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CRISPR Editing of Immortalized Cell Lines with RNPs using Nucleofection

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Protocol status: Working We use this protocol at Synthego and it is working

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Abstract

This protocol describes how to deliver ribonucleoprotein (RNP) complexes that consist of purified Cas9 nuclease duplexed with chemically modified synthetic single guide RNA (sgRNA) to immortalized adherent cells or suspension cells. RNP delivery is accomplished using the Lonza 4D NucleofectorTM unit with 16-well NucleocuvetteTM Strips. A option for knock-in is included. Chemically modified sgRNAs are designed to resist exonucleases and innate intracellular immune cascades that can lead to cell death. Synthego chemically modified synthetic sgRNAs are of exceptional purity and consistently drive high editing efficiencies.

Attachments



Guidelines

Abbreviations:

CRISPR: clustered regularly interspaced short palindromic repeats Cas9: CRISPR associated protein 9 sgRNA: single guide RNA RNP: ribonucleoprotein PCR: polymerase chain reaction ICE: inference of CRISPR edits FACS: fluorescence-activated cell sorting ssODNs: single-stranded donor oligonucleotides HDR: homologydirected repair TE: Tris EDTA PBS: phosphate-buffered saline GFP: green fluorescent protein

Important Considerations

Working with RNA and RNPs

Wearing gloves and using nuclease-free tubes and reagents is recommended in order to avoid RNase contamination.

Always maintain sterile technique, and use sterile, filter pipette tips.

All Synthego and NucleofectorTM reagents should be stored according to the manufacturer's recommendations. Synthetic sgRNA should be dissolved in TE buffer and diluted to a working concentration using nuclease-free water. Please consult the Synthego <u>Quick Start Guide</u> for best practices related to dissolving and storing synthetic sgRNAs.

RNPs can be formed directly in NucleofectorTM solution.

RNP complexes are stable at room temperature for up to 1 hour (may be stored at 4°C for up to one week, or at -20°C for up to 1 month). Note that RNPs stored at 4°C may become susceptible to contamination from microbial growth after long periods of time.

Suggested Controls

| Control | Description | Purpose |
|--------------|---|--|
| Mock | Cells transfected without Cas9 and sgRNA | Wild type sequence for comparison with experimental and other negative controls. |
| | | Control toxicity from RNP, cell death from electroporation or possible viability issues associated with editing the specific gene of interest. |
| Negative | Cas9 complexed with a non-targeting sgRNA or no sgRNA | Ensure that there are no false positives due to contamination (no effect expected=wild type). |
| Positive | sgRNA with high editing efficiency (e.g., CDC42BPB, RELA) | Ensure all reagents, protocol, and equipment are functioning (effect expected). |
| Transfection | pMAX GFP (Lonza), GFP mRNA (SBI) | Assess transfection efficiency (without the use of RNPs). |

Timeline

| Pre-Nucleofection | | Setup & Nucleofection | | Post-l |
|-------------------|-------|---|-------|--------|
| Day 1 | Day 2 | Day 3 | Day 4 | Day |
| Seed Cells | | Prepare Destination Plate | | |
| Incubate (2 days) | | Assemble RNP complexes | | |
| | | Prepare Cell Suspension | | |
| | | Prepare Cell/RNP Solution | | |
| | | Transfer Cell/RNP Solution to the Nucleocuvette [™] Strip | | |
| | | Transfect Cells | | |
| | | Add Recovery Medium | | |
| | | Plate Cells | | |
| | | Incubate (3 days) | | |
| | | | | |

Additional Information

For an up-to-date list of all Synthego Protocols and other resources, please visit synthego.com/resources

For technical assistance, contact our Scientific Support Team:

Ph: 888.611.6883 Email: <u>support@synthego.com</u>

Materials

MATERIALS

- X Chemically modified sgRNA Synthego Catalog #Chemically modified sgRNA
- 🔀 Cas9 2NLS nuclease (S. pyogenes) Synthego Catalog #Cas9 2NLS nuclease
- 🔀 Positive control (optional); Recommended: human RELA sgRNA, CDC42BPB sgRNA Synthego
- X Transfection control (optional); Recommended: pMAX GFP (Lonza), GFP mRNA (SBI)
- X TE buffer (Included with Synthego sgRNA) Synthego
- X Nuclease-free water Thermo Fisher Scientific Catalog #R0581
- **X** 4D-Nucleofector System with X Unit **Lonza Catalog** #AAF-1002X
- X 4D-Nucleofector® X Kit S (32 RCT) specific for cell type Lonza Catalog #V4XC-1032
- X Cell counter Thermo Fisher Scientific
- X Normal growth medium (Cell-type dependent)
- 🔀 TrypLE Express or preferred cell dissociation reagent Thermo Fisher Scientific
- 🔀 1X PBS, cell culture grade Thermo Fisher Scientific
- X 12-well tissue culture plates Corning
- X Microcentrifuge tubes Eppendorf
- 🔀 ssDNA HDR template (optional)

Safety warnings

Please refer to the SDS (Safety Data Sheet) for safety warnings and hazard information.

Pre-Nucleofection - Seed Cells

1 Subculture cells 2 days before nucleofection and seed cells in an appropriately sized vessel so that they are 70-80% confluent on the day of transfection. Each nucleofection reaction will require $\sim 1.5 \times 10^5$ cells.

Note

Culturing cells for additional days may be necessary to reach the desired confluency.

348:00:00 Subculturing cells

Setup & Nucleofection - Prepare Destination Plate

2 Pre-warm 1 ml of normal growth medium in each well of a 12-well cell culture plate per reaction.

 \blacksquare 1 mL normal growth medium

Note

This will serve as the destination plate after nucleofection.

Setup & Nucleofection - Assemble RNP Complexes (9:1 sgRNA to Cas9 ratio)

3 In appropriate plates/tubes, assemble RNP complexes in the order shown below. Synthego recommends sgRNA:Cas9 ratios between 3:1 and 9:1 for RNP formation. Below is an example experiment using a sgRNA:Cas9 ratio of 9:1.

| RNP Components, Molarity, & Volume | | | | |
|------------------------------------|-----------------|-----------------|--|--|
| Component | Molarity | Volume | | |
| Nucleofector [™] Solution | | 18 µl | | |
| sgRNA | 30 µM (pmol/µl) | 6 µl (180 pmol) | | |
| Cas9 | 20 µM (pmol/µl) | 1 µl (20 pmol) | | |
| Total volume | | 25 µl | | |

Note

Knock-in Option: to knock in small inserts (<50 bp), an ssDNA HDR Template can be added. The recommended length of each homology arm is at least 50 bp. Add 1 μ l 60 μ M ssDNA HDR Template per reaction to each well. Optimization may be required. To knock in larger inserts and for more information on designing knock-in experiments, see <u>Tips and</u> <u>Tricks: Design and Optimization of CRISPR Knock-in Experiments.</u>

4 Incubate RNPs for 10 minutes at room temperature.

Setup & Nucleofection - Prepare Cell Suspension

5

Note

For suspension cells: spin down cells before each aspiration of culture medium and washes (step 5). Skip steps 6 and 7 below.

Aspirate cell culture medium and wash cells 1-2 times with appropriate volume of 1X concentration of PBS.

Note Do not shake or hit the flask to dislodge cells, as this may lead to clumping and inaccuracies in cell counting. **Setup & Nucleofection - Prepare Cells** 6 Add appropriate amount of TrypLE Express and incubate the cells for ~5 minutes, or until they detach from the plate completely. 00:05:00 Incubation Setup & Nucleofection - Prepare Cell Suspension 7 Neutralize the dissociation reaction with at least 2X volume of normal growth medium. 8 Count the cells to determine the cell density. 9 Aliquot enough cells to have 1.5×10^5 cells/reaction. 10 Centrifuge cells at 90 x g for 8-10 minutes at room temperature. Note The cell pellets will not be packed tightly, so care is required when removing the supernatant. 🕑 00:08:00 Centrifugation

| Set | up & Nucleofection - Prepare Cell/RNP Solution |
|-----|--|
| 11 | Resuspend the cell pellet in 5 μl Nucleofector TM Solution per reaction. |
| | Note |
| | Work quickly, but carefully, and avoid leaving cells in the Nucleofector TM Solution for longer than 15 minutes. Avoid bubble formation. |
| | \blacksquare 5 µL Nucleofector Solution |
| 12 | Add 5 μl of cell suspension to 25 μl of RNP solution to make 30 μl of cell-RNP solution per reaction. |
| | Δ 5 μL cell suspension |
| | $\stackrel{\text{\ }}{=}$ 25 μ L RNP solution |
| Set | up & Nucleofection - Transfer Cell/RNP Solution to the Nucleocuvette Strip |
| 13 | For each reaction, transfer all 30 μI of cell-RNP solution to a well of the Nucleocuvette TM strip and click the lid into place. |
| 14 | Gently tap the Nucleocuvette TM strip on the benchtop to make sure that each sample covers the bottom of each well and that there are no bubbles in the wells. |
| Set | up & Nucleofection - Transfect Cells |
| 15 | Pre-program the Nucleofector TM depending on the cell type per reaction. |
| | Note Make sure that the entire Nucleofector Supplement is added to the Nucleofector Solution (according to manufacturer's protocol) and that the mixture is not more than 3 months old. |

- 16 Place the NucleocuvetteTM strip with closed lid into the retainer of the 4D-X Core unit. Check for proper orientation of the NucleocuvetteTM strip. Larger cutout is the top (A1 and A2) and smaller cutout is the bottom (H1 and H2).
- 17 Press "Start" on the display of the core unit. After run completion, the screen should display a green "+" over the wells that were successfully transfected. Remove the cuvette strips from the Core unit.

Note

Some cell types require a 10-minute incubation at room temperature after nucleofection. Please consult the optimized Lonza protocol to see if this is a necessary step for your cell line.

Setup & Nucleofection - Add Recovery Medium

- 18 Carefully resuspend the cells in each well of the NucleocuvetteTM strip with 70 μl of prewarmed growth medium, and mix gently by pipetting up and down 2-3 times.
 - $\stackrel{\text{L}}{=}$ 70 µL pre-warmed growth medium

Setup & Nucleofection - Plate Cells

- 19 Transfer all 100 μl to the pre-warmed 12-well tissue culture plate (prepared in step 2)
- 20 Incubate the cells for 2-3 days in a humidified 37°C/5% CO₂ incubator.

37 °C Incubation

48:00:00 Incubation

Post-Nucleofection - Analysis

- 21 Extract DNA from cells 48 hours after transfection.
- 22 Conduct analyses to determine editing efficiency: PCR, Sanger sequencing, and <u>ICE analysis</u>

. Next-Gen Sequencing, FACS, or functional tests may be conducted as alternatives.

Note

Option: If storing cells for future use is desired, split cells into two groups (one for analysis and one for cell culture).