LV-mNIS



Product Description

Product Name: LV-mNIS

Catalog Number: LV008-S (0.25 mL), LV008-L (1 mL)

Lot Number: LV-OA12

Reporter gene: Murine sodium iodide symporter (mNIS)

Quantity: 250 μ L (S), 1 mL (L) Titer: 4.79 x 10⁷ TU/mL* Storage media: Serum free media

Shipping: Dry ice

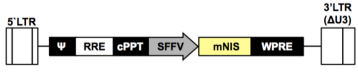
Storage: Store at ≤ -70°C upon receipt. Freeze-thaw

cycles will decrease titer.

Shelf life: One year from date of receipt under proper

storage conditions.

This is a ready-to-use lentivirus preparation. The virus encodes the murine sodium iodide symporter (mNIS) cDNA under control of the spleen focus-forming virus (SFFV) promoter (see below). The lentiviral vectors are self-inactivating (SIN) vectors in which the viral enhancer and promoter have been deleted. Transcription inactivation of the LTR in the SIN provirus increases biosafety by preventing mobilization by replication competent viruses and enables regulated expression of the genes from the internal promoters without *cis*-acting effects of the LTR¹.



5' LTR: 5' long terminal repeat ψ: RNA packaging signal RRE: Rev response element cPPT: Central polypurine tract

SFFV: Spleen focus-forming virus promoter mNIS: Murine sodium iodide symporter

WPRE: Woodchuck hepatitis virus posttranscriptional regulatory element

 3° LTR/ $\!\Delta U3$: 3° self-inactivating long terminal repeat

*Titration by qPCR:

A WPRE probe-based qPCR assay was used to measure the number of copies of lentiviruses stably integrated into the genome after transduction of HeLa H1 cells (transducing units per mL).

Traditional p24 ELISA titrations measure both functional and non-functional lentivirus particles. However, this method overestimates the functional titer, as the p24 protein pool includes a variable amount of free p24 and p24 associated with non-functional vector particles. This ratio can vary greatly between each lot, so the titration is inherently inaccurate. While qPCR titers may appear lower than p24 ELISA, they are more accurate and functional.

Safety Precaution:

All culture work with lentiviruses should be performed by trained personnel and performed under BSL2 containment following NIH guidelines.



Basic Lentivirus Transduction Protocol

Volumes are given for a 6-well plate; increase or decrease as needed. See the Transduction Tips section for additional considerations/modifications.

- 1. Seed cells in complete medium at an appropriate density to achieve 60-70% confluency the next day (e.g. $\sim 2.5 \times 10^5$ HeLaH1 cells). Incubate cells overnight in a $37^{\circ}\text{C}/5\%$ CO₂ incubator.
- 2. Thaw lentivirus stock on ice.
- In a microcentrifuge tube, dilute lentivirus to 1 mL total in serum free media. (See tips below for notes about determining optimal MOIs.)
- Remove culture medium from cells and replace with prepared lentivirus.
- 5. Return cells to 37°C/5% CO₂ incubator.
- After 4 hours add 1 mL complete medium to each well and return cells to 37°C/5% CO₂ incubator.
- 7. 3-7 days after transduction, check transgene expression according to an appropriate protocol.

Transduction Tips

- To determine the optimal MOI for transductions: plate several wells of the target cells and infect with increasing MOIs (e.g 1, 3, 10, and 30). Typically, primary cells require higher MOIs than established cell lines.
- 2. Polybrene® (Imanis #REA001) can be added to the transduction mixture to enhance transduction efficiency². The final concentration of Polybrene® in the transduction medium should be 4-8 µg/mL. Polybrene® can be cytotoxic to some cells and it is not advisable to incubate these cells with Polybrene® overnight; the transduction cocktail may be removed after 3-4 h and replaced with complete media.
- A spin infection can also be used to increase transduction efficiency³. Once the transduction mixture is added to the cells, centrifuge the plate at 800 x g for 30 min at room temperature, before placing the cells in a 37°C/5%CO₂ incubator.
- 4. The presence of serum in the transduction mixture can greatly affect transduction efficiency⁴. In general, lower serum concentrations result in higher transduction efficiencies, though optimizing serum concentrations is recommended for each cell type.

References

¹Miyoshi et al. J Virol. 1998. 72:8150-8157.

²Konopka et al. J Gen Virol 1991. 72: 2685-2696.

³O'Doherty et al. J Virol. 2000. 74:10074-10080.

⁴Andreadis and Palsson. Human Gene Ther. 1997. 8:285-291.

LV-mNIS

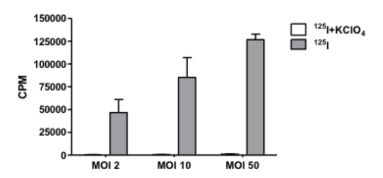


Certificate of Analysis

Testing performed by Imanis Life Sciences

Test description	Result
Virus titer	4.79 x 10 ⁷ TU/mL
Sterility	No contamination detected
NIS expression	Pass QC

NIS Expression:



HeLaH1 cells were transduced with LV-mNIS at the indicated MOIs. After 3 days, uptake of 125 l by 3 x 10 cells was assayed in the presence or absence of KClO₄, an inhibitor of NIS-mediated 125 l uptake.

Legal Disclaimers

Limited Product Warranty
This warranty limits our liability to replacement of this product. No other warranties of any kind, express or implied, including, without limitation, implied warranties of merchantability or fitness for a particular purpose, are provided by Imanis. Imanis shall have no liability for any direct, indirect, consequential, or incidental damages arising out of the use, the results of use, or the inability to use

For in vitro use only. This certificate is a declaration of analysis at the time of manufacture.

PURCHASER NOTIFICATION

LIMITED LICENSE NOTICE - RESEARCH USE ONLY

IMANIS LIFE SCIENCES HAS A LIMTED LICENSE UNDER PATENTS OWNED BY THE SALK INSTITUTE FOR BIOLOGICAL STUDIES THAT PERMITS IMANIS LIFE SCIENCES TO SELL INSTITUTE FOR BIOLOGICAL STUDIES THAT PERMITS IMANIS LIFE SCIENCES TO SELL PRODUCTS CONTAINING WPRE FOR RESEARCH USE ONLY AND NOT FOR ANY COMMERCIAL USES. EXCLUDED COMMERCIAL USES INCLUDE WITHOUT LIMITATION MANUFACTURING, PROVIDING A SERVICE, THERAPEUTIC, DIAGNOSTIC AND PROPHYLACTIC USES, AND ANY OTHER COMMERCIAL USES. USE OF THIS PRODUCT BY A PURCHASER FOR ANY PURPOSE OTHER THAN FOR RESEARCH IS UNAUTHORIZED

The Salk Institute actively licenses its patents for commercial uses, and a commercial use license may be available for Salk's WPRE patents. If you wish to inquire about such a license, please

Office of Technology Development The Salk Institute for Biological Studies 10010 North Torrey Pines Road La Jolla, CA 92037

Phone: (858) 453-4100 extension 1278

Fax: (858) 546-8093

THE NIS GENE AND TECHNOLOGY IS COVERED UNDER AN EXCLUSIVE LICENSE TO IMANIS LIFE SCIENCES. RESEARCHERS MAY USE THIS PRODUCT FOR RESEARCH USE ONLY AND NO COMMERCIAL USE IS ALLOWED. NO OTHER USE OR TRANSFER OF THIS PRODUCT OR DERIVATIVES IS AUTHORIZED WITHOUT THE PRIOR EXPRESS WRITTEN CONSENT OF IMANIS. WITH RESPECT TO ANY USES OUTSIDE THIS LABEL LICENSE, INCLUDING ANY DIAGNOSTIC, THERAPEUTIC OR PROPHYLACTIC USES, PLEASE CONTACT IMANIS LIFE SCIENCES FOR SUPPLY AND LICENSING INFORMATION.

Chief Operations Officer Imanis Life Sciences 221 1st Ave SW STE 102 Rochester MN 55902 Tel: (507) 218-2559 Email: info@imanislife.com

Quality control by: RLV Quality Assurance by: LS Effective Date: 15-Feb-2016