# multichannel\* systems

MEA Application Note: Primary Neurons from Cortex or Striatum



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A major part of this information is based on the instructions provided by the laboratory of Dr. Steve Potter and Dr. Daniel Wagenaar. We also thank Wiebke Fleischer for contributing demo data.

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# 1 Introduction

### **1.1 About this Application Note**

The intention of the MEA Application Notes is to show users how to set up real experiments with the MEA-System on the basis of typical applications that are used worldwide.

The documents have been written by or with the support of experienced MEA users who like to share their experience with new users.

This application note includes a complete protocol for the isolation and cultivation of primary cortical neurons, suggestions for long term cultures, suggestions for MEA-System configurations, and references.

### 1.2 Acknowledgement

Multi Channel Systems would like to thank all MEA users who shared their experience and knowledge with us, especially the following persons.

Daniel Wagenaar

**Steve Potter** 

Wiebke Fleischer

## 2 Material

### 2.1 **Biological Materials**

• Rat embryos E18 (Wistar Kyoto or Sprague Dawley)

### 2.2 Technical Equipment

- MEA-System (with amplifier and data acquisition, please see chapter "Suggested MEA-System")
- Stimulus generator (for USB-MEA-Systems, but integrated in MEA2100-Systems)
- MEAs (microelectrode arrays)
- Sterile workbench
- Incubator set to 35 °C, 65 % relative humidity, 9 % O<sub>2</sub>, 5 % CO<sub>2</sub>
- Water bath at 37 °C
- Stereo microscope
- Inverted microscope
- Micropipettes and pipette tips (20 µl and 1000 µl)
- 15 ml BD Falcon tubes
- 40 µm nylon mesh cell strainer (BD Falcon)
- Sharp forceps
- Curved forceps
- Small scissors
- Teflon membranes (ALA Scientific Instruments Inc.)

### 2.3 Chemicals

- NaOH
- MgCl<sub>2</sub>
- CaCl,
- HEPES
- Phenol Red
- Na<sub>2</sub>SO<sub>4</sub>
- K<sub>2</sub>SO<sub>4</sub>
- Kynurenic acid
- DL-2-amino-5-phosphonovaleric acid (APV)
- Polyethylenimine (PEI)
- Laminin
- Sodium pyruvate
- Insulin
- Glutamate
- Phosphate buffered saline (PBS)
- Bovine serum albumin (BSA)
- Trypsin (Sigma)
- Papain suspension (Roche Applied Science, catalog No. 10108014001)
- DNAse (Sigma)
- Horse serum (Donor Equine Serum from HyClone)
- Dulbecco's Modified Eagle Medium (DMEM) (Gibco/Invitrogen)
- L-Alanyl-L-Glutamine (GlutaMAX from Gibco/Invitrogen)
- Hanks' Balanced Salt Solution (HBSS) without Calcium / Magnesium (Gibco/Invitrogen)

#### 2.4 Media

#### 2.4.1 Culture Medium for Primary Cultures

- DMEM (may contain GlutaMax, depending on the supplier)
- 10 % horse serum
- 0.5 mM GlutaMax (final concentration)
- 1 mM sodium pyruvate
- 2.5 µg/ml insulin

Note: Some work groups use trypsin, other papain for the enzymatic digestion of the tissue.

#### 2.4.2 Enzyme Solution (Trypsin)

- HBSS
- 0.25 % trypsine
- 0.02 mg/ml DNAse

#### 2.4.3 Alternative Enzyme Solution (Papain)

- 2 ml Segal's medium (see below)
- 200 µl papain suspension
- NaOH for adjusting the pH to 7.3

#### 2.4.4 Segal's medium

(Banker & Goslin, p309ff.)

	Conc. (mM)	FW (g/mol)	mg (for 500 ml)
MgCl <sub>2</sub> ·6H <sub>2</sub> O	5.8	203.31	590
CaCl <sub>2</sub> ·6H <sub>2</sub> O	0.25	147.02	18.4
HEPES	1.6	238.3	191
Phenol Red	0.001 %		5
Na <sub>2</sub> SO <sub>4</sub> ·10H <sub>2</sub> O	90	322.21	14500
K <sub>2</sub> SO <sub>4</sub>	30	174.26	2610
Kynurenic acid	1	189.2	95.6
APV	0.05	197.1	4.92

Note: Kynurenic acid takes a **lot** of stirring to dissolve. It is also recommended to add more NaOH while dissolving to keep the pH reasonable.

- 1. Use 0.1N NaOH (about 1 ml) for adjusting the pH to 7.3 before adding APV and Kynurenic acid.
- 2. Again, use 0.1N NaOH (about 1 ml) for adjusting the pH to 7.3 after having added APV and Kynurenic acid.
- 3. Bring up to 500 ml after the final pH adjustment.
- 4. Sterile filter, aliquot, and freeze in liquid nitrogen.

#### 2.4.5 BSA / PBS

- Phosphate buffered saline (PBS)
- Bovine serum albumine (BSA)

### 3 Methods

#### 3.1 MEA Coating

Depending on the type of selected MEA, various coatings may be applied. Standard MEAs should be coated with Polyethylenimine or Laminin. Suggestions for coating methods can be found in the MEA manual available in the "Download section" of the MCS web site.

#### 3.2 Dissection

- 1. Euthanize the animal with carbon dioxide.
- 1. Decapitate the animal with large sharp scissors or with a guillotine.
- 2. Open the skull carefully with a scissor and remove the brain.
- 3. Isolate the cortex (or dissect the striata) with a scalpel.

### 3.3 Enzymatic Digestion

- 1. Chop the isolated sample into pieces of approximately 1 mm<sup>3</sup> and transfer the fragments into a Falcon tube with 2 3 ml enzyme solution pre-warmed at 37 °C.
- 2. Digest the fragments with the enzyme solution at 37 °C for 15 min. Gently swirl the suspension every 5 min.
- 3. Gently wash the fragments in the culture medium twice: Gently swirl or invert the suspension of brain pieces in medium to wash away the protease, allow the pieces to settle, remove the supernatant with a pipette, and add 2 3 ml fresh medium. After the second wash, add 1 ml medium to the fragments.
- 4. Gently triturate the fragments by passing the preparation five times through the 0.78 mm wide opening of a 1000  $\mu$ l pipette tip. The majority of cells should now be in suspension.
- 5. Transfer the supernatant containing the suspended cells into a fresh Falcon tube.
- 6. Add 1 ml medium to the remaining fragments, and triturate the remaining fragments once more.
- 7. Combine the supernatants from the two triturations in one tube, giving 2 ml cell suspension.
- 8. Remove the debris by gravity flow filtering the cell suspension through a 40 μm nylon mesh cell strainer (Falcon) into a 15 ml DB Falcon tube filled with BSA/PBS.

### 3.4 Plating und Culturing the Cells

- 1. Centrifuge the cell suspension at 160 g for 5 min.
- 2. Discard the supernatant and resuspend the cells in approximately 0.5 ml culture medium.
- 3. Count the cells under a microscope to determine the cell density by using a Neubauer chamber or an automated cell counter.
- 4. Plate the cells in a density of 1000 5000 cells per mm<sup>2</sup> (depending on your application) onto the recording field of the MEA.
- 5. Maintain the cells for 4 5 days in an incubator set to 35 °C, 65 % relative humidity, 9 %  $O_2$ , 5 %  $CO_2$  before recording, depending on your application.
- 6. Check the pH of medium daily by eye, and change it as soon as the color indicates that the medium is going acidic (shifting from pink to orange color).

Note: The choice of plating density is really up to the investigator. The denser, the sooner the activity will be observed (as soon as 4 days *in vitro*), but the more often the culture will need to be fed. 5000 cells/mm<sup>2</sup> is very dense, and would require feeding about every 2 days, while 1000 – 3000 may only require feeding weekly or every 5 days. The best time for recording depends on the application: Many studies may be aimed at the development of the cultures, and therefore require recording as soon as possible. For other applications, it may be better to keep the cells longer in culture before starting the experiment. Cells are still developing, but the culture is more stable after about one month in culture. The culture can be used for several months or years. Application Example: Drug Testing (NMDA)

Data was recorded from a neuronal cell culture with striatum cells from rat (Wistar, E15). This example shows the application of N-methyl-D-aspartate (NMDA).



The MC\_Rack **Analyzer** extracts the spike number in 100 ms bins: Under control conditions, over 60 % of these bins are empty, that is, without spike activity. This demonstrates the organization of the spontaneous spiking activity in bursts. After NMDA application, the spiking activity is more random, demonstrated by the fact that only 8 % of the bins are empty. The overall spiking activity is increased, too, but the main difference is how the activity is organized.

(Data kindly provided by W. Fleischer, University of Düsseldorf)

#### **Control experiment without drug**

This screen shot demonstrates the raw data recording and spike analysis. The window on the lower left shows the spike detection. Spike waveforms are shown in green. On the lower right, the data is analyzed in 100 ms bins and plotted as a false color raster plot. Low or no activity (with respect to the 100 ms bins) is plotted in cyan; highest activity is plotted in magenta, with the color gradient following the activity level. You see that most of the bins are empty (cyan color), because the spike activity is organized in bursts rather than randomly distributed over the bins. This is also reflected by the spike raster plot on the top right, which plots the time points of spike detection events as rows of green vertical bars. The length of each row is 1 s. Most display space remains white, with a regular pattern of green bars, that is, the activity is highly organized in bursts.

#### 5 µM NMDA application

This screen shot shows signals from the same cells, but after the drug application. You see that a majority of electrodes record a more overall, randomly distributed spike activity with only few empty bins. The raster plot shows no regular pattern anymore.



### 4 Longterm Culturing



In order to allow long term cultivation and recording, Multi Channel Systems MCS GmbH recommends the use of teflon membranes (fluorinated ethylene-propylene, 12.5 microns thick) developed by Potter and DeMarse (2001). The ALA-MEA-MEM membrane is produced in license by ALA Scientific Instruments Inc., and distributed via the world-wide network of MCS distributors.

The sealed MEA culture chamber with transparent semipermeable membrane is suitable for all MEAs with glass ring. A hydrophobic semipermeable membrane from Dupont that is selectively permeable to gases ( $O_2$ ,  $CO_2$ ), but not to fluid, keeps your culture clean and sterile, preventing contaminations by airborne pathogens. It also greatly reduces evaporation and thus prevents a dry-out of the culture.

# 5 Suggested MEA-System

#### 5.1 System Configurations

Depending on the throughput and the analysis requirements desired in your laboratory, different system configurations are recommended for the recording from cultured neurons.

- **MEA2100-System**: For the MEA2100-HS60(2x60)-System, the same MEAs can be used as for the MEA1060 amplifiers, with the same advantages and disadvantages. In addition, for the MEA2100-HS120 a 120-channel MEA layout is available (120MEA200/30iR), which provides a larger electrode field with a resolution identical to the 60MEA200/30iR. This might be interesting for a mapping applications. The MEA2100-System allow to operate a maximum of two headstages with up to two 60 channel MEAs each. If all those slots are used with 6 well MEAs 24 wells can be tested simultaneously.
- MEA60-System: 60-channel MEA recording system for inverted or upright microscopes. The temperature controller TC02 regulates the temperature of the MEA and of the perfusion fluid via the perfusion cannula PH01. The peristaltic perfusion system PPS2 is provided respectively. A MEA1060 amplifier allows recording up to 60 channels from a 60-channel MEA. This is the standard configuration for low-throughput academic research.

#### 5.2 Microelectrode Arrays

Available MEAs differ in electrode material, diameter, and spacing. For an overview on available MEA types please see the MEA Manual, or the Multi Channel Systems MCS GmbH web site or contact your local retailer.

The microfold structures formed by titanium nitride (TiN) result in a large surface area that allows the design of small electrodes with a low impedance and an excellent signal to noise ratio. For recording from cultured neurons, a medium spatial resolution with an electrode diameter of 30  $\mu$ m and a spacing of 200  $\mu$ m is generally sufficient.

Recommended MEAs include:

- **60MEA200/30iR-Ti** standard 8 x 8 layout, TiN electrodes for recording and stimulation, with substrate-integrated reference electrode.
- **60ThinMEA** for high-resolution imaging and combination with intracellular calcium measurements. ThinMEAs are only 180 µm "thick" and mounted on a robust ceramic carrier. Tracks and contact pads are made of transparent ITO.

#### **Ring Options**

The standard glass ring is best for running a perfusion, and will also allow using the ALA MEA-Insert. A plastic ring should only be used if parallel patch clamp experiments are planned.

### 6 References

- F.J. Arnold, F. Hofmann, C.P. Bengtson, M. Wittmann, P. Vanhoutte, H. Bading, Microelectrode array recordings of cultured hippocampal networks reveal a simple model for transcription and protein synthesis-dependent plasticity, J Physiol 564 (2005) 3-19.
- Hulata, E., R. Segev, et al. (2000). "Detection and sorting of neural spikes using wavelet packets." <u>Phys Rev Lett</u> 85(21): 4637-40.
- Hardingham, G. E., F. J. Arnold, et al. (2001). "A calcium microdomain near NMDA receptors: on switch for ERK-dependent synapse-to-nucleus communication." <u>Nat Neurosci</u> 4(6): 565-6.
- Potter, S. M. (2001). "Distributed processing in cultured neuronal networks." <u>Prog Brain Res</u> 130: 49-62.
- Potter, S. M. and T. B. DeMarse (2001). "A new approach to neural cell culture for long-term studies." J Neurosci Methods **110**(1-2): 17-24.
- Segev, R., Y. Shapira, et al. (2001). "Observations and modeling of synchronized bursting in two-dimensional neural networks." Phys Rev E Stat Nonlin Soft Matter Phys **64**(1 Pt 1): 011920.
- Streit, J., A. Tscherter, et al. (2001). "The generation of rhythmic activity in dissociated cultures of rat spinal cord." <u>Eur J Neurosci</u> **14**(2): 191-202.
- Marom, S. and G. Shahaf (2002). "Development, learning and memory in large random networks of cortical neurons: lessons beyond anatomy." <u>O Rev Biophys</u> **35**(1): 63-87.
- Segev, R., M. Benveniste, et al. (2002). "Long term behavior of lithographically prepared in vitro neuronal networks." <u>Phys Rev Lett</u> **88**(11): 118102.
- Wagenaar, D. A. and S. M. Potter (2002). "Real-time multi-channel stimulus artifact suppression by local curve fitting." <u>J Neurosci Methods</u> **120**(2): 113-20.
- 0Eytan, D., N. Brenner, et al. (2003). "Selective adaptation in networks of cortical neurons." J Neurosci 23(28): 9349-56.
- Jimbo, Y., N. Kasai, et al. (2003). "A system for MEA-based multisite stimulation." <u>IEEE Trans</u> <u>Biomed Eng</u> **50**(2): 241-8.
- Otto, F., P. Gortz, et al. (2003). "Cryopreserved rat cortical cells develop functional neuronal networks on microelectrode arrays." <u>J Neurosci Methods</u> **128**(1-2): 173-81.
- Segev, R., M. Benveniste, et al. (2003). "Formation of electrically active clusterized neural networks." <u>Phys Rev Lett</u> **90**(16): 168101.
- Baruchi, I. and E. Ben-Jacob (2004). "Functional holography of recorded neuronal networks activity." <u>Neuroinformatics</u> **2**(3): 333-52.
- Gortz, P., W. Fleischer, et al. (2004). "Neuronal network properties of human teratocarcinoma cell line-derived neurons." <u>Brain Res</u> **1018**(1): 18-25.
- Gortz, P., A. Hoinkes, et al. (2004). "Implications for hyperhomocysteinemia: not homocysteine but its oxidized forms strongly inhibit neuronal network activity." J Neurol Sci **218**(1-2): 109-14.
- Hulata, E., I. Baruchi, et al. (2004). "Self-regulated complexity in cultured neuronal networks." <u>Phys Rev Lett</u> 92(19): 198105.
- Segev, R., I. Baruchi, et al. (2004). "Hidden neuronal correlations in cultured networks." <u>Phys Rev</u> Lett **92**(11): 118102.
- Wagenaar, D. A., J. Pine, et al. (2004). "Effective parameters for stimulation of dissociated cultures using multi-electrode arrays." J Neurosci Methods **138**(1-2): 27-37.
- Wagenaar, D. A., Madhavan, R., Pine, J. and Potter, S. M. (2005). "Controlling bursting in cortical cultures with closed-loop multi-electrode stimulation." J Neurosci 25(3): 680-8.
- Evans MS, Collings MA, Brewer GJ (1998). Electrophysiology of embryonic, adult and aged rat hippocampal neurons in serum-free culture. Journal of Neuroscience Methods 79:37-46.