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Protocol status: Working We use this protocol and it's working

Wnt-3a and R-spo1 conditioned media reporter assay V.2

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ABSTRACT

Protocol designed to measure the activity of Wnt-3a or R-spondin-1 (Rspo1) conditioned media.

A reporter HEK cell line expressing luciferase under Wnt-3a stimulation is cultured with conditioned media followed by cell lysis and a luciferase reporter assay. Activity is compared to previous media batches or references.

IMAGE ATTRIBUTION

Created with BioRender.com

MATERIALS

Assay-reagent solution:

[M] 20 millimolar (mM)Tricine (MW 179,2)[M] 1.07 millimolar (mM)Mg Carbonate x 5 H20 (MW 485,7)[M] 2.67 millimolar (mM)Mg Sulfate x 7 H20 (MW 246,5)[M] 0.1 millimolar (mM)EDTA (MW 372,2)

Preparation (warm up for preparation):

Dissolve:

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Lysis solution

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luciferase solution stocks preparation

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Co Enzym A: . Sigma C3019-100 mg

Dissolve 100 mg in 1,27 ml in water. Make aliquotes of 100 µL

Keywords: Wnt-3a, conditioned media, activity assay, luciferase reporter assay, R-spondin conditioned

media

Luciferin

Dissolve 25 mg D-Luciferin-Na salt in 4,135 ml water. [Stock]=20 mM. Make aliquotes of 500 µL

0.5 M CDTA pH 8:

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Dissolve 8,7 g in 50 ml. Buffer the solution first with NaOH-pearls and fine adjustment with liquid NaOH ca. 10 M

0.5 M MgS04-Stock, pH 8:

Dissolve 6,15 g in 50 ml in water

1M EDTA-Stock:

Dissolve 29,2 g in 100 ml water

0.5 M Tris-Stocksolution, pH 7,8:

Dissolve 15,14 g Tris-Base(MW 121,14) in 250 ml water. Adjust pH with phosphoric acid

1M DTT-Stock:

Dissolve 1,54 g (MW 154,3) in 10 ml water

Lysis solution stocks

125 mM Tris-Phosphate 10 mM DTT 10 mM CDTA S0% Glycerin 5 % Triton X 100

5x Lysis-Solution preparation

12,5 ml Tris-Phosphat 0,5 M, pH 7,8 500 µL DTT 1 M 1 ml CDTA 0.5 M, pH 8 25 g Glycerin 2,5 g Triton X 100

Bring to a total vol. of 50 ml with water. Make10 ml aliquotes and store at-20°C

HEK-medium

DMEM F12 HEPES (Gibco Cat. No. 11330), 20% FCS 200 $\mu g/ml$ G418 material for cell counting

Microplatte for luminescence readers

Greiner Bio-one Cat. no. 655094

BEFORE START INSTRUCTIONS

Be sure to have access to:

- Luminometer with or without injection system.
- 96 well plates appropriate for luminescence signal measurements
- Reagents for culturing reporter cell line. See reference (1)

*reagents might take a while to gather and prepare but once prepared they last for a long time

*time to produce conditioned media around 3 weeks. See reference protocols (2), (3).

Day 1: Seeding of Hek 293 STF cells

1 Culture HEK 293 STF CRL-3249[™] (from now own referred as HEK-STF) according to the company specifications (1) until confluent and not too many passages old.

Seed 3 wells of HEK-STF cells per sample and following controls:

- -negative control: conditioned media from L-cells (not transfected with luciferase construct) (4)
- -positive control: previous batch with known activity, HEK-STF cells with recombinant Wnt-3a or agonist stimulation
- -control lysates as blank for luminescence: HEK-SFT cells with HEK-medium only
- 1.1 Start with one almost confluent T75 culture bottle of HEK-STF cells

1.2 1x wash with DPBS: take out medium, add 5 mL DPBS, turn gently the bottle, take out DPBS 1.3 Detach cells with 1-2 mL Trypsin/EDTA (§ 37 °C) and transfer to a conical tube with 8-9 5m mL HEK-medium 1.4 Centrifuge at 100-200 g for 00:05:00 5m count cells (Neubauer chamber or automatic cell counter) 1.5 count cells (Newbauer chamber or automatic cell counter) 5m 1.6 15m Seed cells in a 24 well plate. HEK-STF: 24 well Platte ($0.05x10^6$ cells/well). $1,3x10^6$ cells / 13ml for the whole plate cover each well with A 0.5 mL of HEK medium. 1d Day 2: cell stimulation 2 30m Aspirate medium and discard. Add a total of A 500 µL of HEK medium with the following amount of conditioned medium (CM) to test: ■ For R-spondin CM: [M] 12.5 % volume Wnt-3a CM + [M] 2.5 % volume Rspo1 CM (3) ■ For Wnt3a CM . [м] 50 % volume 2.1 Incubate at stadard culture conditions for aprox. (24:00:00) 20m **Day3: luicferasse reaction**

3 Luiferase solution

Prepare on the day luciferase solution. Keep in the fridge until ready to use When using a plate reader with automatic injection, calculate the extra volume needed for it (ca 3 mL)

fill to final volume with **Assay reagent** solution

A	В	С	D
Reagent	units	[stock]	[working]
DTT	М	1	0,033
СоА	mM	10	0,266
Luciferin	mM	20	0,467
ATP	mM	100	0,633

4 Prepare 1x Lysis medium from the stock with destiled water

20m

Cell lysisAspirate medium from HEK-STF cells

2m

5.1 Add \underline{A} 150 μL /well of 1x Lysis buffer.

- 2m
- Leave the plate for 00:20:00 at Room temperature. (Note: pippeting seems to make cell aggregates and bubbles)
- 20m

- 7 Check under the microscope that most cells are lysed
- <u>\$</u>

- 9 Transfer Δ 20 μL of each sample to a 96 well plate for luminescence read.

 Avoid bubbles or cell clumps. Recommended pipetting scheme as in the 24 well plate of HEK-STF culture-
- 10m

If using a plate reader without automatic injection:

20m

For accuracy, use a multichanel pipette and add the solution covering not one sample at a time but one of the triplcates of all samples at one time.

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(signal decreases ca. 20% in the first 10 minutes)

- 12 Check that the relative luminescence units from negative controls are orders of magnitude lower that test samples. Average values from triplicates.
 - Values below more than 50-60% lower than a working batch of conditioned media are considered of poor quality.
 - Examples of equipment and methods: Berthold Tristar-GAS ISRE Luciferase Assay, SpectraMax i3x-SpectraMax Glo Steady-Luc Reporter Assay