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Preparing primary sandwich hippocampal neuron cultures for cryo-electron tomography

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Abstract

Primary sandwich hippocampal neuron cultures are adapted from the Kaech and Banker protocol (Kaech and Banker, 2006) and provide neuronal cultures with almost no glial cells, which may facilitate the targeting of neurons for cryoelectron tomography (see accompanying protocol by Siegert, Petrovic, Do et al.). In addition, the cells can be seeded at a very low density.

Attachments

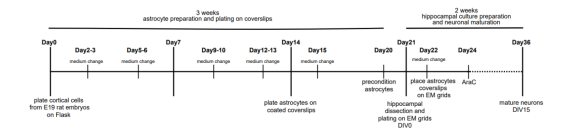


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Guidelines





Materials

Astrocyte feeder layer

- T75 flasks
- 10 mL serological pipettes
- 15 mL sterile tubes
- 25 mm sterile coverslips

A	В
MEM 1X (Gibco) supplemented with	
L-Glutamine	2 mM
Horse Serum (Gibco)	10 %
Glucose (MEM10%H S)	4.65 g/L

- 0.1 mg/mL poly-L-lysine;
- Sterile water;
- 0.05 % Trypsin/0.02 % EDTA
- DMSO
- Centrifuge for 15 mL tubes.

Primary neuron cultures with astrocyte feeder layer

Sterile tweezers

A	В
Neurobasal medium supplemented with	
B27	2 %
L-Glutamine	2 mM

- DMEM10%FCS
- Arabinofuranoside (cytosine-β-arabinofuranoside hydrochloride)



Preparation of the EM grids

1 For preparation of EM grids, follow the accompanying protocol by Siegert, Petrovic, Do et al.

Note

Only 35 mm dishes with 4 inner rings are suitable for the preparation of primary sandwich hippocampal neuron cultures.

Preparation of primary sandwich hippocampal neuron culture on EM grids - Astrocyte feeder layer

- 2 Plate 5 million cortical cells from an E19 rat embryo cortical suspension in T75 flask in MEM10%HS.
- Change the medium of the astrocytes twice a week with fresh MEM10%HS preequilibrated at and 5 % CO₂. Slam the flask against the bench surface before aspirating the medium to remove the microglial cells. Two weeks after the dissection, the astrocyte cultures should be more than 50 % confluent.

Note

The choice of a specific medium as well as the removal of microglia (as described in the previous step) favours the proliferation of astrocytes over neurons and other glia in the cortical cell suspension.

4 6-7 days before the hippocampal dissection (2 weeks after the cortical dissection; Fig.1), coat 25 mm sterile coverslips with μ1 0.1 mg/mL poly-L-lysine for 00:15:00 at

15m

Room temperature



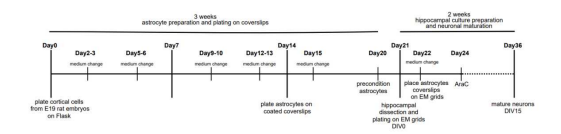


Fig.1: Timeline of primary sandwich hippocampal neuron culture preparation. Astrocytes are cultured for 3 weeks prior to plating on coverslips. Following the hippocampal dissection, primary hippocampal cultures are plated on EM grids, and neuronal maturation takes place in the presence of the astrocyte coverslips and arabinofuranoside.

- 4.1 Wash the coverslips 3 times with sterile water and replace the water with MEM10%HS.
- 5 Harvest the astrocytes from the flask: Slam the flask and aspirate the medium.
- 5.1 Quickly wash the flask with 0.05 % Trypsin/0.02 % EDTA.
- 5.2 Add <u>Add</u> 2 mL of trypsin/EDTA and incubate 000:02:00 at 37 °C until the cells detach.
- 5.3 Stop the trypsination by adding \perp 5 mL of MEM10%HS.
- 5.4 Release the cells by multiple rounds of pipetting with a 10 mL serological pipette.
- 5.5 Transfer the cells to a 15 mL sterile tube and centrifuge for \bigcirc 00:05:00 at \bigcirc 500 x g .
- 5.6 Resuspend the cells in 4 2 mL of MEM10%HS complete medium.

2m



5.7 Count the cells and plate them dropwise on each of the coverslips: use 200,000 cells in a total volume of 4 100 µL of medium per coverslip.



Note

The remaining astrocytes in MEM10%HS complete medium with 10 % DMSO can be frozen and kept in liquid nitrogen storage for more than a year. They can be thawed six days before the hippocampal dissection.

One day after plating, change the medium of the astrocytes again with fresh preequilibrated MEM10%HS.

Preparation of primary sandwich hippocampal neuron culture on EM grids - Primary neuron cultures with astrocyte feeder layer

- One day before the hippocampal dissection, precondition the astrocyte feeder layer in Neurobasal medium + 2 % B27 + [M] 2 millimolar (mM) L-Glutamine pre-incubated at 37 °C and 5 % CO₂ for few hours.
- 8 Dilute the hippocampal suspension to reach a density of 200,000 to 350,000 cells per mL in DMFM10%FCS.
- 9 Immediately after removing the water from the compartments of the dish plate Δ 100 μL of cell suspension dropwise on each EM grid (Fig. 2, left). Incubate at 37 °C and 5 % CO₂.





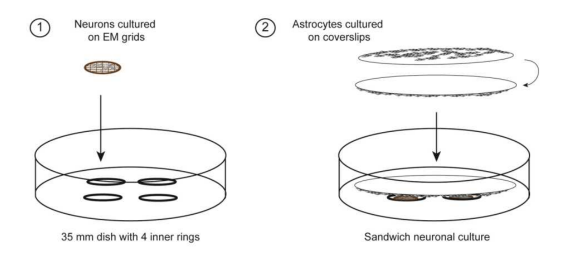


Fig 2: Schema of the sandwich cultures on EM grids

- After two hours, add 500 μL of pre-heated DMEM10%FCS per dish.
 Incubate Overnight at 37 °C and 5 % CO₂.
 Replace the plating medium with 2 mL of pre-conditioned Neurobasal medium from the astrocyte feeder layers' dish.
- With sterile tweezers take the coverslip with the astrocyte layer and flip the coverslip so that the neurons are facing the astrocytes (Fig. 2, right).



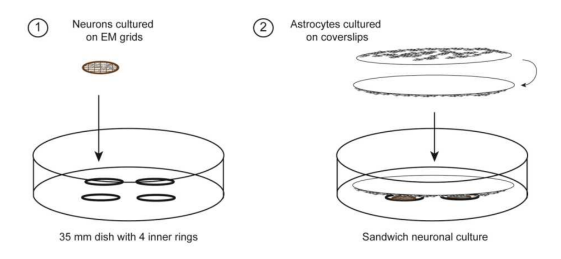


Fig 2: Schema of the sandwich cultures on EM grids

- To stop the proliferation of glial cells on the grids, treat the primary neuron cultures with the anti mitotic agent arabinofuranoside (cytosine-β-arabinofuranoside hydrochloride) three days after plating the hippocampal cells.
- 14.1 Add arabinofuranoside to a final concentration of [M] 2.45 micromolar (µM) to each dish and distribute evenly by gently moving the dish in a circular motion.
- 15 The neurons are considered mature starting from 15 days in vitro (DIV15).

Protocol references

Kaech, S., and Banker, G. (2006). Culturing hippocampal neurons. Nature Protocols 1, 2406-2415.