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### Western blot - alpha-synuclein

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#### **ABSTRACT**

This protocol describes how to detect alpha-synuclein in protein derived from STC-1 cells by western blot using DAB/peroxidase or ECL to visualise the bands.

#### **MATERIALS**

RIPA buffer (50 mM Tris-HCl, pH 8.0, with 150 mM sodium chloride, 1.0% Igepal CA-630 (NP-40), 0.5% sodium deoxycholate, and 0.1% sodium dodecyl sulphate)

Protease inhibitors (0.8 µM Aprotinin, 40 µM Bestatin, 140 µM E-64 at, Leupeptin at 20 μM and Pepstatin A at 15 μM, 1mM phenylmethanesulfonyl fluoride)

Towbin transfer buffer: 25 mM Tris, 192 mM glycine, pH 8.3, with 20% methanol (vol/vol).

To prepare 1 L of buffer:

- -800 mL distilled H20
- 200 mL methanol
- -3.03 q Tris base
- 14.4 g Glycine

Sample preparation

1d 4h 16m

Sample preparation using either protein pellet from an extraction (step 1.1) or direct lysis (step 1.2) 1.1 30m 30s Dissolve protein pellets from the TRI Reagent® extraction in 2% SDS containing 8 M urea. Vortex 00:00:30 twice, then orbital shaker 05 260 rpm 00:30:00 1.2 Dissolve cells in RIPA buffer containing protease inhibitors (see Materials) | On ice 30m 00:30:00 2 15m 20s Mix samples with 4x NuPAGE™ LDS sample buffer containing 50 mM dithiothreitol Vortex 00:00:20 twice, then orbital shaker 5 260 rpm 00:15:00 3 10m 4 1m 50s Centrifuge samples 00:00:30 to collect condensation from tube lid. Vortex 00:00:20 then centrifuge again 600:01:00 1d 4h 16m **Electrophoresis and transfer** 45m 5 Separate samples (~ 10 ug total protein) by polyacrylamide gel electrophoresis using precast Bolt™ or NuPAGE™ 12%, Bis-Tris, 1.0 mm, Mini Protein Gels at 180 V 👏 00:45:00 or until dye front reaches the bottom of the gel. Run with pre-stained or biotinylated size markers 6 Wet PVDF membrane with methanol and then equilibrate in Towbin buffer (see materials)

7 Soak transfer sandwich components (4 sheets of filter paper and 5 blotting pads) in Towbin transfer buffer and assemble in the transfer cassette in the following order:

Starting with the cathode plate

2 x blotting pads

2 x filter paper

gel

**PVDF** 

2 x filter paper

3 x pad (use 3 because there is less chance of the gel slipping)

Use a roller to remove any air bubbles

#### anode



2X Blotting pads Filter paper Membrane Gel Filter paper 2X Blotting pads

cathode

8 Place cassette in transfer tank and transfer protein onto 0.2 mm PVDF (polyvinylidene difluoride) membranes (30 V, 100:00 in Towbin buffer

9 Rinse membranes with PBS (5) 00:00:30

30s

## Immunodectection of protein bands

1d 1h 5m 30s

10

Fix proteins to membrane for 00:20:00 with 4% paraformaldehyde



20m

- 11 5m Wash with PBST (PBS containing 0.01% (v/v) Tween™-20) 4 x ♦ 00:05:00 12 Block non-specific binding sites with block solution (PBST containing 2% (w/v) BSA and 0.005 % (w/v) thiomersal) for 01:00:00 13 20h Incubate membranes in block solution containing rabbit monoclonal antibody against α-synuclein (1:1250) (ab212184) and β-actin (1:5000) (ab241153) (20:00:00 Room temperature 14 Wash with PBST 4 x 00:05:00 5m 15 Incubate membrane with secondary antibody in block solution (step 15.1 or step 15.2)
  - 15.1 Incubate membrane with biotinylated goat anti-rabbit IgG (1:1000) (Stratech) 01:15:00 and go to step 16
  - OR incubate membrane with goat anti-rabbit conjugated to peroxidase (1:25,000) (Stratech)

    15.2

    OR incubate membrane with goat anti-rabbit conjugated to peroxidase (1:25,000) (Stratech)

    15.2

    OR incubate membrane with goat anti-rabbit conjugated to peroxidase (1:25,000) (Stratech)
- 16 Wash with PBST 4 x 00:05:00

- 17 45m incubated with peroxidase conjugated streptavidin (1:1000) (Roche) 00:45:00 18 5m Wash with PBST **4** x • 00:05:00 19 Wash with PBS 2 x 600:05:00 5m 20 Visualise bands with DAB if biotinylated secondary and streptavidin-peroxidase were used (step 20.1) or use ECL plus (Amersham) if anti-rabbit peroxidase secondary was used (step 20.4) 20.1 30s For DAB detection incubate in PBS containing 0.05 % (w/v) 3,3'-diaminobenzidine, 0.015 % (v/v)  $H_2O_2$  and 0.05 % (w/v) nickel ammonium sulphate for  $\leq$  00:00:30 20.2 5m Rinse in running tap water > (5) 00:05:00 20.3 Dry membrane 60°C 1 hour 20.4
  - **20.4** For chemiluminescent detection use an Amersham ECL plus kit according to the manufacturer's instructions
  - 21 Visualize ECL or digitize DAB stained membrane using a ChemiDoc™ MP imaging system (Biorad)

22 ECL reagents can be washed off the membrane and the membrane then processed for DAB staining if desired