Simultaneous Measurement of Neural Activities of Acute Mouse Hippocampal Slices Using Multi-Electrode Array System and Laser Confocal Calcium Imaging

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Abstract
Recently, non-invasive, real-time and multi-point measurement of neural activities has become possible by using a multi-electrode array (MEA). Another method for multi-point measurement is the fluorescent imaging technique using voltage indicator dyes or calcium indicator dyes. Especially, calcium imaging using fluorescent calcium indicator dyes is often more useful, because they exhibit larger changes in the fluorescence intensity than voltage indicator dyes and their fluorescence changes can be detect easily. Additionally, calcium signals play key roles in the brain function, such as the long-term potentiation (LTP) in the hippocampus, and calcium imaging can be a powerful tool to elucidate the brain function. In this study, we constructed a measurement apparatus combining the MEA system and laser confocal calcium imaging and simultaneously measured electric signals and calcium signals in acute mouse hippocampal slices. The obtained results showed the availability of the present method.

Keywords
Multi-Electrode Array, Laser Confocal Calcium Imaging, Hippocampus, Acute Slice, Long-Term Potentiation

1. Introduction
The brain function is based on complicated interactions among electric neural activities, intracellular calcium signals, intercellular communications by neuro-
transmitters, and so on. In the hippocampus, the long-term potentiation (LTP), a persistent increase in synaptic strength following high-frequency stimulation of a chemical synapse, is a widely accepted cellular model for learning and memory, and an activity-dependent intracellular Ca\(^{2+}\) increase represents a key signal for the activation of the mechanism. At the synapse between the CA3 region and CA1 region, where the neurotransmitter is glutamate, Ca\(^{2+}\) influx through NMDA (N-methyl-D-aspartate) receptors in postsynaptic membrane induces increases in the activity and number of AMAP (\(\alpha\)-amino-3-hydroxy-5-methyl-4-isoxazolepropionicacid) receptors and the quantity of released glutamate following the activation of a series of second messengers [1]. The involvement of calcium signals in the LTP mechanism was shown by using Ca\(^{2+}\) chelator [2] and caged Ca\(^{2+}\) [3], and calcium imaging technique [4] [5] mentioned below.

Electric signals in the brain can be measured by the conventional electrophysiological methods. By intracellular or extracellular recording, direct measurement from a single neuron or neural area is possible using microelectrodes. For the LTP measurement in the hippocampus, for example, electric stimulation such as the tetanic stimulation or theta burst stimulation is applied by a microelectrode inserted around the Shaffer collateral, and field excitatory postsynaptic potential (fEPSP) is extracellularly measured by another microelectrode inserted in the CA1 region.

The electrophysiological methods, however, are not suitable for measurement from many neurons or neural areas. They are also invasive and not suitable for long-term measurement. Recently, non-invasive, real-time and multi-point measurement of neural activities has become possible by using a multi-electrode array (MEA) [6] [7] [8] [9] [10]. The MEA has been also used for the LTP measurement in the hippocampus [11] [12] [13], in which some of the MEA electrodes were used for electric stimulation. Another method for multi-point measurement is the fluorescent imaging technique using voltage indicator dyes [14] [15] [16] [17] or calcium indicator dyes [18] [19]. Especially, calcium imaging using fluorescent calcium indicator dyes is often more useful, because they exhibit larger changes in the fluorescence intensity than voltage indicator dyes and their fluorescence changes can be detected easily. Additionally, calcium signals play key roles in the brain function, such as the LTP in the hippocampus, as mentioned above and calcium imaging can be a powerful tool to elucidate the brain function.

The MEA system [8] [11] [12] [13] and calcium imaging [4] [5] [19] have been individually used for multi-point measurement of neural activities in the hippocampus. In this study, we constructed the combined apparatus and simultaneously measured electric signals and calcium signals in acute mouse hippocampal slices. Here, we used calcium imaging using laser confocal microscopy which enables us to measure the calcium signals with single-neuron resolution [20] [21] [22].
2. Materials and Methods

2.1. Chemicals

Cal-520 AM, a fluorescent calcium indicator dye, was purchased from AATBiocquest (CA, USA). All other chemicals were of the highest purity commercially available.

2.2. Slice Preparations

Male ddY mice (1-week-old) were purchased from Sankyo Labo Service (Tokyo, Japan). They were killed by cervical dislocation followed by decapitation. The brain was rapidly removed and placed into an ice-cold oxygenated artificial cerebrospinal fluid (ACSF) (composition in mM: NaCl 124, KCl 5, CaCl2 2, NaHCO3 22, MgSO4 2, NaH2PO4 1.24, glucose 10, pH 7.4, bubbled with 95% O2/5% CO2). Transverse hippocampal slices (thickness 350 µm) were then prepared using a microslicer (DTK-1000, Dosaka-EM, Kyoto, Japan) in an ice-cold oxygenated sucrose solution (composition in mM: KCl 2.6, CaCl2 0.5, NaHCO3 27, MgSO4 7, NaH2PO4 1.5, ascorbic acid 0.1, sucrose 222, pH 7.4, bubbled with 95% O2/5% CO2) which can prevent cell damages [21]. The slices were recovered in ACSF at 30˚C for 60 min and held at room temperature.

The care of animals and experimental procedures were carried out in accordance with national and institutional guidelines, and all experimental protocols were approved by Nihon University Animal Care and Use Committee.

2.3. Laser Confocal Calcium Imaging

The prepared hippocampal slices were stained with Cal-520. The slices were loaded for 60 min at room temperature with about 4 µM of the dye in the presence of 0.005% Cremophor EL and 0.01% Pluronic F-127 in 4.0 mL ACSF. After dye loading, the slices were washed in ACSF for 30 min. The stained slices were placed on a planar glass MEA chip mentioned below, and then the chip was placed on the stage of an upright microscope (E600FN, Nikon, Tokyo, Japan) and perfused with a modified ACSF (composition in mM: NaCl 124, KCl 5, CaCl2 2, NaHCO3 22, MgSO4 2, NaH2PO4 1.24, glucose 10, pH 7.4, bubbled with 95% O2/5% CO2) kept at 30˚C. They were illuminated through 10% ND filter by an Ar laser (488 nm; 532-BS-AO4, Melles Griot, NM, USA). The laser power was set to 15 mW. Then, the 520 nm fluorescence images were acquired through a 16x fluorescence objective (0.75 NA, S Fluor, Nikon, Tokyo, Japan), a Nipkow confocal unit (CSU-10, Yokogawa, Tokyo, Japan) and an EM-CCD camera (iXonX3 897, Andor, Belfast, UK). In each acquisition trial, consecutive fluorescence images were acquired at a rate of 150 ms/frame on a personal computer (PRECISION T3500, DELL). The obtained fluorescence images were analyzed with Andor Solis (Ver. 4.18, Andor, Belfast, UK).

2.4. Multi-Electrode Array Recording

The planar glass MEA chip used comprised 64 electrodes (50 × 50 µm) with 150
µm spacing in 8 × 8 grid arrangement (MED-P515A, Alpha MED Scientific, Japan). The extracellular electric signals detected through the electrodes were amplified by 64-channel main amplifier (MED-D64A32, Alpha MED Science, Japan) and 64-channel head amplifier (MED-A64HE1S, Alpha MED Science, Japan) and acquired on a personal computer (PRECISION T1700, DELL) at a rate of 20 kHz/channel. In some experiments, the electrodes were used for electric stimulation which was applied for the LTP measurement. A schematic illustration of the present apparatus is shown in Figure 1.

2.5. Data Analysis

We obtained raster plots from the peaks of the measured calcium signals as follows. From each pixel of the camera, the photocount was acquired as the fluorescence intensity (F). The change in F from the baseline was calculated as ΔF/F₀ = (F − F₀)/F₀. Here, F₀ is the baseline fluorescence intensity. We next detected cell bodies in the fluorescence images by their profiles, and regions of interest (ROIs) were put on the individual cell bodies. The fluorescence intensity changes, ΔF/F₀, of the pixels in each ROI were averaged as the calcium signals. Figure 2(a) shows a time course of the calcium signals. The peaks were detected using a custom program for MATLAB (Mathworks, Natick, NY, USA).

We also obtained raster plots from the peaks of the measured electric signals. Figure 2(b) shows a time course of the electric signals. The upwards and downwards peaks were also detected using a custom program for MATLAB (Mathworks, Natick, NY, USA).

3. Results and Discussion

We combined the MEA system and laser confocal calcium imaging (Figure 1) and simultaneously measured extracellular electric signals and intracellular
Figure 2. Time courses of (a) the calcium signals and (b) electric signals. The blue and black dots are detected peaks of the calcium signals and electric signals, respectively.

calcium signals in acute mouse hippocampal slices. As the calcium indicator dye, we used Cal-520 which is recently developed and exhibits a large fluorescence change [23] [24]. Although we used an upright microscope in the present study, an inverted microscope could be suitable for another study. Especially in such a case, we must detect the calcium signals through the glass MEA chip and therefore Cal-520 will be preferable. In our previous experiments, the average peak amplitude of calcium signals and SNR of Cal-520 were estimated to be about 11.5% and 3.4, respectively, while those of Oregon green which has been much often used for laser confocal calcium imaging [21] [22] were estimated to be 6.5% and 2.0 (not published data).

We first measured spontaneous electric signals and calcium signals simultaneously using the present method. Here, no stimuli were applied anywhere. Sometimes such spontaneous neural activities occurred without any stimuli. The electric signals were measured in the whole of hippocampus (Figure 3(a) (upper panel) and Figure 3(c)) and the calcium signals were measured in the CA1 region (Figure 3(a) (lower panel) and Figure 3(b)) which is shown with blue rectangles in Figure 3(a) (upper panel) and Figure 3(c). The supplemental movie can be seen in the web site (The URL is in the figure legend of Figure 3). Figure 4 shows the raster plots for the electric signals and calcium signals. The peak positions of signals are shown by black or blue dots. The electric signals in the
Figure 3. (a) Upper panel: A hippocampal slice on the MEA chip. The small black squares are the 64 electrodes. Lower panel: Fluorescence image of the CA1 region for laser confocal calcium imaging (b), which is shown as a blue rectangle in the upper panel and (c). (b) Consecutive images of $\Delta F/F_0$ in the CA1 region. The images are shown with a gray scale. The time proceeds from the upper left to lower right image at an interval of 150 ms. (c) Consecutive electric signals from 64 MEA electrodes. 64 windows of each panel correspond to the MEA electrodes. The full scales of the horizontal and vertical axes of each window are 1 s and 0.1 $\mu$V, respectively. The time proceeds from the upper left to lower right panel at an interval of 150 ms. The blue rectangle in each panel corresponds to each image of (b), and the image was acquired at the center time (dotted line) of each window. See also the supplemental movie (http://saitolab-chaos.com/Papers/jbbs2017/Supplemental-movie1.html; double speed reproduction).

Figure 4. Raster plots for the electric signals from 64 MEA electrodes under the DG, CA3 and CA1 regions and the calcium signals from 135 ROIs in the CA1 region. The black and blue dots show the peak positions of electric signals and calcium signals, respectively. The slice is the same one used for Figure 3.
whole of hippocampus, where larger burst-like signals were seen in the CA3 region, and the calcium signals of the CA1 region seem to occur simultaneously. Expanding the time scale, however, the calcium signals occur with a time delay of a few hundred of milliseconds from the electric signals (Figure 5). On the other hand, no distinct time delay was detected among the electric signals of those regions and therefore it was unclear where the spontaneous neural activities originated.

We next measured the electric signals and calcium signals during the LTP measurement. The LTP was induced by applying the theta burst stimulation to an electrode under the Shaffer collateral. In the theta burst stimulation, the burst of 4 pairs of pulses (each pulse strength: −100 μA or +100 μA, each pulse width: 0.1 ms) with 10 ms interval was repeated 10 times with 200 ms interval, and the train of 10 bursts was repeated 20 times with 5 s interval. The electric signals were measured for the whole of hippocampus, and the LTP induction was confirmed by an increase in the slope of fEPSP in the CA1 region for the test stimulus (a pair of pulses (each pulse strength: −100 μA or +100 μA, each pulse width: 0.1 ms)) repeated with 20 s interval to the Shaffer collateral (Figure 6), where the calcium signals were measured simultaneously. Figure 7 shows consecutive images of calcium signals in the CA1 region during the LTP measurement. The supplemental movies can be seen in the web site (The URL is in the figure legend)

Figure 5. A time relation between the electric signals and the peak positions (blue dots) of calcium signals. The electric signals were measured by one of electrodes under the DG, CA3 or CA1 region. The slice is the same one used for Figure 3.
Figure 6. (a) fEPSPs before and after the theta burst stimulation; (b) Change in the fEPSP slope by the theta burst stimulation. The slope is normalized by that before the stimulation.

Figure 7. Consecutive images of $\Delta F/F_0$ in the CA1 region during the LTP measurement. The images are shown with a gray scale. The time from the onset of the theta burst stimulation is shown in each image. The yellow circle shows the timing for application of the test stimulus, and the red circle shows the timing for application of a burst train in the theta burst stimulus. No circle shows the timing without any stimuli. The slice is the same one used for Figure 6. See also the supplemental movie (http://saitolab-chaos.com/Papers/jbbs2017/Supplemental-movie2.html; double speed reproduction).
of Figure 7. Figure 8 shows the raster plots of the calcium signals. In this slice, obvious spontaneous neural activities could not be seen; the average peak number of calcium signals was 0.82/min per one ROI before the theta burst stimulation. The test stimuli did not also induce more calcium signals. However, as expected, the calcium signals remarkably increased during the theta burst stimulation; the average peak number of calcium signals was 7.82/min per one ROI. After the theta burst stimulation, it became 0.83/min per one ROI, which did not much increase compared with that before the stimulation.

4. Conclusions

In this study, we constructed a measurement apparatus combining the MEA system and laser confocal calcium imaging. For the calcium indicator dye, we used Cal-520 which is recently developed and exhibits a large fluorescence change. By the present method, we simultaneously measured extracellular electric signals and intracellular calcium signals in acute mouse hippocampal slices. We showed the availability of the present method by obtaining the following results:

1) The calcium signals in the CA1 region occur with a time delay of a few hundred of milliseconds from the electric signals in the DG, CA3 and CA1 regions, while no distinct time delay was detected among the electric signals of those regions.
2) During the LTP induction by applying the theta burst stimulation to the Shaffer collateral, the calcium signals in the CA1 region remarkably increased as expected, while the calcium signals after the theta burst stimulation did not much increase compared with those before the stimulation.

![Figure 8](image)

**Figure 8.** Raster plots for the calcium signals from 112 ROIs in the CA1 region during the LTP measurement. The tick marks on the horizontal axis show the timings for the test stimuli. The blue dots show the peak positions of calcium signals. The slice is the same one used for Figure 6.
To elucidate the brain function, we need to measure complicated interactions among electric neural activities, intracellular calcium signals, intercellular communications by neurotransmitters, and so on. The present method will be the first step to realize it, and we hope that the present method will contribute to the future studies on the brain function including the LTP mechanism in the hippocampus.

**Acknowledgements**

This study was supported by Nihon University Research grant (Total Research 14-002, 15-002 to M.S.).

**References**


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