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Analysis of laminar profile of evoked potentials in rat visual cortex

Master's degree thesis field of study Applications of Physics in Biology and Medicine speciality Neuroinformatics

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Summary

Key words

electrophysiology, visual system, primary visual cortex, visual stimulation, local field potential, current source density, temporal frequency elektrofizjologia, układ wzrokowy, pierwszorzędowa kora wzrokowa, stymulacja wzrokowa, lokalny potencjał polowy, current source density, częstotliwość czasowa po polsku czy po ang?

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The title of the thesis in Polish

Analiza profilu warstowego potencjałów wywołanych w korze wzrokowej szczura

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Goal

The main goal of this work was to study responses of neurons from particular layers of visual cortex to visual stimulation with different temporal frequency.

Chapter 1

Introduction

1.1. Structure of the visual system

Visual perception is an ability to process information coded by light. There are two largely independent visual pathways. Both begin in retina where light is converted into electrical discharge by photoreceptors and then the information is transferred to ganglion cells and then further via the optic nerve (Fig. 1.1). Here the pathways split. The first one, known as geniculate visual pathway, leads via the dorsal part of lateral geniculate nucleus (LGN) to primary visual cortex (VCx), and from there further to higher-order cortex. The second pathway, known as extrageniculate visual pathway, leads through superior colliculus and the lateral posterior nucleus-pulvinar complex and then—similarly to the first pathway—reaches primary visual cortex and the higher-order cortex (Waleszczyk et al., 2005; Thomson, 2010).

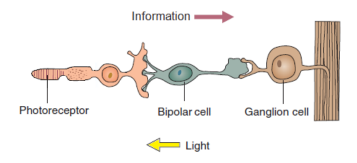


Figure 1.1: Neural circuitry in the retina (modified from Carlson, 2013).

1.2. Parallel processing in the visual system

Retina is the first place where visual information is decoded and segregated. Three main morphological types of ganglion cells recognize different properties of visual stimuli (Fig. 1.2).

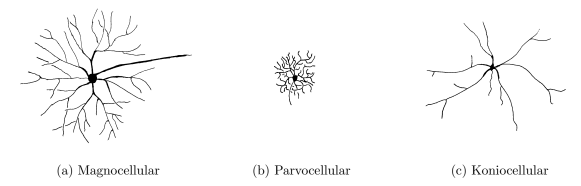


Figure 1.2: Three most important morphological classes of ganglion cells (Drawing on the basis of Stone, 1983).

Magnocellular cells (Fig. 1.2a) have big soma with large dendritic tree. Thanks to thick axon this type of cells can recognize fast stimuli but have poor spatial frequency. Parvocellular cells are cells with medium soma (Fig. 1.2b). They are characterized by high resolution spatial frequency due to small dendritic size, but weak temporal frequency resolution. Koniocellular cells are heterogeneous group of cells with small-to-medium somata of different dendritic morphologies (Fig. 1.2c; Stone, 1983; Waleszczyk et al., 2005).

1.3. Architecture of primary visual cortex

Primary visual cortex can be divided into six vertical layers differing from one another in function and morphology. Sometimes layers II and III are considered to be one layer II/III. Parallel processing is continued from retina via LGN to VCx. Magnocellular channel enters mainly to layer IV but also to layer VI; parvocellular channel enters layers IV (lower than magnocellular) and VI (higher than magnocellular); koniocellular channel enters layers I, II/III and V (Fig. 1.3; Stone, 1983).

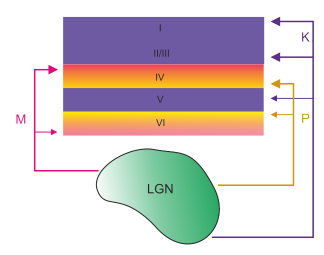


Figure 1.3: The main connections made by axons from the dorsal lateral geniculate body to primary visual cortex.

Other division can be made based on granule cells located in layer IV. These cells are characterized by very small cell bodies. Layers above layer IV are called *supragranular* while layers below are referred to as *infragranular*. Around layer IV there is a polarization change observable on averaged evoked potentials (Maier et al., 2010). In layer I cell bodies are absent but there are plenty of axons, dendrites, and synapses. Two major classes of cortical cells are pyramidal cells which occur in all layers except I and IV, and stellate cells which are found in all layers (Fig. 1.4).

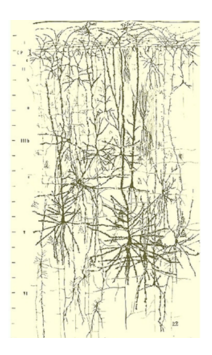


Figure 1.4: Composite figure of a mosaic of camera lucida drawings showing the morphological features (size, location, and distribution) of the principal neuronal types of the human cortex. From rapid-Golgi preparations. Scale (on the left): 100 μ m (Schmolesky, 2007).

1.4. Local field potential

Electrophysiology is a method of recording brain activity characterized by high temporal resolution but weaker spatial resolution. Thanks to electrodes inserted in different places in the brain we can gather information from them. Electrophysiology can be used to examine responses to various stimuli at different locations of the brain. Usage of vertical electrodes allows to record simultaneously signal across the brain.

Local field potentials (LFP)—i.e. the low-frequency part of extracellular electrical recordings from electrodes inserted into the examined tissue—are a measure of the neural activity reflecting dendritic processing of synaptic inputs to neuronal populations. LFP is a superposition of postsynaptic and action potentials (Fig. 1.5).

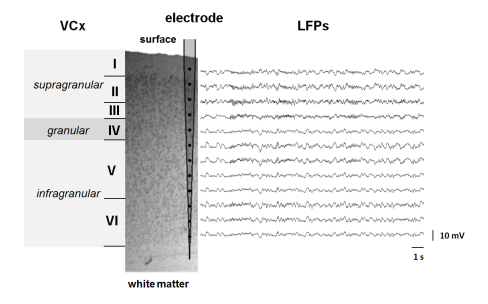


Figure 1.5: An example of LFP signals recorded in VCx.

Postsynaptic potentials are potentials generated in the postsynaptic membrane of a dendrite of a nerve cell. They can be either excitatory (Excitatory Post-Synaptic Potentials, EPSPs)—that enhance chance of induces action potentials—or inhibitory (Inhibitory Post-Synaptic Potentials, IPSPs)—making action potential less likely to happen. There are lots of potentials reaching every neuron. When the sum of them exceeds a threshold the neuron becomes excited and generates an action potential which propagates along axon (Fig. 1.6).

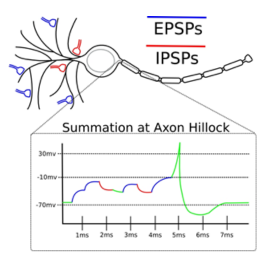


Figure 1.6: Excitatory and inhibitory post-synaptic potentials.

Chapter 2

Materials and methods

2.1. Subjects

For electrophysiological experiments presented in this study, we used 6 adult male Wistar rats (250-300g). All animals were housed with free access to food and water and maintained on a 12 h light/dark cycle. All experimental procedures were conducted in accordance with the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research and the EC Directive 86/609/EEC for animal experiments using protocols and methods accepted by the First Warsaw Local Ethical Commission for Animal Experimentation. The experiments took place in Laboratory of Visual Neurobiology in Nencki Institute with great help from Katarzyna Kordecka.

2.2. Surgical procedures

Animals under deep urethane anesthesia (1.5 g/kg, Sigma-Aldrich, Germany, 30% aqueous solution, i.p) were placed in a stereotaxic apparatus. Additional doses of urethane (0.15 g/kg) were administered when necessary. Body temperature was maintained between 36 and 38 °C using an automatically controlled electric heating blanket. Every hour fluid requirements were fulfilled by subcutaneous injections of 0.9% NaCl (1ml/hour) and eyes were humidified with Vidisic (Polfa Warszawa S.A., Poland) to prevent cornea from drying. The skin on the head was disinfected with iodine and local anesthetic with 1% lidocaine hydrochloride (0.5 ml, Polfa Warszawa S.A., Poland) was injected over the scalp. The skull was opened to expose areas of the binocular primary VCx (7.5 mm posterior to bregma, 5.0 mm lateral) in left hemisphere. Coordinates of electrode were chosen based on the rat brain atlas (Paxinos and Watson, 2007).

2.3. Stimulation electrode

For all experiments the same custom vertical electrode made of tungsten was used (Fig. 2.1). The electrode had 12 channels but only recordings from 7 of them were used for analysis. Particular channels of the electrode were chosen independently for each rat and the choice was made after the recording. This procedure was motivated by previous experiments which had shown that it is very difficult to insert the electrode in such a way that a predetermined channel corresponds to layer IV. The cause for this difficulty is that layer IV is very thin. It is also the reason why upper contacts of the electrode were located more densely (0.1 mm intervals) than the lower ones (0.2 mm intervals).

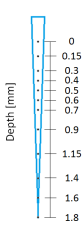


Figure 2.1: Scheme of 12-channel vertical electrode with intervals between contacts expressed in mm.

Although the same electrode was used for all the experiments, the exact depth of each channel varied. This was due to anatomical differences between subjects and imperfect spatial accuracy of the applied method. Therefore channel numbers do not correspond to the same actual physical channels in all experiments. Table 2.1 presents the exact depth of every channel for every rat. For further explanation of channel choice see Section 3.1.

Table 2.1: Assignment of channels numbers to actual electrode channels for each animal. Channels without any number were discarded.

Depth [mm]	Rat1	Rat2	Rat3	Rat4	Rat5	Rat6
0	1	1	1	1	1	1
0.15			2		2	
0.3	2	2		2		2
0.4			3			
0.5	3	3	4	3	3	3
0.6		4		4		4
0.7	4				4	
0.9		5	5	5	5	5
1.15	5				6	6
1.4		6	6	6	7	
1.6	6					7
1.8	7	7	7	7		

2.4. Stimulation procedure

In the experiments stimuli were presented with interstimulus intervals of 1, 0.5, 0.25, 0.2, 0.143, 0.08 [s] which correspond to frequencies 1, 2, 4, 7, 10, 12 [Hz], respectively. An individual stimulus was a 2-millisecond-long LED flash at 560 cd/m² luminance. It was dark in the room and rats had both eyes opened. For each frequency the stimuli were applied in 40 repetitions of 5-second-long series with random (2-3 s) breaks between them (Fig. 2.2).

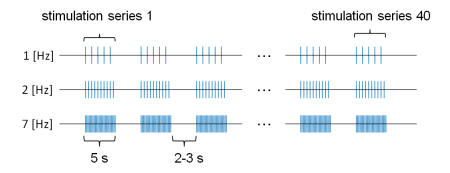


Figure 2.2: Stimulation protocol. The stimuli were presented in 5 s long 40 series at given frequency (from 1 to 12 Hz). The number of stimuli in a series depends on frequency.

Chapter 3

Data analysis

3.1. Data preparation

Signal was filtered by a bandpass filter in range of 0.3–500 Hz with 500 gain using differential AC amplifiers (A-M 140 Systems, US) and recorded with sampling frequency 1000 Hz. For further analysis hardware filter turned out to be insufficient so three other Butterworth filters were used: 1st order lowpass at cut off 100 Hz, 1st order bandstop at cut off between 45 and 55 Hz and 1st order highpass at cut off 0.5 Hz. To avoid change of phase of the signal filtfilt function was used. After filtering each channel of data was normalized by subtracting the mean of all recordings for each channel and divided by their standard deviation. Butterworth filters were used because they are characterized by low deformation of signal giving smooth and monotonic transfer function. It is at cost of low efficiency of filtering, but for this data it was sufficient. After this preparation data was cut into 6-s-long chunks from -0.5 s to 5.5 s where 0 corresponded to the beginning of stimulation and averaged across trials. For further analysis 7 channels out of 12 from contralateral (in respect of stimulation side) VCx were chosen to obtain representative profile: 3 of supragranular, 3 of infragranular and one channel from granular layer (layer IV; Fig 3.1).

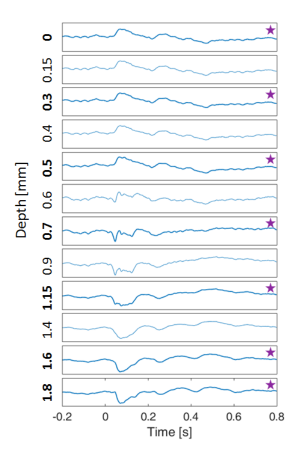


Figure 3.1: Averaged evoked potentials from VCx from Rat1. On channels at 0.6 and 0.7 mm depth repolarization can be observed. Channels chosen to further analysis are marked by purple stars. Time 0 s means an occurrence of stimulus. Y axis the same for all subfigures.

Reducing the number of channels was necessary due to two reasons:

- sometimes short circuits between channels occurred,
- there were differences of depth of electrodes and layer IV was recorded by different channels.

The choice was made basing on analysis on the averaged evoked potentials. The referential depth was established by a channel where repolarization was observed right after the occurrence of the stimulus. This channel was assumed to be recorded from layer IV. Then 3 other channels above the reference and 3 more below were chosen, possibly maximally scattered, but avoiding channels which exhibited any distortion.

3.2. Averaging visual potentials

3.2.1. Description of the method

Visual Evoked Potential (VEP) is a case of steady state evoked potential where light is the stimulus. Theoretically, spontaneous activity of ECoG is a stochastic process (independent, stationary noise with mean equal to zero) and brain response to each stimulus is constant.

Assuming this signal in each realization can be expressed as follows:

$$x_i(t) = s(t) + n_i(t), \tag{3.1}$$

where s(t) is a real signal, $n_i(t)$ noise part. For white noise with mean zero, expected value equals:

$$E\left[\frac{1}{N}\sum_{i=1}^{N}n_i(t)\right] = 0, (3.2)$$

which in turn for averaged signal gives us:

$$E\left[\bar{x}(t)\right] = s(t) \tag{3.3}$$

The effect of averaging is shown in Figure 3.2. Visual response to a single visual stimulus is usually too weak to distinguish it from the background of spontaneous activity of visual cortex. Averaging across several repetitions makes evoked potential stand out.

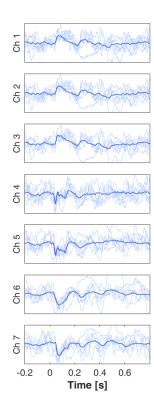


Figure 3.2: Comparison between single trials and their average for Rat1. Lighter blue lines represent few examples of responses to a single repetition of a stimulus. Bold and darker line is the average across all trials. Y axis is the same for all subfigures.

3.2.2. Measurement of peak amplitude in VEP

For stimulation with frequencies 1 and 2 Hz it was possible to observe visual evoked potentials. There were always two deflections from zero: first almost immediately after the stimulus and second about 0.3 s later (Fig. 3.3). Further they are called first and second peak (no matter whether they are negative or positive).

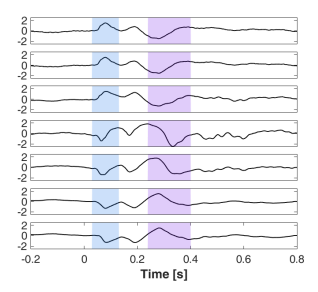


Figure 3.3: Averaged potentials for Rat4 stimulated with 1 Hz. Amplitude of the first peak was calculated for the period marked with blue background (0.03–0.13 s after stimulus). Amplitude of the second peak was calculated for the period marked with purple background (0.24–0.4 s after stimulus).

In Figure 3.4 first channel for the same rat as in Figure 3.3 is presented. Amplitude was taken to be the most extreme value (i.e. having the highest absolute value) in a specified time after an occurrence of a stimulus.

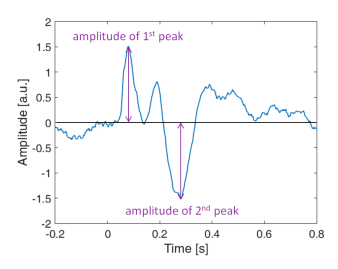


Figure 3.4: Measurement of the peaks amplitude for channel 1 of Rat4 stimulated with 1 Hz.

3.3. Estimation of signal spectrum

Spectra of the signals were estimated by a method introduced by Welch (Welch, 1967). It relies on splitting the signal into overlapping time windows, estimation of the periodogram for each of the windows, and then averaging of the estimates. In this experiment, data was

averaged by 5-second-long chunks for 7 channels and estimation of power spectrum density was calculated. 3-second-long Hamming window with overlap of 2 seconds was used. In Figure 3.5 measurement of the amplitude of fundamental frequency and second harmonic is presented. Amplitudes are the exact values of power spectrum in specific frequencies.

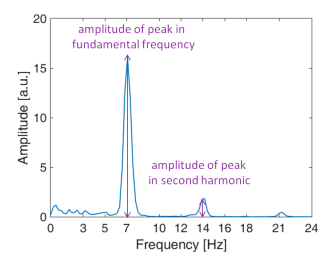


Figure 3.5: Measurement of the peak amplitude for channel 1 of Rat4 stimulated with 7 Hz.

3.4. Current Source Density

To localize synaptic dynamics it is convenient, whenever possible, to estimate the density of transmembrane current sources (CSD) generating the LFP. Let us assume that we record potential in infinitive, homogeneous ans isotropic conductive medium. Current I in that tissue has current density \overrightarrow{J} given by equation:

$$\vec{J} = \frac{I\vec{r}}{4\pi r^2} \tag{3.4}$$

radially at a distance r from the source. In conductive medium with conductivity σ Ohm's law describes relation between current density and potential:

$$\overrightarrow{J} = -\sigma \nabla V. \tag{3.5}$$

Multitude of currents I_j located at $\overrightarrow{r_j}$ induce potential:

$$V(\vec{r}) = \sum_{j} \frac{I_{j}}{4\pi\sigma |\vec{r} - \vec{r'}|}.$$
(3.6)

Introducing current source density equals to:

$$C(\vec{r}) = \sum_{j} I_{j} \delta(\vec{r} - \vec{r'}), \tag{3.7}$$

we can find a relation between potential and CSD:

$$V(\vec{r}) = \frac{1}{4\pi\sigma} \int d^3 \vec{r'} \frac{C(\vec{r'})}{|\vec{r} - \vec{r'}|},$$
(3.8)

where V is potential generated by sources, σ is a conductivity, \overrightarrow{r} is a distance between potential and source. By inverting this relation we get Poisson equation:

$$C(\vec{r}) = -\sigma \Delta V. \tag{3.9}$$

Equation 3.9 can be generalized for arbitrary conductivity tensor fields σ :

$$C(\overrightarrow{r}) = -\nabla \cdot [\sigma \nabla V]. \tag{3.10}$$

In this work to calculate CSD the method introduced by Jan Potworowski called *kernel Current Source Density* (kCSD) was used (Wójcik, 2014; Potworowski et al., 2012).

Chapter 4

Results

In Sections 4.1 and 4.2 all results were presented in the same way. On Figure 4.1 an example of used method is shown on the evoked potentials in channel 4 of Rat1.

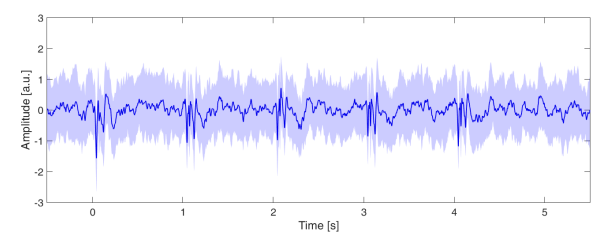


Figure 4.1: Method of presenting results on an example of stimulation of 1 Hz in time domain. Bold line is the average evoked potential for all animals and blue background is the dispersion from the mean by one standard deviation.

4.1. Overview of results obtained in the time domain

Next, there are all experiments with 6 different stimulation frequency presented one after another. In the Figure 4.2 results of stimulation with 1 and 2 Hz frequencies are presented—there is a clear peak of response after each repetition of stimulus.

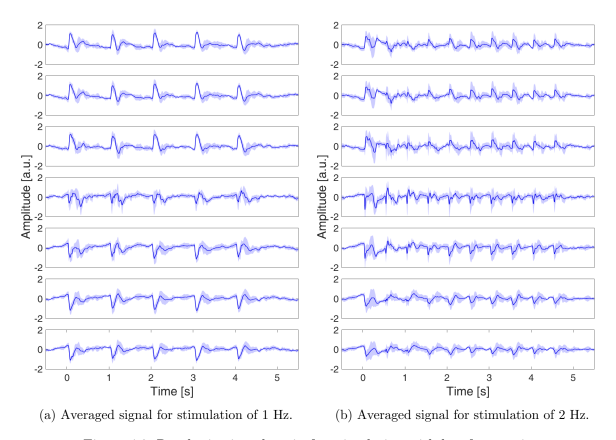


Figure 4.2: Results in time domain for stimulation with low frequencies.

For frequencies higher than 2 Hz signals look different (Fig. 4.3). Firstly the response for stimuli is blurred: stimuli occurs so often that it is impossible to determine where response for certain stimulus starts and ends. The signal forms a continuous oscillation. During the first 1.5 s of the stimulation: the amplitude of oscillations is higher than for the rest. At around 5.15 s there is noticeable peak probably as a response to end of stimulation.

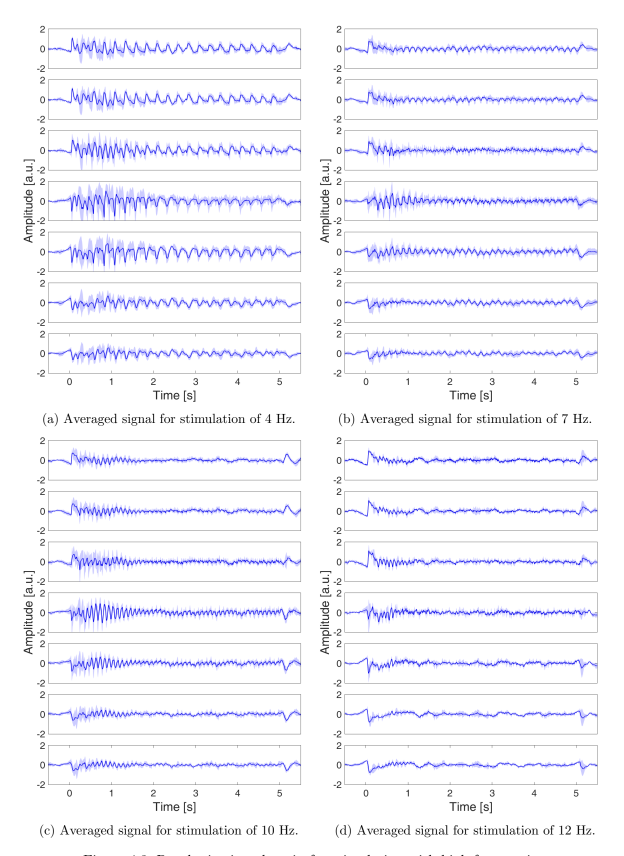


Figure 4.3: Results in time domain for stimulation with high frequencies.

4.2. Overview of results obtained in the frequency domain

Power spectra calculated by Welch's methods are shown in Figures 4.4 and 4.5. For stimulation of 1 Hz there are up to 5 harmonic, and for both 1 and 2 Hz there is so high dispersion that for lowest values there is no difference between peak and zero (Fig. 4.4).

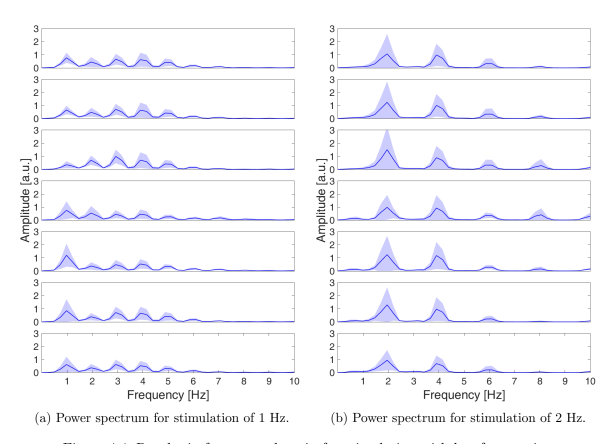


Figure 4.4: Results in frequency domain for stimulation with low frequencies.

For frequencies from range 4-10 Hz, the fundamental frequency and the second harmonic are clean and much bigger than others. Response for 12 Hz in power spectrum is noticeably weaker – peak in the fundamental frequency is at the same level as low frequency noise (Fig. 4.5).

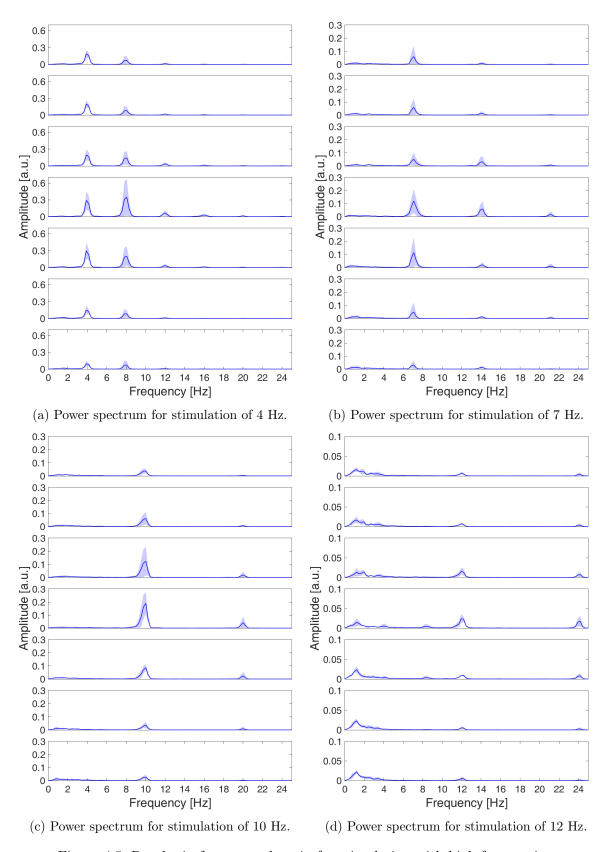


Figure 4.5: Results in frequency domain for stimulation with high frequencies.

4.3. Analysis of current source density

Current source density analysis is shown in Figure 4.6 and 4.7. For stimulation of 1 and 2 Hz there is a change in polarization to each repetition of stimulus (Fig. 4.6).

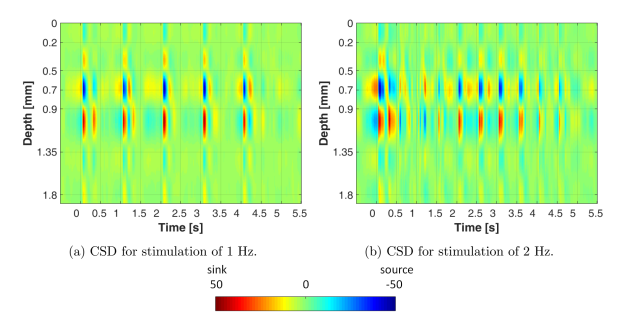


Figure 4.6: Results of CSD for stimulation with low frequencies.

For stimulation of frequency higher than 2 Hz there is a change of polarization just after end of stimulation (around 5.2 s; Fig. 4.7).

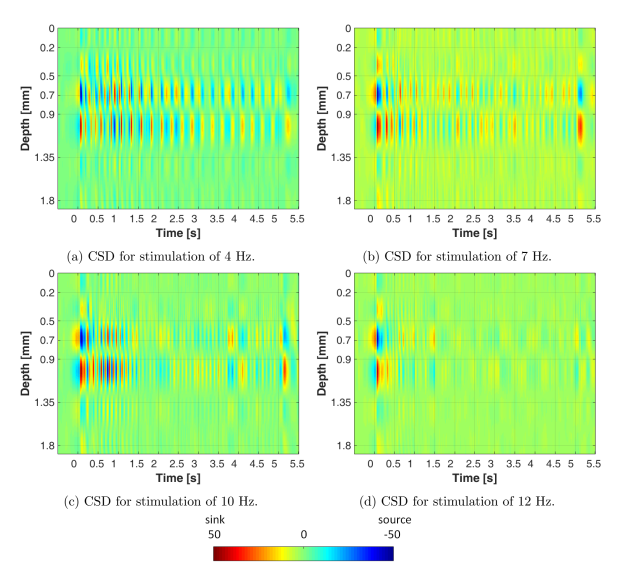


Figure 4.7: Results of CSD for stimulation with low frequencies.

4.4. Analysis of laminar profile for stimulation with 1 Hz

The reverse of potential in layer IV of VCx can be observed (Fig. 4.8). Granular layer (always 4^{th} subfigure) is characterized by the most diverse response to the stimulation within this group of animals. It can be noticed that first peak of response is stable whereas second peak is more varied.

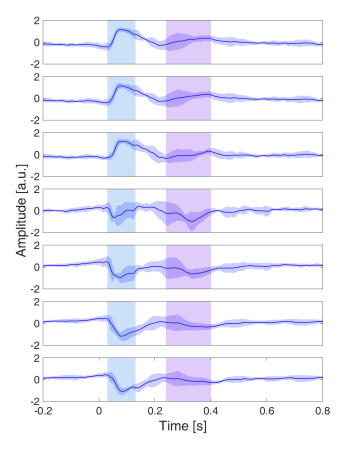
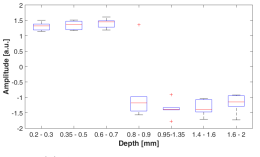
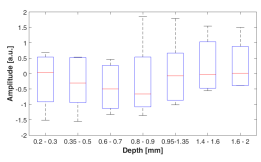


Figure 4.8: Laminar profile in function of depth of the electrode for 1 Hz stimulation. All experiments averaged by 40 trials in time domain – bold line. Differences between animals are by the standard deviation. Time 0 s indicates beginning of 5 s constant stimulation (here only 0.8 s is shown). Both peaks are marked by color backgrounds.

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In the Figure 4.9 laminar profile of amplitude in dependence of depth of the electrode is shown. Amplitude of the first peak is very similar among animals and there is clear change of sign between third and forth channel (Fig. 4.9a). Amplitude of the second peak is characterized by a high variance and there is not difference between layers (Fig. 4.9b).

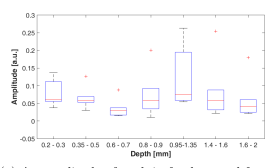


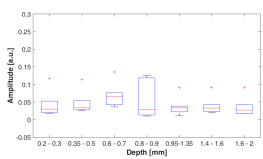


- (a) Amplitude of the first peak.
- (b) Amplitude of the second peak.

Figure 4.9: Amplitude of the averaged signal in function of the depth of the electrode for stimulation of 1 Hz. On the left there is an amplitude of first peak (blue extremum in Fig. ??), on the right – amplitude of second peak (red extremum in Fig. ??). Blue section is a 95% confidence interval, red line is a mediana, red pluses are outliers.

In the Figure 4.10 laminar profile of amplitude of the peak fundamental and frequency and second harmonic in dependence of depth of the electrode is shown. Amplitude of the first peak is the biggest for fifth channel than for the others (Fig. 4.10a). Amplitude of the second peak is constant for all channels except the forth one. This channel also has the highest variance (Fig. 4.10b).





(a) An amplitude of peak in fundamental frequen-(b) An amplitude of peak in the second harmonic.

Figure 4.10: Laminar profile of frequency domain in function of depth of the electrode for 1 Hz stimulation. On the left there is an amplitude of peak in fundamental frequency, on the right – amplitude of second harmonic. Blue section is a 95% confidence interval, red line is a mediana, red pluses are outliers.

4.5. Analysis of laminar profile for stimulation with 2 Hz

In the Figure 4.11 laminar profile of amplitude of response to stimulus in dependence of depth of the electrode for stimulation of 2 Hz is shown. Amplitude of the first peak is very similar among animals except channel 4 where variance is the biggest. (Fig. 4.11a). Tendency

of amplitude of the second peak is like for 1 Hz: it has a high variance and there is not difference between layers (Fig. 4.11b).

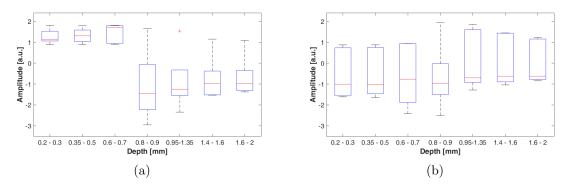


Figure 4.11: Laminar profile in function of depth of the electrode for 2 Hz stimulation.

In the Figure 4.12 laminar profile of amplitude of peak in 2 and 4 Hz in dependence of depth of the electrode for stimulation of 2 Hz is shown. There is no clear difference for amplitude of the first peak. (Fig. 4.12a). Median for second peak is the lowest for forth channel but all channels have the same, high variance (Fig. 4.12b).

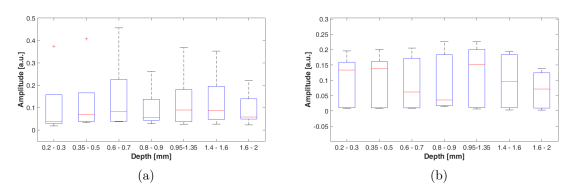


Figure 4.12: Laminar profile in function of depth of the electrode for 2 Hz stimulation.

4.6. Analysis of laminar profile for stimulation with 4 Hz

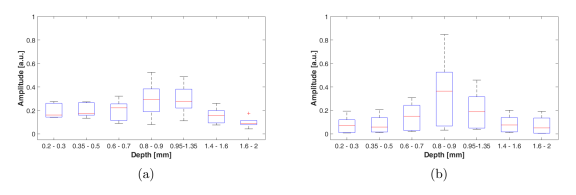


Figure 4.13: Laminar profile in function of depth of the electrode for 4 Hz stimulation.

4.7. Analysis of laminar profile for stimulation with 7 Hz

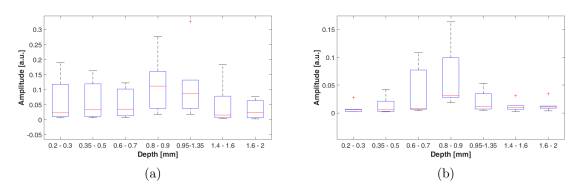


Figure 4.14: Laminar profile in function of depth of the electrode for 7 Hz stimulation.

4.8. Analysis of laminar profile for stimulation with 10 Hz

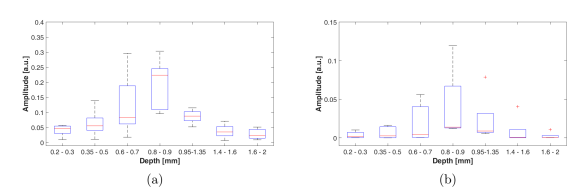


Figure 4.15: Laminar profile in function of depth of the electrode for 10 Hz stimulation.

4.9. Analysis of laminar profile for stimulation with 12 Hz

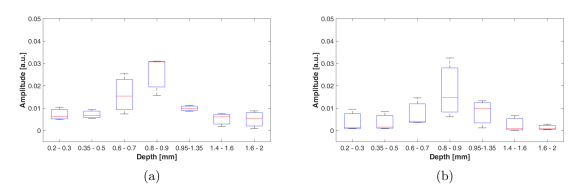


Figure 4.16: Laminar profile in function of depth of the electrode for 12 Hz stimulation.

Chapter 5

Discussion

First of all, it is worth mentioning that even though all recordings were performed on the same structure, using the same electrode, brain anatomy differs between subjects. Individual channels were chosen for analysis based on the pattern of visual potential so there is no guarantee that one channel represents exactly the same structure for every animal. The same concerns the method of recording signal—local field potential is spatially an inaccurate method because it gathers signal from quite a big area of the brain, possibly spanning over multiple structures. On the other hand, this experiment was performed on a group of six animals and the obtained results are coherent and repeatable. Especially the latter confirms that the conclusions are correct.

Analysis of averaged signal for 1 Hz and 2 Hz indicates that there is a clear difference in amplitude of the first peak between supra- and infragranular layers. It can be a result of the fact that magnocellular channel enters from LGN mainly to layer IV of VCx and information from there propagates the further the bigger amplitude gets. For both stimulations there was a high variance of the amplitude of the second peak between animals, but there was no significant difference across the layers. Although the results of averaged potentials are similar for both low-frequency stimulations, there are substantial differences in the peaks for fundamental frequency and second harmonic. For 1 Hz stimulation channel 5 exhibited the highest amplitude and variance of fundamental frequency, while for 2 Hz stimulation channel 4 had the lowest value and variance. The same concerns second harmonic: for 1 Hz stimulation all channels had similar values except channel 4 which stood out having the highest variance. For 2 Hz stimulation the situation was quite opposite—channel 4 exhibited the lowest variance. Those contradictory effects may be a result of improper method of analysis. On the other hand, they could be due to high amplitude of low frequencies which are present in a sleeping animal so that low frequency of stimulation may be not enough to stand out (Wierzbicka et al., 2017).

For stimulation with frequency higher than 2 Hz the amplitude of peak in averaged signal was not calculated because stimulus occurred to often to generate full response and it was not possible to determine the peak. During analysis in the frequency domain the same tendency for all frequencies was observed. It is noticeable already for fundamental frequency, but for second harmonic it is much more apparent: channel 4 is characterized by the biggest amplitude and has lower variance than other channels. It also may be a cause of magnocellular entrance in layer IV in VCx. ...

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