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Abstract

This protocol details the immunofluorescence staining in mouse brain tissue sections.



Materials

Pyrex® Staining Dish **Ted Pella Inc. Catalog #**36754-60

⊗ 8-Section Staining Nets **Ted Pella Inc. Catalog #**36154-64

⊠ Fisherbrand[™] Superfrost[™] Plus Microscope Slides **Fisher Scientific Catalog #**12-550-15

⊠ ProLong[™] Diamond Antifade Mountant **Invitrogen - Thermo Fisher Catalog #**P36961

Primary Antibodies:

A	В	С	D
Target	Species	Conc	Manufacturer
TH	Rabbit	1:2000	Novus Biological N300109
TH	Chicken	1:750	Abcam ab76442
GFP	Chicken	1:2000	Aves Labs- GFP-1010
GFP	Rabbit	1:500	Thermofisher A-11122
GFP	Mouse	1:1000	Roche 11814460001
P62	Guinea pig	1:2000	Progen GP62-C
LAMP2	Rat	1:1000	Abcam ab13524
GAD67	Mouse	1:500	Millipore Sigma MAB5406
parvalbumin	Rabbit	1:500	Abcam ab11427
TFE3	Rabbit	1:500	Abcam ab93808

Secondary Antibodies (From Thermofisher):

A	В	С	D
Antibody	Fluorophore	Concentration	Cat Number
goat-anti rabbit	488	1:500	A-11008
goat-anti rabbit	546	1:500	A-11010



A	В	С	D
goat-anti rabbit	647	1:500	A-21245
goat-anti mouse	488	1:500	A-11029
goat-anti mouse	546	1:500	A-11003
goat-anti rat	647	1:500	A-21247
goat-anti chicken	488	1:500	A-11039
goat-anti chicken	647	1:500	A-21449



Day 1

30m

- Staining of 35μm free-floating mouse brain sections is performed in glass staining dishes (Pyrex 36754-60) using 8-section staining nets (Ted Pella 36154-64).
 - dx.doi.org/10.17504/protocols.io.5jyl8pzk9g2w/v1
- 1.1 The sections can be transferred between wells using a paint brush.
- 1.2 Volume of solution:
 - 🚨 20 mL 🚨 30 mL for antibodies.
 - <u>A</u> 50 mL for washing.
- Wash sections 3 times (5 min each wash) in Phosphate Buffer Saline (PBS) at
 - Room temperature to remove cryo-protectant solution.
- 2.1 Wash sections for 00:05:00 in Phosphate Buffer Saline (PBS) at Room temperature to remove cryo-protectant solution (1/3).



5m

2.2 Wash sections for 00:05:00 in Phosphate Buffer Saline (PBS) at Room temperature to remove cryo-protectant solution (2/3).



5m

2.3 Wash sections for 00:05:00 in Phosphate Buffer Saline (PBS) at Room temperature to remove cryo-protectant solution (3/3).



5m

Wash sections 3 times (5 min each wash) in PBS + 0.1% triton X-100.



3.1 Wash sections for 00:05:00 in PBS + 0.1% triton X-100 (1/3).

5m

3.2 Wash sections for 00:05:00 in PBS + 0.1% triton X-100 (2/3).

5m

3.3 Wash sections for 00:05:00 in PBS + 0.1% triton X-100 (3/3).

5m

4 Block sections for (2) 01:00:00 in PBS + 10% Normal Goat Serum (NGS) + 0.4% Bovine Serum 2h Albumin (BSA) and 0.2% triton X-100 for 01:00:00 at Room temperature. 5 Incubate sections with primary antibodies in blocking solution (PBS + 0.1% triton X-100, 0.4% 1h BSA) Overnight at 4 °C. Day 2 2d 2h 5m 6 Wash sections 3 times (10 min each wash) in PBS + 0.1% triton X-100 at Room temperature 6.1 Wash sections for 00:10:00 in PBS + 0.1% triton X-100 at Room temperature (1/3). 10m 6.2 Wash sections for 00:10:00 in PBS + 0.1% triton X-100 at Room temperature (2/3). 10m 6.3 Wash sections for 00:10:00 in PBS + 0.1% triton X-100 at Room temperature (3/3). 10m 7 Incubate with fluorescent secondary antibodies (1:500) diluted in PBS + 0.1% triton X-100 + 0.4% 1d BSA for 2 hours at \$\mathbb{{\mathbb{g}}}\$ Room temperature or \(\mathbb{C}\) 24:00:00 at \$\mathbb{{\mathbb{g}}}\$ 4 °C . Note Sections should be protected from light at this step and all proceeding steps. 8 Wash sections 3 times (10 min each wash) in PBS + 0.1% triton X-100 at 🖁 Room temperature 8.1 Wash sections for 00:10:00 in PBS + 0.1% triton X-100 at Room temperature (1/3). 10m

8.2 Wash sections for 00:10:00 in PBS + 0.1% triton X-100 at Room temperature (2/3). 10m 8.3 Wash sections for 00:10:00 in PBS + 0.1% triton X-100 at Room temperature (3/3). 10m 9 Incubate with DAPI in PBS (1:5000) for 00:30:00 at 8 Room temperature. 30m 10 Wash sections 3 times (10 min each wash) in PBS at | Room temperature |. 10.1 Wash sections for 00:10:00 in PBS at Room temperature (1/3). 10m 10.2 Wash sections for 00:10:00 in PBS at Room temperature (2/3). 10m 10.3 Wash sections for 00:10:00 in PBS at Room temperature (3/3). 10m 11 Mount sections: 11.1 Use a petri dish filled with PBS to mount the sections on superfrost plus slides (Fisher Scientific 12-550-15). 11.2 This is easiest to do using a medium sized paint brush. 11.3 Let sections dry (5) 00:02:00 - (5) 00:05:00 . 7m Note Coverslip using ProLongTM Diamond Antifade Mountant (Thermofisher P36961).



12 Dry slides for 👏 24:00:00 at 🖁 Room temperature .



- 1. Protect slides from light at this step.
- 2. Store slides at 🔓 4 °C .

