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Viral infection

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ABSTRACT

Protocol for generation and precipitation of retrovirus and infection of LHeLa cells

OPEN ACCESS



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

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Day 1

1 Seed HEK293T cells into a 10cm tissue culture plate (6.1 million cells/plate), seeding one plate per construct you are generating virus for.

Day 2

2 Transfect cells with viral and helper vectors using lipofectamine LTX. In a 15mL falcon tube, combine the following:

	Reagent	Amount
1	Viral vector construct containing cDNA of interest	9 mg
2	Gag-pol vector	6 mg
3	VSV-G vector	3 mg
4	Opti-MEM	3 mL

20m 20s

5m 20s

- Add \triangle 54 μ L of Lipofectamine LTX , vortex for \sim \bigcirc 00:00:20 . Incubate at \bigcirc Room temperature for \bigcirc 00:20:00 .
- **5** Once the 20 minute incubation starts, replace the media on the HEK cell plates with

7 mL media/plate

At the end of the incubation, add the lipofectamine/OptiMEM mix to the plate using a sterile transfer pipette, in a drop wise fashion across the plate. Gently rock the plate to mix, and return the plate to the incubator.

7	In the morning, remove the media from the HEK cell plate and dispose of the media in a beaker
	containing bleach. Add 🔼 6 mL of fresh growth media to the HEK cell plate.

Day 4

- In the late afternoon, collect the media from the HEK cell plate into a falcon tube. Add back 8 mL fresh growth media to the HEK cell plate.
- Centrifuge the media collected at maximum speed for 00:05:00 at Room temperature to pellet any debris. Pass the media through a 0.45 mm syringe filter into a clean falcon tube. Parafilm the falcon tube, and place store the media at 4 °C in the fridge.

Day 6

- In the morning, collect the media from the HEK cell plate into a falcon tube. Centrifuge the collected media at maximum speed for 00:05:00 at 8 Room temperature. Pass the media through a 0.45 mm syringe filter into a clean falcon tube.
- 11 Combine the media collected in step 9 and in step 10 into one tube. Add Lentivirus Precipitation Solution (ALSTEM) to the media at a ratio of 1:4 (Lentivirus precipitation solution : viral media).

Day 7

13 If intending to infect cells with fresh virus, seed the cells to be infected into a 6 well plate (~220K cells/well).

Day 8

2d

5m

After the 48 hour rocking incubation, centrifuge the viral precipitation mix at for 01:00:00

- Remove the supernatant, disposing of it into a beaker containing bleach. Resuspend the precipitated pellet in 3 6 mL of fresh growth media (3 mL if you require high construct
- Add polybrene at a final concentration of 8 mg/mL to each well, and rock the plate to mix. Place the cells back into the incubator.
- Any leftover virus can be frozen at 8 -80 °C for use at a future date

Day 9 - 10

expression).

After 24 – 48h (depending on the construct and expression required), remove the media from the infected cells and place in bleach to dispose. Add fresh growth media back to the cells.

Day 15 - 17

1h

After 5 - 7 days after the virus has been removed (and after three passages), using FACS, sort the cells populations to the desired fluorescent expression level.