

VirusGEN® AAV Transfection Kit

Quick Reference Protocol

Instructions for MIR 6750

Full protocol, SDS and Certificate of Analysis available at mirusbio.com/virusgen



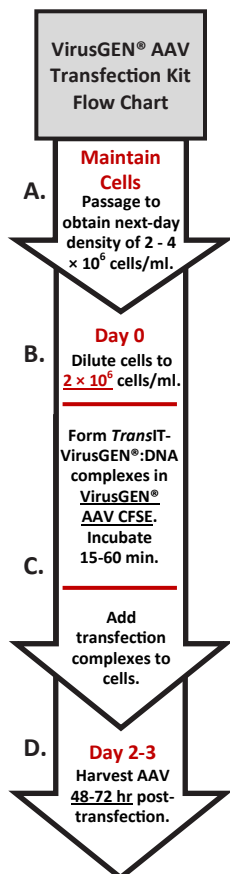
SPECIFICATIONS

Storage	Store <i>TransIT-VirusGEN®</i> Transfection Reagent at -10 to -30°C, tightly capped. Before each use , warm to room temperature and vortex gently. Store VirusGEN® AAV Complex Formation Solution and Enhancer at 2 to 10°C.
Product Guarantee	6 months from the date of purchase, when properly stored and handled.

► PROTOCOL FOR ADENO-ASSOCIATED VIRUS (AAV) GENERATION IN SUSPENSION HEK 293 CULTURES



Full protocol and additional documentation available at mirusbio.com/virusgen



Fill in volumes below based on total culture volume (Table 1).

A. Maintain cells

1. Passage suspension HEK 293 cells 18-24 hours prior to transfection to obtain a density of $2 - 4 \times 10^6$ cells/ml the next day. Do NOT proceed with transfection if cells are not doubling every 24 hours or are < 95% viable.
2. Incubate cells overnight at appropriate temperature and CO₂ levels.

B. Prepare *TransIT-VirusGEN®* Reagent:DNA complexes

1. Immediately prior to transfection, seed cells at a density of 2×10^6 cells/ml.
2. Warm *TransIT-VirusGEN®* Reagent to room temperature and vortex.
3. Place ___ ml of VirusGEN® AAV Complex Formation Solution and Enhancer (CFS&E) in a sterile tube.
4. Add ___ µl AAV plasmid DNA mixture to the tube. Mix gently by pipetting.
5. Add ___ µl of *TransIT-VirusGEN®* Reagent. Vortex gently to mix.
6. Incubate at room temperature for 15-60 minutes to allow transfection complexes to form.

C. Distribute complexes to cells in complete growth medium

1. Add *TransIT-VirusGEN®* Reagent:DNA complexes (from Step B) to cells.
2. Incubate cultures in appropriate conditions (i.e. 37°C, 5% CO₂, shaking) for 48 -72 hours prior to AAV harvest.

D. Harvest virus

1. Following the 48-72 hour incubation, prepare 10X Cell Lysis Buffer (500 mM Tris pH 8, 10% Tween® 20, 20 mM MgCl₂).
2. Transfer the total volume of cell suspension (___ ml) to a sterile conical tube or appropriate vessel.
2. Add 0.1X volume (___ ml) of 10X Cell Lysis Buffer and 100 U/ml (___ µl) of Benzonase®. Mix completely and incubate at 37°C for 1.5 hours with shaking.
3. Add 0.1X volume (___ ml) of 5 M NaCl. Mix completely and incubate at 37°C for 30 minutes with shaking.
4. Centrifuge the mixture at 4,100 × g for 10 minutes to remove cell debris.
5. Transfer the AAV-containing supernatant to a new tube. Store at -80°C.

Table 1. Volume scaling worksheet for VirusGEN® AAV Transfection Kit

Starting conditions per milliliter of complete growth medium	Total culture volume		
	Per 1 ml	x	_____ ml = _____ ml
VirusGEN® AAV CFS&E	0.1 ml	x	_____ ml = _____ ml
Total Plasmid DNA (1 µg/µl)	2 µl	x	_____ ml = _____ µl
<i>TransIT-VirusGEN®</i> Reagent	3 µl	x	_____ ml = _____ µl

Total Plasmid DNA refers to the combined mass of packaging plasmids and the transfer plasmid containing the gene-of-interest. Premix the plasmids together prior to adding to the complex formation medium.



Reagent Agent®

Reagent Agent® is an online tool designed to help determine the best solution for nucleic acid delivery based on in-house data, customer feedback and citations.

Learn more at: [mirusbio.com/ra](https://www.mirusbio.com/ra)

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