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Protocol for performing PINK1 siRNA knockdown in mouse embryonic fibroblasts (MEFs)

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We use this protocol and it's

working

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Abstract

This protocol details the siRNA knockdown in mouse embryonic fibroblasts (MEFs) for PINK1, but applicable for any other target.

Attachments



926-2384.docx

17KB

Guidelines

72-Hr siRNA knockdown in MEFs in a 6-well plate

- If using Immortalised MEFs seed 100,000 cells per well.
- If using Primary MEFs seed 200,000 cells per well.

Materials

Key Reagents

• siRNA as purchased from Horizon Discovery as ON-TARGETplus siRNA Reagents (DharmaconTM siRNA solutions).

Note

Protocol for 5 nmol quantities of purchased siRNA

- 🔀 Lipofectamine™ RNAiMAX Transfection Reagent Thermo Fisher Catalog #13778150
- Oligomycin
- Antimycin
- MLi2



Day 1 - Cell Seeding and siRNA preparation

- 1 For Primary MEFs seed 200,000 cells per well in a 6-well plate at a total volume of 🚨 2 mL .

3

Day 1 - Cell Seeding and siRNA preparation

- 4 Resuspend the dried siRNA protocol for 5 nmol of purchased siRNA to reconstitute at [IM] 10 micromolar (µM)].
- 4.1 Add 400 µL of RNA-free water.
- 4.2 Add 🚨 100 µL of 5x siRNA Buffer.
- 4.3 Incubate in hood for $\bigcirc 00:05:00$, vortex vigorously and store at $\boxed{\$}$ -20 $^{\circ}$ C .

Note

The protocol can be amended if the nmol quantity of siRNA purchased is higher/lower by adjusting the resuspension to ensure a final [$_{\mbox{\scriptsize IM}}$] 10 micromolar ($_{\mbox{\scriptsize IM}}$)] concentration.

Day 2 - siRNA Knock-Down

5m

5m

Prepare mastermix according to the number of wells. Four tubes have to be prepared, 2 with Optimem and lipofectamine and one each with the PINK1 and scramble siRNA respectively.



Note

For PINK1 knockdown a final concentration of 25 nM of siRNA in each well is used.

5.1 Tube1 PINK1 siRNA:

> Per well - Δ 5 μL of PINK1 siRNA at [[M] 10 micromolar (μM)] diluted in Δ 100 μL OPTI-MEMTube 3

5.2 Tube 2:

Per well - Δ 10 μL of Lipofectamine diluted in Δ 100 μL of siRNA-OPTI-MEM

5.3 Tube 3 scramble siRNA

> Per well - Δ 5 μL of scramble siRNA at [[M] 10 micromolar (μM)] diluted in Δ 100 μL OPTI-MEM

5.4 Tube 4 (same as tube 2)

Per well - Δ 10 μL of Lipofectamine diluted in Δ 100 μL of siRNA-OPTI-MEM

- 6 Vortex slowly and combine the content of Tube 1 with tube 2 (PINK1 siRNA) and similarly with tube 3 and 4.
- 7 Vortex again slowly and incubate for 00:05:00 at Room temperature.

5m

8 Add drop by drop, Add drop by drop by drop, Add drop by drop b the scramble siRNA for controls.



Note

For an experiment with 4x 6 well plates (WT and mutant cells treated with scramble or

the following volumes can be used:

- Tube1 and tube 3: 🚨 60 µL siRNA + 🚨 1200 µL OPtiMEM
- Tube 2 and 4:
 △ 120 µL of lipofectamine + △ 1200 µL of otpimem
- This gives \bot 2580 μ L of each mix (\bot 180 μ L spare)

Day 4 - Mitochondrial depolarization

5m

- 9 Make a 500x of Oligomycin/antimycin solution. The final concentration in the well is:
 - oligo: [M] 1 micromolar (µM)
 - antimycin: [M] 10 micromolar (μM)
- 9.1 Add \triangle 4 μ L of this 500x solution will be to each well.

Note

For a \perp 100 μ L of 500x solution, we use:

	A	В
Г	Compound	Quantity
Г	Antimycin A 50 mM solution	10 ul
Г	Oligomycin 6.4 mM solution	7.8 ul
	DMSO	82.2 ul



10 Treat cell with Oligomycin/antimycin A for 24:00:00 . OA should be added 48:00:00 3d after the addition of the siRNA. Include DMSO control. Day 5 - Cell lysis 1h 50m 11 Prepare working stock of MLi2 at a concentration of [M] 100 micromolar (µM) in DMSO. 12 Treat cells with MLi2([M] 10 nanomolar (nM)) for (2 ul/well). 1h 30m 13 Lyse cells using \perp 50 μ L of Lysis buffer/well 14 Lysate can be precleared by centrifugation 3 17000 x g for 00:15:00 at 4 °C. 15m 8 15 Perform protein estimation and subject lysate to immunoblotting.