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DAB staining protocol for subsequent stereological cell counting

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Protocol status: Working

We use this protocol and it's working



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Abstract

This protocol describes the steps for performing a double chromogen staining using DAB and SK- 4700. Stained sections can subsequently be imaged and used for stereological cell quantification with Stereo Investigator software from MBF Bioscience or other Stereology software.

Attachments



[n75mb93sf.pdf](#)

97KB

Materials

Materials:

- Pre-cut brain sections (30-50 μm thick)
- 0.1 M PB
- Triton-X100 (detergent)
- Methanol (e.g. from Sigma-Aldrich)
- Ethanol absolute (e.g. from Sigma-Aldrich)
- 30% H_2O_2 (e.g. from Merck)
- Xylene (e.g. from J.T.Baker)
- Na-hypochlorite (e.g. Sigma-Aldrich)
- Normal donkey serum (S30-100ML, Sigma-Aldrich)
- Primary antibodies (e.g. Anti-ChAT AB144P, Merck; anti-TH ab113, abcam; anti-p62, ab109012, abcam; anti-alpha-synuclein(phosphoS129), ab51253, abcam)
- Biotinylated secondary antibody
- Vectastain® Elite® ABC HRP Kit (Vector Laboratories)
- 3,3'-diaminobenzidine (DAB) (e.g. from Serva)
- SK-4700 SG Peroxidase Kit (Vector Laboratories)
- 12-well-plates with netwell inserts (e.g. Corning Costar Netwell)
- Quick-hardening Mounting Medium (e.g. Eukis, Sigma Aldrich)
- Microscopy slides
- Glass coverslips
- Designated container for storage of microscopy slides
- Orbital shaker (e.g. Heidolph Duomax 1030)
- Plajorm shaker (e.g. Heidolph Unimax 1010)
- Microscope system configured for brightfield (and multi-channel fluorescent work) with Stereo Investigator software from MicroBrightField (MBF) (e.g. Zeiss Axio Imager.M2 with MBF extension modules needed for Stereo Investigator software)

Recommended PPE:

- Lab coat/disposable gown
- Safety goggles
- Examination gloves
- FFP2 mask/respirator

Primary antibodies:

 Goat anti-ChAT antibody **Merck Millipore (EMD Millipore) Catalog #AB144P**

 Anti-Tyrosine Hydroxylase antibody **Abcam Catalog #ab113**

 Recombinant Anti-SQSTM1 / p62 antibody [EPR4844] - Autophagosome Marker **Abcam Catalog #ab109012**

 Recombinant Anti-Alpha-synuclein (phospho S129) antibody **Abcam Catalog #ab51253**



DAB staining – Before the procedure:

12h 47m

- 1 Prepare [IM] 0.1 Mass Percent PB (PB).
- 2 Prepare PBT ([IM] 0.1 Mass Percent PB with 0.3% Triton X-100).
- 3 Prepare a quenching solution (for 40 mL solution: 32 mL [IM] 0.1 Mass Percent PB, 4 mL methanol (100%), 4 mL H₂O₂ (30%)).
- 4 Prepare NDS solution (5% normal donkey serum diluted in PBT).

DAB Staining Procedure

12h 47m

- 5 Place brain sections (30 µm - 50 µm) in 12-well-plates with netwell inserts (up to 6-10 sections per netwell insert, depending on the size of the sections).
- 6 Place 12-well-plates on platform shaker and wash sections for 3× 5 min in [IM] 0.1 Mass Percent PB. Exchange PB solution in between washing steps.
- 6.1 Place 12-well-plates on platform shaker and wash sections for 00:05:00 in [IM] 0.1 Mass Percent PB. Exchange PB solution in between washing steps (1/3).
- 6.2 Place 12-well-plates on platform shaker and wash sections for 00:05:00 in [IM] 0.1 Mass Percent PB. Exchange PB solution in between washing steps (2/3).
- 6.3 Place 12-well-plates on platform shaker and wash sections for 00:05:00 in [IM] 0.1 Mass Percent PB. Exchange PB solution in between washing steps (3/3).
- 7 Quench sections for 00:15:00 at Room temperature in quenching solution in 12-well-plates on platform shaker.

5m



5m



5m



15m

**Note**

We recommend  4 mL solution per well for good results.

8 Thereafter, wash sections for 4× 5 min in  0.1 Mass Percent PB on platform shaker. Exchange PB solution in between washing steps.



8.1 Thereafter, wash sections for  00:05:00 in  0.1 Mass Percent PB on plajorm shaker. Exchange PB solution in between washing steps (1/).

5m



8.2 Thereafter, wash sections for  00:05:00 in  0.1 Mass Percent PB on plajorm shaker. Exchange PB solution in between washing steps (2/4).

5m



8.3 Thereafter, wash sections for  00:05:00 in  0.1 Mass Percent PB on plajorm shaker. Exchange PB solution in between washing steps (3/4).

5m



8.4 Thereafter, wash sections for  00:05:00 in  0.1 Mass Percent PB on plajorm shaker. Exchange PB solution in between washing steps (4/4).

5m

9 Block sections for  01:00:00 at  Room temperature in NDS solution in 12-well-plates.

1h

Note

We recommend  4 mL solution per well for good results.

10 For incubation with primary antibody (e.g. anti neuronal nuclei (NeuN), Merck Millipore, MAB377, 1:1000), transfer sections in a new 12-well-plate but without netwell inserts. This allows better shaking  Overnight . Solution for incubation with primary antibody should contain NDS solution and the respective primary antibody diluted according to manufacturer recommendation. Incubate sections on orbital shaker with gentle shaking at  4 °C  Overnight .

2h



**Note**

We recommend at least 1 ml solution for each well.

- 11 On the next day, transfer sections into 12-well-plate with netwell inserts and wash sections 4× 5 min in PB on a platform shaker at Room temperature . Change washing solution after each washing step.



- 11.1 On the next day, transfer sections into 12-well-plate with netwell inserts and wash sections for 00:05:00 in PB on platform shaker at Room temperature . Change washing solution after each washing step (1/4).

5m

- 11.2 On the next day, transfer sections into 12-well-plate with netwell inserts and wash sections for 00:05:00 in PB on platform shaker at Room temperature . Change washing solution after each washing step (2/4).

5m

- 11.3 On the next day, transfer sections into 12-well-plate with netwell inserts and wash sections for 00:05:00 in PB on platform shaker at Room temperature . Change washing solution after each washing step (3/4).

5m

- 11.4 On the next day, transfer sections into 12-well-plate with netwell inserts and wash sections for 00:05:00 in PB on platform shaker at Room temperature . Change washing solution after each washing step (4/4).

5m

- 12 Next, transfer sections back in 12-well-plate without netwell inserts for incubation with biotinylated secondary antibody (e.g., biotinylated donkey anti-mouse, Jackson ImmunoResearch, 715-065-151, 1:500) diluted in NDS solution. Incubate for 01:00:00 on orbital shaker at Room temperature .

1h

**Note**

We recommend at least 1 ml solution for each well.

- 13 Next, without washing, incubate sections in 12-well-plate without netwell inserts on orbital shaker at Room temperature for 01:00:00 in avidin-biotin-peroxidase solution using the Vectastain ABC Kit, diluted according to manufacturer's recommendation.

1h





- 14 Prepare 5% DAB working solution fresh before color reaction (for 40 ml solution: 35.2 mL 0.1 Mass Percent PB, 4 mL DAB solution (5 μ L), 0.8 mL H_2O_2 (1%)). Wear an FFP2/N95 mask or a respirator for this step and all further steps in which you work directly with DAB. Only carry out this work under a fume hood.
- 15 Without washing, transfer brain sections back in 12-well-plates with netwell inserts and start DAB color reaction by incubating sections in DAB working solution for 3- 00:06:00 at Room temperature under fume hood, depending on desired color intensity. Gently shake sections during incubation to avoid sections sticking together. 6m
- 16 Stop color reaction by washing sections 4 $\times$ 5 min. in 0.1 Mass Percent PB on plajorm shaker. Exchange PB solution in between washing steps. Use a fresh 12-well-plate which was not previously in contact with DAB solution. While washing use diluted Na-hypochlorite solution to neutralize DAB solution on used 12-well-plate and inserts. Let plastics neutralize in diluted Nahypochlorite solution Overnight . 5m
- 16.1 Stop color reaction by washing sections for 00:05:00 in 0.1 Mass Percent PB on plajorm shaker (1/4). 5m
- 16.2 Stop color reaction by washing sections for 00:05:00 in 0.1 Mass Percent PB on plajorm shaker (2/4). 5m
- 16.3 Stop color reaction by washing sections for 00:05:00 in 0.1 Mass Percent PB on plajorm shaker (3/4). 5m
- 16.4 Stop color reaction by washing sections for 00:05:00 in 0.1 Mass Percent PB on plajorm shaker (4/4). 5m
- 17 Block sections again for 01:00:00 at Room temperature in NDS solution in 12-well-plates on orbital shaker. 1h

Note

We recommend 4 mL solution per well for good results.



- 18 For incubation with primary antibody (e.g. anti-tyrosine hydroxylase (TH), Merck Millipore, AB152, 1:1000), transfer sections in a new 12-well-plate but without netwell inserts. This allows better shaking Overnight . solution for incubation with primary antibody should contain NDS solution and the respective primary antibody diluted according to manufacturer recommendation. Incubate sections on orbital shaker with gentle shaking at 4 °C Overnight .

2h

**Note**

We recommend at least 1 mL solution for each well.

- 19 On the next day, transfer sections into 12-well-plate with netwell inserts and wash sections 4× 5 min in PB on platform shaker at Room temperature . Change washing solution after each washing step.



- 19.1 On the next day, transfer sections into 12-well-plate with netwell inserts and wash sections for 00:05:00 in PB on platform shaker at Room temperature . Change washing solution after each washing step (1/4).

5m

- 19.2 On the next day, transfer sections into 12-well-plate with netwell inserts and wash sections for 00:05:00 in PB on platform shaker at Room temperature . Change washing solution after each washing step (2/4).

5m

- 19.3 On the next day, transfer sections into 12-well-plate with netwell inserts and wash sections for 00:05:00 in PB on platform shaker at Room temperature . Change washing solution after each washing step (3/4).

5m

- 19.4 On the next day, transfer sections into 12-well-plate with netwell inserts and wash sections for 00:05:00 in PB on platform shaker at Room temperature . Change washing solution after each washing step (4/4).

5m

- 20 Next, transfer sections back in 12-well-plate without netwell inserts for incubation with biotinylated secondary antibody (e.g., biotinylated donkey anti-rabbit, Jackson

1h





ImmunoResearch, 711-065-152, 1:500) diluted in NDS solution. Incubate for

01:00:00 on orbital shaker at Room temperature .

Note

We recommend at least 1 mL solution for each well.

- 21 Next, without washing, incubate sections in 12-well-plate without netwell inserts on orbital shaker at Room temperature for 01:00:00 in avidin-biotin-peroxidase solution using the Vectastain ABC Kit, diluted according to manufacturer's recommendation.
- 22 Prepare Peroxidase Substrate Kit (SG Peroxidase Substrate Kit, Vector Labs, SK-4700) as recommended by the manufacturer. Wear an FFP2/N95 mask or a respirator for this step and all further steps in which you work directly with SK-4700. Only carry out this work under a fume hood.
- 23 Without washing, transfer brain sections back in 12-well-plates with netwell inserts and start SK-4700 color reaction by incubating sections in prepared peroxidase substrate solution for 3- 00:06:00 at Room temperature under fume hood, depending on desired color intensity. Gently shake sections during incubation to avoid sections sticking together.
- 24 Stop color reaction by washing sections 4× 5 min. in 0.1 Mass Percent PB on platform shaker. Exchange PB solution in between washing steps. Use a fresh 12-well-plate which was not previously in contact with SK-4700 solution. While washing use diluted Na-hypochlorite solution to neutralize SK-4700 solution on used 12-well-plate and inserts. Let plastics neutralize in diluted Na-hypochlorite solution Overnight .
- 24.1 Stop color reaction by washing sections for 00:05:00 in 0.1 Mass Percent PB on platform shaker (1/4).
- 24.2 Stop color reaction by washing sections for 00:05:00 in 0.1 Mass Percent PB on platform shaker (2/4).



- 24.3 Stop color reaction by washing sections for 00:05:00 in **IMJ 0.1 Mass Percent** PB 5m
- 24.4 Stop color reaction by washing sections for 00:05:00 in **IMJ 0.1 Mass Percent** PB 5m
- 25 After washing, mount sections on microscopy slides using a fine brush.
- 26 Let sections dry Overnight . 5m
- 27 Immerse dried slides in a series of ethanol (70%, 96%, 100%) 30 sec. each for stepwise dehydration. Work under a fume hood.
- 28 Let slides rest for 00:10:00 in 100% xylene solution. Work under a fume hood. 10m
- 29 Next, quickly apply a small amount of hard-drying mounting medium (e.g., EukIS) sufficient to cover the sections. Carefully avoid the formation of air bubbles. Gently apply a coverslip over the sections and the mounting medium. Work under a fume hood.

DAB staining – After the procedure:

- 30 Let slides cure for 2 days under a fume hood.
- 31 Dispose of waste and excess reagents/solution according to institutional guidelines.
- 32 Clean tools/working station.
- 33 Microscopy slides should be stored in a designated container at room temperature until time of observation.

Stereological cell counting – Before the procedure:

- 34 Turn on microscope and computer according to specific manuals/instructions.



- 35 Turn on MBF Stereo Investigator software.

Stereological cell counting – Procedure:

- 36 Load microscopy slides in the designated stage.
- 37 Use brightfield settings for imaging.
- 38 Carefully follow all steps of the Stereo Investigator Workflow according to the manual. The workflow is step-by-step and very intuitive.
- 39 We typically use a 60x oil immersion lens for imaging.
- 40 After cell counts have been collected export counting data into an excel sheet for further data analysis.

Stereological cell counting – After the procedure:

- 41 Clean immersion objectives with a lens wipe and the appropriate cleaning solution.
- 42 Turn off software/microscope/laser according to specific instructions.