings (for example a meta-analysis [1]).

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Cohen’s d will be calculated from the means and standard deviations for FAA measures (subtracting mean log transformed left midfrontal alpha from mean log transformed right midfrontal alpha $\ln(F4) - \ln(F3)$), or a similar measure. Possible covariates including age, gender, handedness, year of publication, country of residence, depression severity, medication, EEG recording length, keeping eyes either open, closed or both, EEG reference, and used alpha frequency will be explored. A study will be included if the article (1) reports on both depressed and healthy individuals, (2) provides an FAA measure involving F3 and F4, and (3) provides all data regarding above mentioned covariates (reported either directly or obtained through contact with corresponding authors).

Preliminary results of our currently ongoing meta-analysis will be presented. On the one hand, previous studies have reported relative more left-sided alpha in MDD (sometimes only for higher frequency alpha and not for every EEG montage). On the other hand, non-significant and even opposite results have been reported, showing no baseline FAA differences between depressed patients and controls, or finding relatively more right-sided frontal alpha. Our expectation is that there will be no difference in FAA between MDD and non-MDD groups, based on more recent studies reporting contradicting results, as well as today’s largest investigated sample regarding this topic, the iSPOT-D study [2], showing non-significant results. If non-significance is indeed demonstrated, the use of FAA as a diagnostic tool can be questioned. Nevertheless, its contribution to other applications (such as treatment prediction) could be further explored.

References

A85 Frontal alpha asymmetry in depression: fact or fiction?

A meta-analysis

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A85

In major depressive disorder (MDD) research, frontal alpha asymmetry (FAA) has frequently been reported as a potential discriminator between depressed and healthy individuals, although contradicting studies and non-significant results have been published [1, 2]. Locating an MDD biomarker could benefit many people, as MDD is predicted by the WHO to become the second most debilitating disease by 2020. The aim of the current meta-analysis is to clarify the relationships between MDD and FAA further, through analyzing new research from the last decade and put it in perspective by comparing current and past findings (for example a meta-analysis [1]).

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References

A86 Neuropsychological profile of selected areas responsible for the inhibition of P50 wave: from the P50 wave to off-label treatment of schizophrenia

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Neuropsychiatric Electrophysiology 2016, 2(Suppl 1):A86

Patients suffering from schizophrenia have been shown to exhibit impaired P50 ERP amplitude-reduction to the second (S2) relative to the first (S1) of identical brief auditory stimuli. This reduction is often mentioned in connection with the inability to filter redundant sensory stimuli typically manifested as inability to gate neuronal responses related to the P50 wave [1, 2]. The key neuronal structure responsible for the sensory gating process is the hippocampus. Inhibition of redundant stimuli in the hippocampus is affected via the release of glutamate from excitatory pathways, which is controlled by GABAB receptors. It is closely connected with a physiological deficit of hippocampal GABAergic interneurons, which demonstrates neuropathological changes in schizophrenia. Several drugs are able to improve sensory gating, the effect of which is explained by their ability to disinhibit GABAAergic neurons in the hippocampus. The effect of setrons may be an example of such effective GABAergic interneurons disinhibition. This antagonist of 5-HT 3 receptors increased (by disinhibition of GABAergic interneurons) release of acetylcholine, which by agonism of alpha7 nicotinic receptors improved auditory gating [3]. Besides the hippocampus the prefrontal cortex is an important neuronal part of the sensory gating. Patients with a prefrontal damage fail to suppress irrelevant sensory information, which leads to increased neural noise and inability to inhibit task-irrelevant information during behavioral tasks requiring performance over a delay. Some of the P50 source analysis leads to the conclusion that while the temporal cortices are the main generator of the P50 component, the prefrontal cortex seems to be a main contributor to the process of sensory gating (P50 amplitude reduction) [4]. As in the case of the hippocampus, there are drugs that improve sensory gating by acting on the prefrontal cortex. Clonidine acts as an agonist of alpha2 noradrenergic receptors and has a proven restorative effect on sensory gating. Stimulation of alpha2 noradrenergic receptors on PFC spines by clonidine leads to strengthening of network connectivity, increase in neuronal PFC firing, and thus improves PFC regulation of sensory gating [5]. The aim of our poster is to interlink a pharmacological profile of neuronal areas that are involved in the inhibition of P50 wave with clinical treatment of schizophrenia. We believe that the neuropsychological aspects of P50 wave offer an interesting hypothesis relating mainly to the pharmacological augmentation strategies. Some of them are suggested and explained further in our poster communication.

This work was supported by the project „National Institute of Mental Health (NIMH–CZ)“, under grant number 281.1.00/03.0078, the European Regional Development Fund, the Charles University research program PRVOUK P34, and the travel grant of the Czech neuropsychopharmacological society.
Dopamine under the influence of sunlight? Transitions in solar irradiation explaining attentional performance in DRD4 7R carriers

Background
Previous research suggests that high exposure to solar irradiation has a preventive effect on the development of attention-deficit/hyperactivity disorder (ADHD) [1]. Note that the Dopaminergic DRD4 receptor is involved in phototransduction in the retina. Interestingly, being a DRD4 7R carrier while being born in spring and summer has been demonstrated to result in a 2.8 higher likelihood of developing hyperkinetic disorder, equivalent to ADHD [2]. These findings suggested a possible gene X environment interaction between the DRD4 7R allele and season of birth. The current study focused on the influence of solar irradiation exposure around birth on adult attentional performance.

Methods
We used an RDoC approach focusing on “inattention” operationalized as false negative errors, i.e. missed targets, from two cognitive tasks; the auditory oddball task and the continuous performance task. DRD4 genotype was regarded a vulnerability to develop ADHD, i.e. high inattention. We specifically aimed to test hypotheses that we generated based on previous studies. We distinguished the solar irradiation data.

Results
Results showed an interaction between DRD4 genotype and transition in solar irradiation following birth on the number of inattention errors made (F(1, 269) = 6.785, p = .010). More specifically, a one-way ANOVA for the DRD4 7R carrier group showed a significant difference between positive and negative transition in solar irradiation (1.86) = 8.602, p = .004, d = −0.449), while data from participants lacking the DRD4 7R genotype did not differ.

Conclusions
These results provide evidence that factors around birth influence adult performance and may strengthen or weaken the risk to develop attention related problems once already genetically at risk. Results also further strengthen the hypothesis that a relationship between solar irradiation and ADHD exists, possibly mediated by the dopamine DRD4 receptor.

References