

Takara Bio USA, Inc.

Lenti-X™ Tet Express™ Inducible Expression System User Manual

Cat. Nos. 631189, 631355, 631356, 631357
(011018)

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I. Introduction

A. Summary

The **Tet-Express Systems** are inducible gene expression systems for mammalian cells. Target cells that contain a gene of interest (GOI) under the control of a TRE3G promoter (P_{TRE3G}) will express high levels of your GOI, when cultured in the presence of Tet-Express, a tetracycline transactivator with protein transduction properties (Figure 1).

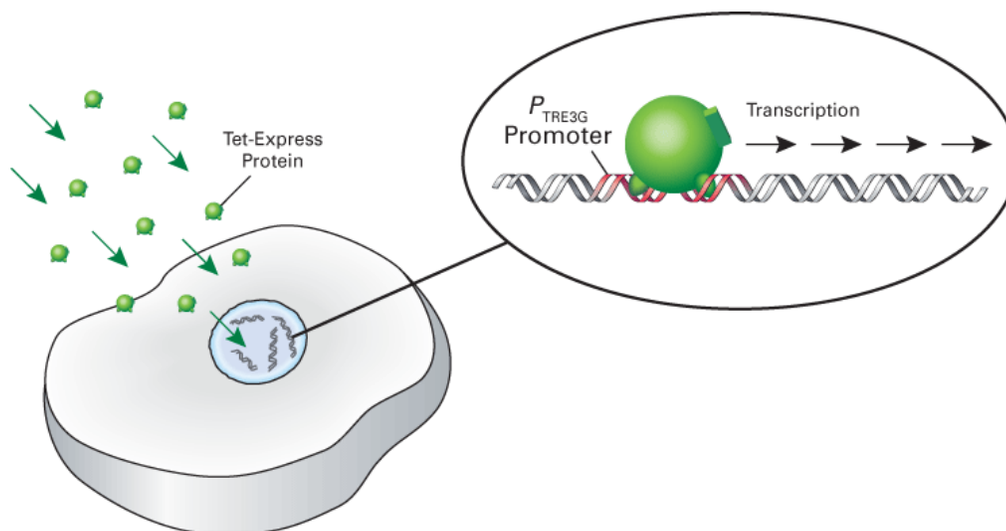


Figure 1. The Tet-Express Systems allow inducible gene expression when the self-transducing Tet-Express transactivator protein is added directly to your cells.

B. Elements of Lenti-X Tet-Express

Tet-Express Transactivator Protein

Based on the transcriptional regulators described by Gossen & Bujard (1992) and Urlinger *et al.* (2000), Tet-Express is a modified form of the Tet-Off® Advanced transactivator protein selected for its ability to transduce itself into mammalian cells. The Intensifier Reagent enhances this intrinsic protein transduction ability.

P_{TRE3GV} Inducible Promoter

The inducible promoter P_{TRE3G} provides for very low basal expression and high maximal expression after induction (Loew *et al.*, 2010). It consists of 7 repeats of a 19 bp tet operator sequence located upstream of a minimal CMV promoter. P_{TRE3GV} is a version of P_{TRE3G} that was modified at Takara Bio for higher performance in lentiviruses and retroviruses. Tet-Express binds specifically to P_{TRE3GV} and activates transcription of the downstream GOI. P_{TRE3GV} lacks binding sites for endogenous mammalian transcription factors, so it is virtually silent in the absence of induction.

4th Generation Lentiviral Packaging System

Our **Lenti-X Packaging Single Shots (VSV-G)**, provided with the **Lenti-X Tet-Express Inducible Expression System** (Cat. No. 631189), can generate lentiviral titers that are superior to most other commercially available lentiviral packaging systems. The concerted effects of multiple components in an optimized five-vector plasmid mix, pre-aliquoted and lyophilized with Xfect™ Transfection Reagent, allow the **Lenti-X 293T Cell Line** (sold separately; Cat. No. 632180) to produce the highest amounts of safe, replication-incompetent lentivirus (see takarabio.com).

II. List of Components

Store Lenti-X GoStix™ Plus at room temperature and all other components at –20°C.

A. Lenti-X Tet-Express Plasmid Systems

<u>Cat. No.</u>	<u>System Name</u>
631189	Lenti-X Tet-Express Inducible Expression System
631357	Lenti-X Tet-Express Inducible Expression System (Bicistronic Version)
631355	Lenti-X Tet-Express Inducible Expression System (with mCherry)
631356	Lenti-X Tet-Express Inducible Expression System (with ZsGreen1)

B. General System Components

All systems listed in Section II.A contain the following components (store all components at -20°C):

- 25 rxns Tet-Express (includes Tet-Express Transactivator and Intensifier Reagent)
- 10 µg response vector (see Section II.C)
- 10 µg pLVX-TRE3G-Luc Control Vector
- 16 rxns Lenti-X Packaging Single Shots (VSV-G) (Cat. No. 631275)
- Lenti-X GoStix Plus (Sample) (Cat. No. 631279; not sold separately)

C. System-Specific Response Vectors

<u>Cat. No.</u>	<u>Response Plasmid</u>
631189	pLVX-TRE3G
631357	pLVX-TRE3G-IRES
631355	pLVX-TRE3G-mCherry
631356	pLVX-TRE3G-ZsGreen1

D. Tet-Express (available separately)

<u>Cat. No.</u>	<u>Product Name</u>	<u>Size</u>
631177	Tet-Express	25 rxns
631178	Tet-Express	100 rxns

III. Additional Materials Required

A. Tetracycline-Free Fetal Bovine Serum

Contaminating tetracyclines, often found in serum, will significantly elevate basal expression when using Tet-Express. The following functionally tested tetracycline-free sera are available from Takara Bio:

<u>Cat. No.</u>	<u>Serum Name</u>
631106	Tet System Approved FBS (500 ml)
631107	Tet System Approved FBS (50 ml)
631367	Tet System Approved FBS (3 x 500 ml)
631101	Tet System Approved FBS, US-Sourced (500 ml)
631105	Tet System Approved FBS, US-Sourced (50 ml)
631368	Tet System Approved FBS, US-Sourced (3 x 500 ml)

B. Antibiotics for Selecting Stable Cell Lines

Table 1. Recommended Antibiotic Concentrations for Selecting & Maintaining Stable Cell Lines

		Recommended Concentration (µg/ml)	
<u>Cat. No.</u>	<u>Antibiotic</u>	<u>Selecting Colonies¹</u>	<u>Maintenance</u>
631306	Puromycin (100 mg)	0.25–10	0.25
631305	Puromycin (25 mg)		

¹ When selecting for single colonies, the appropriate dose must be determined empirically for your specific cell line. Test a dosage range using dishes of untransfected cells and choose the dose that kills all of the cells in 3–5 days. If all the cells die in less than 24 hr, you should use a lower dose.

C. Tet-Express

<u>Cat. No.</u>	<u>Product Name</u>	<u>Size</u>
631177	Tet-Express	25 rxns
631178	Tet-Express	100 rxns

D. Mammalian Cell Culture Supplies

- **Medium for Lenti-X 293T Cells:**
90% Dulbecco's Modified Eagle's Medium (DMEM) with high glucose (4.5 g/L), 4 mM L-glutamine, and sodium bicarbonate (Sigma-Aldrich, D5796); 10% Fetal Bovine Serum (FBS); 100 units/ml penicillin G sodium & 100 µg/ml streptomycin sulfate.
- Culture medium, supplies, and additives specific for your target cells
- Trypsin/EDTA (e.g., Sigma, Cat. No. T4049)
- Cloning cylinders or discs for isolating colonies of adherent cell lines (Sigma, Cat. No. C1059) for freezing 293T cell lines.
- Cell Freezing Medium, with or without DMSO (Sigma, Cat. Nos. C6164 or C6039)
- 6-well, 12-well, and 24-well cell culture plates; 10 cm cell culture dishes

E. Lenti-X 293T Cells

Getting the most from any lentiviral packaging system requires a host 293T cell line that transfects easily and supports high-level expression of viral proteins. Our Lenti-X 293T Cell Line was clonally selected to meet these requirements, allowing you to produce the highest possible lentiviral titers when combined with Lenti-X Packaging Single Shots (VSV-G), an optimized fourth-generation packaging system, pre-mixed and lyophilized with Xfect Transfection Reagent.

- Lenti-X 293T Cell Line (Cat. No. 632180)

F. High-Titer Packaging System

<u>Cat. No.</u>	<u>Lentiviral Packaging System</u>
631275	Lenti-X Packaging Single Shots (VSV-G) (16 rxns)
631276	Lenti-X Packaging Single Shots (VSV-G) (96 rxns)

G. Lentiviral Titer Determination

For accurate and consistent transductions, we highly recommend titrating your lentiviral stocks. Various technologies are available from Takara Bio; visit takarabio.com for details.

<u>Cat. No.</u>	<u>Lentiviral Titration Technology</u>
632200	Lenti-X p24 Rapid Titer Kit (96 rxns)
631235	Lenti-X qRT-PCR Titration Kit (200 rxns)
631280	Lenti-X GoStix Plus (20 tests)

H. Lentivirus Concentration

Use Lenti-X Concentrator to simply increase your available titer up to 100-fold without ultracentrifugation—visit takarabio.com for details.

<u>Cat. No.</u>	<u>Concentrator</u>
631231	Lenti-X Concentrator (100 ml)
631232	Lenti-X Concentrator (500 ml)

I. Transduction Enhancers

Use Polybrene (hexadimethrine bromide; Sigma-Aldrich, No. H9268), Lenti-X Accelerator (see below), or RetroNectin® (see below).

- Lenti-X Accelerator is a magnetic bead-based technology designed to accelerate lentiviral and retroviral transduction experiments; visit takarabio.com for details.
- RetroNectin is a multivalent molecule that simultaneously binds virus particles and cell surface proteins, maximizing cell-virus contact. RetroNectin, in particular, is recommended for increasing the transduction efficiency of suspension cells and stem cells; visit takarabio.com for details.

<u>Cat. No.</u>	<u>Transduction Enhancer</u>	<u>Size</u>
631256	Lenti-X Accelerator	400 µl
631257	Lenti-X Accelerator	1,000 µl
631254	Lenti-X Accelerator Starter Kit	each
T110A	RetroNectin Precoated Dish	10 dishes
T100B	RetroNectin Recombinant Human Fibronectin Fragment	2.5 mg
T100A	RetroNectin Recombinant Human Fibronectin Fragment	0.5 mg

J. Xfect Transfection Reagent

Xfect provides high transfection efficiency for most commonly used cell types.

<u>Cat. No.</u>	<u>Transfection Reagent</u>
631317	Xfect Transfection Reagent (100 rxns)
631318	Xfect Transfection Reagent (300 rxns)

K. In-Fusion® HD Cloning System

In-Fusion is a revolutionary technology that greatly simplifies cloning.

For more information, visit takarabio.com/infusion

<u>Cat. No.</u>	<u>In-Fusion Cloning Kit</u>
638909	In-Fusion HD Cloning Plus (10 rxns)
638910	In-Fusion HD Cloning Plus (50 rxns)
638911	In-Fusion HD Cloning Plus (100 rxns)

L. Stellar™ Competent Cells

We recommend Stellar Competent Cells for cloning of lentiviral and retroviral vectors. Propagation of vectors containing repeat sequences such as viral LTRs using other strains of *E.coli* may result in plasmid rearrangements. Stellar Competent Cells are sold separately and provided with all In-Fusion HD Plus Systems.

<u>Cat. No.</u>	<u>Competent Cells</u>
636763	Stellar Competent Cells (10 x 100 µl)
636766	Stellar Competent Cells (50 x 100 µl)

M. Luciferase Assay and Luminometer

These items are required when using the pLVX-TRE3G-Luc Vector as a control to test for induction (Section VI.B). Use any standard firefly luciferase assay system and luminometer.

IV. Protocol Overview

Please read each protocol completely before starting. Successful results depend on understanding and performing the following steps correctly.

A. General Cell Culture

This user manual provides only general guidelines for mammalian cell culture techniques. For users requiring more information on mammalian cell culture, transfection, and creating stable cell lines, we recommend the following general reference:

Freshney, R.I. (2005). *Culture of Animal Cells: A Manual of Basic Technique, 5th Edition* (Wiley-Liss, Hoboken, NJ).

B. Safety Guidelines for Working with Lentiviruses

The protocols in this User Manual require the production, handling, and storage of infectious lentivirus. It is imperative to fully understand the potential hazards of, and necessary precautions for, the laboratory use of lentiviruses.

The National Institute of Health and Center for Disease Control have designated recombinant lentiviruses as Level 2 organisms. This requires the maintenance of a Biosafety Level 2 facility for work involving this virus and others like it. The VSV-G pseudotyped lentiviruses packaged from the HIV-1-based vectors described here are capable of infecting human cells. The viral supernatants produced by these lentiviral systems could, depending on your insert, contain potentially hazardous recombinant virus. Similar vectors have been approved for human gene therapy trials, attesting to their potential ability to express genes *in vivo*.

IMPORTANT: For these reasons, due caution must be exercised in the production and handling of any recombinant lentivirus. **The user is strongly advised not to create VSV-G pseudotyped lentiviruses capable of expressing known oncogenes.**

For more information on Biosafety Level 2 agents and practices, download the following reference:

Biosafety in Microbiological and Biomedical Laboratories (BMBL), Fifth Edition (February 2007) HHS Pub. No. (CDC) 93-8395. U.S. Department of Health and Human Services Centers for Disease Control and Prevention and NIH.

Available on the web at <http://www.cdc.gov/od/ohs/biosfty/bmbl5/bmbl5toc.htm>

Biosafety Level 2: The following information is a brief description of Biosafety Level 2. It is neither detailed nor complete. Details of the practices, safety equipment, and facilities that combine to produce a Biosafety Level 2 are available in the above publication. If possible, observe and learn the practices described below from someone who has experience working with lentiviruses.

Summary of Biosafety Level 2:

- **Practices:**
 - Standard microbiological practices
 - Limited access to work area
 - Biohazard warning signs posted
 - Minimize production of aerosols
 - Decontaminate potentially infectious wastes before disposal
 - Use precautions with sharps (e.g., syringes, blades)
 - Biosafety manual defining any needed waste decontamination or medical surveillance policies
- **Safety equipment:**
 - Biological Safety Cabinet, preferably a Class II BSC/laminar flow hood (with a HEPA microfilter) used for all manipulations of agents that cause splashes or aerosols of infectious materials; exhaust air is unrecirculated
 - PPE: protective laboratory coats, gloves, face protection as needed
- **Facilities:**
 - Autoclave available for waste decontamination
 - Chemical disinfectants available for spills

C. Protocol Summary

The following are the steps required to create a Tet-Express-inducible expression system using lentivirus (see Figure 2).

1. Clone your gene of interest into the pLVX-TRE3G Vector using In-Fusion HD (Section V).
2. Pilot test Tet-based induction of your construct (Section VI).
3. Produce lentiviral supernatants using Lenti-X Packaging Single Shots (VSV-G) (Section VII).
4. Infect your target cells with TRE3G virus and induce expression with Tet-Express (Section IX).

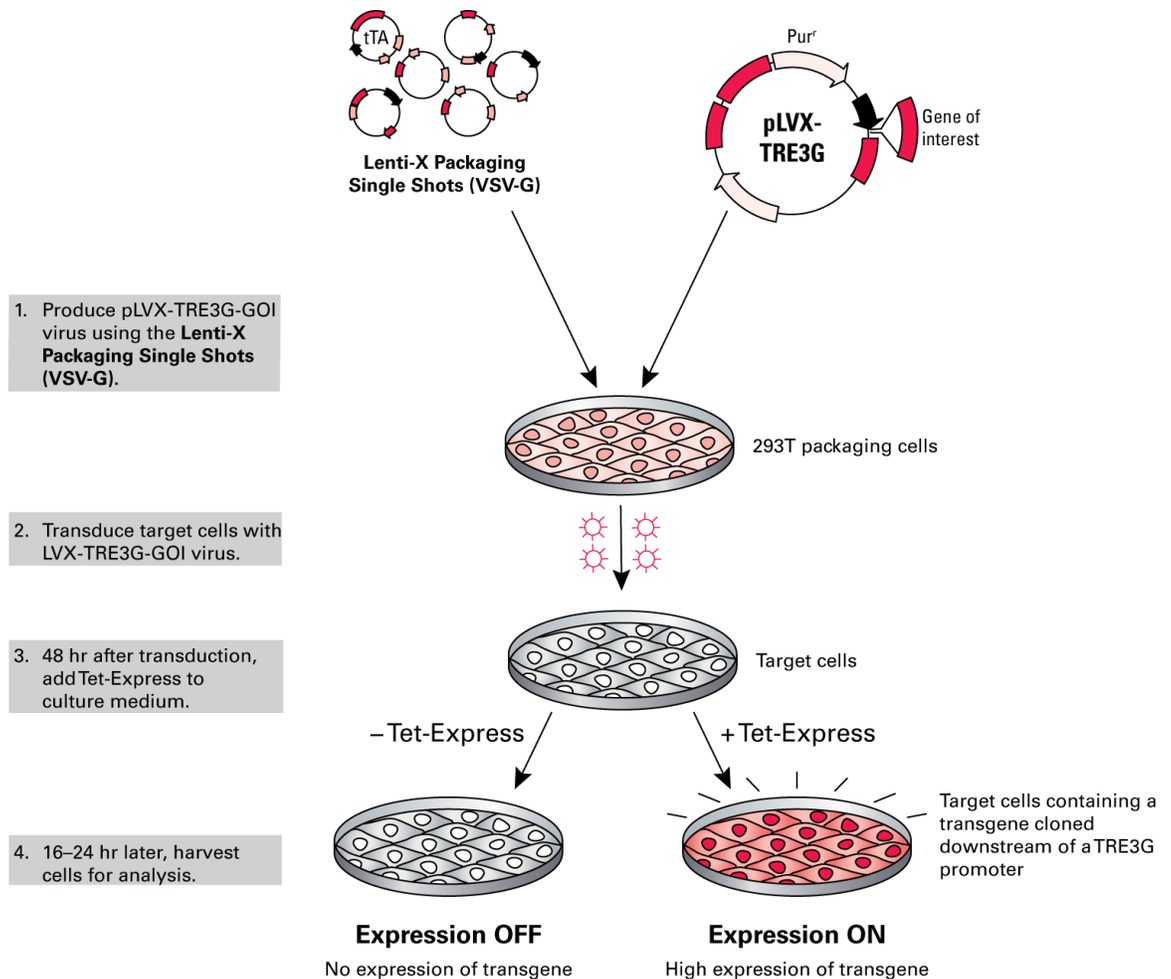


Figure 2. Establishing an inducible expression system in target cells with Lenti-X Tet-Express. Lenti-X Packaging Single Shots (VSV-G), an optimized packaging pre-mix lyophilized with Xfect Transfection Reagent, and 293T cells are used to generate high-titer lentiviral supernatants from the pLVX-TRE3G Vector, which contains your gene of interest. Target cells are then transduced with the lentivirus (~8 hr). After culturing for an additional 48 hr, Tet-Express is added to the culture medium to induce expression, which can be detected within a few hours after addition of Tet-Express.

V. Cloning Your Gene of Interest into the pLVX-TRE3G Vector using In-Fusion HD

We recommend using In-Fusion HD for all cloning. Follow the protocol outlined in the In-Fusion HD user manual at takarabio.com/infusion

NOTE: We recommend Stellar Competent Cells (Section III.K) for cloning of lentiviral and retroviral vectors. Propagation of vectors containing repeat sequences such as viral LTRs using other strains of *E. coli* may result in plasmid rearrangements. Stellar Competent Cells are provided with all In-Fusion HD Cloning Systems.

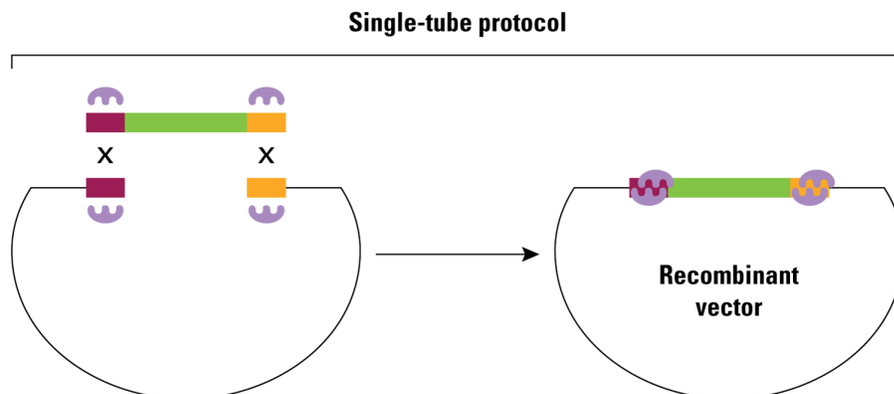


Figure 3. The In-Fusion HD Single-Tube Cloning Protocol.

Depending on which pLVX-TRE3G vector you are using, the recommended linearization sites and forward/reverse primer designs are as follows:

Response Plasmid	Linearize with	Forward Primer*	Reverse Primer**
pLVX-TRE3G	BamHI & MluI	tcttatacttgatcc 111 222 333 444 555 666 777 888	ttcccatatgacgct SSS NNN NNN NNN NNN NNN NNN NNN
pLVX-TRE3G-IRES (MCS-1)	BamHI & NotI	tcttatacttgatcc 111 222 333 444 555 666 777 888	cgcccgggcgccgc SSS NNN NNN NNN NNN NNN NNN NNN
pLVX-TRE3G-IRES (MCS-2)	MluI & EcoRI	gccccgggacgct 111 222 333 444 555 666 777 888	ctaccggtagaattc SSS NNN NNN NNN NNN NNN NNN NNN
pLVX-TRE3G-mCherry	MluI & EcoRI	gccccgggacgct 111 222 333 444 555 666 777 888	ctaccggtagaattc SSS NNN NNN NNN NNN NNN NNN NNN
pLVX-TRE3G-ZsGreen1	MluI & EcoRI	gccccgggacgct 111 222 333 444 555 666 777 888	ctaccggtagaattc SSS NNN NNN NNN NNN NNN NNN NNN

*111 = Start codon of your gene; 222 = 2nd codon of your gene; etc.

**SSS = reverse complement of the stop codon of your gene; NNN = reverse complement of the end of your gene.

VI. Pilot Testing Tet-Based Induction of Your Construct

Your pLVX-TRE3G construct should be tested for inducible expression prior to creating a Tet-Express-inducible stable cell line. Transiently transfect your pLVX-TRE3G-GOI into an easy-to-transfect cell line such as HeLa or HEK 293, or your target cell line, and test for GOI induction with Tet-Express. You will need an appropriate gene-specific assay to test for induction, such as:

- Western blot
- Northern blot
- qRT-PCR
- Gene-specific functional assay

A. Materials Required

1. pLVX-TRE3G Vector containing your gene of interest (Section V)
2. Tet-Express
3. Host cell line
4. Xfect Transfection Reagent (Section III.I)
5. Mammalian cell culture supplies (Section III.D)
6. Tet Approved FBS (Section III.A)

B. Protocol

1. Perform an overnight transfection of pLVX-TRE3G-GOI (GOI = gene of interest) into your target cells (in a 6-well plate) using Xfect transfection reagent. Follow the Xfect Transfection Reagent Protocol-At-A-Glance. (Locate this protocol by searching at takarabio.com/manuals). You can use pLVX-TRE3G-Luc as a positive control.
 - Transfect 5 µg of pLVX-TRE3G-GOI (or pLVX-TRE3G-Luc) into each well.
 - We recommend performing the test in duplicate with negative controls.
2. The next day, prepare 2 vials as follows:

Tube 1		Tube 2	
3 µl	Tet-Express Transactivator	2.5 µl	Intensifier Reagent
97 µl	serum-free medium (e.g., DMEM)	97.5 µl	serum-free medium (e.g., DMEM)
100 µl	Total Volume	100 µl	Total Volume

3. Vortex each tube to mix.
4. Replace the serum-containing media that your target cells are growing in with 1 ml of serum-free media (e.g., DMEM, just like in the Tet-Express/Intensifier Reagent mixture).

NOTE: You can add Tet-Express in the presence of serum, but the transduction level will be reduced.

5. Combine the contents of both vials and add the entire 200 µl dropwise to your overnight transfected cells. Rock the plate gently to mix and distribute Tet-Express evenly across the cells.
6. Incubate for 1 hr and then add 1 ml of serum-containing media.
7. Incubate for 2 hr to overnight, and assay for induction of your gene of interest (compare to pLVX-TRE3G-GOI-transfected cells that have not been exposed to Tet-Express).

VII. Producing Lentivirus from the Lenti-X Vectors

Follow the **Lenti-X Packaging Single Shots (VSV-G) Protocol-At-A-Glance**. (Locate this protocol by searching at takarabio.com/manuals).

VIII. Lentivirus Titration

A. Summary

1. Instant Qualitative Titer Test

You can quantify your lentivirus stock in ten minutes with **Lenti-X GoStix Plus** (Cat. Nos. 631280, 631281) and the related smartphone app. The GoStix detect lentiviral p24 in only 20 µl, and can be used to determine whether virus production is within a usable range or for selecting the best time to harvest your virus. A 3-prep sample is supplied for free with many of Takara Bio's Lenti-X systems. Visit takarabio.com/gostixhelp for details.

2. Quantitative Titer Test

- a. **Determining the viral titer** is necessary to obtain the following information:
 - Confirmation that viral stocks are viable
 - The proper transduction conditions for your particular cell type by adjusting the MOI for the desired transduction efficiency. MOI = No. of infectious virus particles per target cell
 - The maximum number of target cells that can be transduced by a given virus volume.
- b. To transduce using a known multiplicity of infection (MOI), it is necessary to titrate your lentiviral stocks. We recommend the **Lenti-X qRT-PCR Titration Kit** (Cat. No. 631235) or **Lenti-X p24 Rapid Titer Kit** (Cat. No. 632200) for very rapid quantitative titrations of virus stocks (~4 hr), or a standard method that relies on infection.
- c. The **standard viral titration protocol** consists of infecting cells with serial dilutions of the stock, selecting for stable transductants with antibiotic and counting the resulting cell colonies (Section VIII.B).
 - Freshly harvested virus can be titered immediately, or frozen in aliquots at –80°C and then titrated. Note that each freeze-thaw cycle can reduce the functional titers of infectious virus by up to 2–4 fold.
 - Absolute titers will depend heavily on the cell type used for titration, and there may be significant differences between the titer values determined in cells typically used for lentiviral titration (i.e. HT-1080) and the number of target cells transduced by the titered virus. However, titrations serve to determine the relative virus content of different viral stocks prepared from different vectors.

B. Protocol: Determining Viral Titer by Colony Formation

NOTE: This protocol can be completed in 7–14 days.

1. Plate HT-1080 cells (or other) in 6-well plates the day before performing the titration infections. Plate 2×10^5 cells/well, in 2 ml of medium. Allow at least one well to be used as a “no infection” control.
2. Prepare 20 ml of complete medium and add 60 μ l of 4 mg/ml Polybrene. This will be diluted 3-fold for a final Polybrene concentration of 4 μ g/ml.

NOTE: Polybrene is a polycation that reduces charge repulsion between the virus and the cellular membrane. The optimum final concentration of Polybrene may be determined empirically but generally falls within a range of 2–12 μ g/ml. Excessive exposure to Polybrene (>24 hr) can be toxic to cells.

3. Prepare filtered viral supernatant from packaging cells (Section VII). This is the virus stock.
4. Prepare six 10-fold serial dilutions of the virus stock as follows:
 - a. Add 1.35 ml of medium containing Polybrene (Step 2) to each of six sterile and numbered 1.5 ml microfuge tubes.
 - b. Add 150 μ l of the virus stock (Step 3) to the tube 1. Mix.
 - c. Transfer 150 μ l tube 1 to tube 2 and mix. Continue making serial dilutions by transferring 150 μ l from each successive dilution into the next prepared tube.
5. Infect the HT-1080 cells by adding 1 ml of each viral dilution (Step 4) to each appropriate well. The final Polybrene concentration will be 4 μ g/ml in ~3 ml. Centrifuge the cultures to improve infection efficiency*.

***NOTE: CULTURE CENTRIFUGATION INCREASES INFECTION EFFICIENCY.** Centrifuging the plate at 1,200 x g for 60–90 min at 32°C can significantly increase infection efficiency. A room temperature centrifuge is acceptable if a 32°C unit is not available.

6. After infecting for 8–24 hours, remove supernatants and subject the cells to puromycin selection using the selection concentrations that are optimal for your cell line (Section III.B).
7. Allow colonies to form for 7–14 days. Stain the colonies with 1% crystal violet solution (in 10% ethanol) and count.
8. The titer of virus corresponds to the number of colonies generated by the highest dilution, multiplied by the dilution factor. For example, the presence of 4 colonies in the 10^6 dilution would represent a viral titer of 4×10^6 colony forming units.

IX. Transducing Target Cells with TRE3G Lentiviruses and Inducing Expression with Tet-Express

A. Protocol: Transducing Target Cells with Lenti-X TRE3G Lentiviruses

NOTE: This protocol can be completed in 2–3 days.

1. Plate target cells in complete growth medium 12–18 hr before transduction.
2. Thaw aliquots of your LVX-TRE3G-GOI lentiviral stock, or use filtered virus stocks freshly prepared from packaging cells (Section VII). Mix gently, but do not vortex.
3. Add Polybrene to the cell cultures to obtain the desired final concentration during the transduction step (e.g., 4 µg/ml).

NOTE: Lenti-X Accelerator (Cat. Nos. 631256, 631257 & 631254) and **RetroNectin** (Cat. Nos. T110A, T100B & T100A) may be used as transduction enhancers instead of Polybrene (see Section III.H).

4. Test different amounts of your lentiviral supernatant, but make certain that the total volume of supernatant used makes up no more than 1/3 the final volume of culture medium used in the transduction. Centrifuge the cultures to improve transduction efficiency (see Section VIII.B).
5. Transduce the cells for 8–24 hr. If you are concerned that exposure to either the Polybrene or to the viral supernatant (which contains medium conditioned by the packaging cells) may adversely affect your target cells, limit the transduction to 6–8 hr.
6. Remove and discard the virus-containing medium and replace it with fresh growth medium, with or without Tet-Express, as appropriate (Section IX.B). Alternatively, passage the cultures and subject the cells to selection using puromycin to create a TRE3G stable cell line. See Appendix C for information on freezing and thawing cell line stocks.

B. Protocol: Inducing Expression with Tet-Express

1. Seed your LVX-TRE3G-GOI-transduced cells (Section IX.A) into two wells of a 6-well plate.
2. The next day, prepare 2 vials as follows:

Tube 1		Tube 2	
3 µl	Tet-Express Transactivator	2.5 µl	Intensifier Reagent
97 µl	serum-free medium (e.g., DMEM)	97.5 µl	serum-free medium(e.g., DMEM)
100 µl	Total Volume	100 µl	Total Volume

3. Vortex each tube to mix.
4. Replace the serum-containing media that your target cells are growing in with 1 ml of serum-free media (e.g., DMEM, just like in the Tet-Express/Intensifier Reagent mixture).

NOTE: You can add Tet-Express in the presence of serum, but the transduction level will be reduced.

5. Combine the contents of both vials and add the entire 200 µl dropwise to one well of your transduced cells. Leave one well untreated as a negative control. Rock the plate gently to mix and distribute Tet-Express evenly across the cells.
6. Incubate for 1 hr and then add 1 ml of serum-containing media.

7. Incubate for 2 hr to overnight, and assay for induction of your gene of interest (compare to pLVX-TRE3G-GOI-transfected cells that have not been exposed to Tet-Express).

X. References

Takara Bio's Tet Systems were developed in cooperation with Dr. Bujard and his colleagues at the Center for Molecular Biology in Heidelberg (ZMBH) and in Dr. Wolfgang Hillen's laboratory at the University of Erlangen, Germany. Additional background information on Tet-regulated gene expression systems and an extensive bibliography are available at the website maintained by TET Systems: <http://www.tetsystems.com> (Please note that Takara Bio is not responsible for the information contained on this website.)

Freshney, R.I. (2005). *Culture of Animal Cells: A Manual of Basic Technique, 5th Edition* (Wiley-Liss, Hoboken, NJ).

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Appendix A. Troubleshooting Guide

Table 2. Troubleshooting Guide for the Lenti-X Tet-Express Inducible Expression System

Problem	Possible Explanation	Solution
A. Vector Cloning		
Plasmid is difficult to grow or clone	Some viral vectors may undergo rearrangement between the 5' and 3' LTRs when propagated in less-than-optimal <i>E. coli</i> host strains	Use Stellar Competent Cells (Cat. No. 636763) to produce high DNA yields and to minimize the potential for DNA rearrangements.
B. Lenti-X 293T Packaging Cells		
Poor viability upon thawing	Improper thawing techniques	Use thawing procedure in Appendix C, and/or consult the Lenti-X 293T Cell Line Protocol-at-a-Glance
	Incorrect culture medium	Use DMEM with additives listed in Section III.D. Use 10% Tet System Approved FBS (Tc-free).
	Improper tissue culture plasticware	Use collagen I-coated plates to aid cell adherence during initial seeding.
Slow growth	Incorrect culture medium	Use DMEM with additives listed in Section III.D. Use 10% Tet System Approved FBS (Tc-free).
Cells do not attach to plate	Improper tissue culture plasticware	Use collagen I-coated plates to aid cell adherence during initial seeding.
Cells appear morphologically different	Passage of cell culture is too high (old cells)	Thaw/purchase new aliquot of Lenti-X 293T cells.
C. Virus Production		
Poor transfection efficiency (as determined by GOI or marker expression in the Lenti-X 293T cell line)	Cells plated too densely	Plate 4–5 x 10 ⁶ cells/100 mm plate, or fewer if the cells divide rapidly. Use at 50–80% confluency. See Section VII.
	Transfection is toxic to cells	Use the optimized conditions provided in Section VII.
	Cells harvested or analyzed too soon after transfection	Wait 48 hr after transfection for maximal expression of GOI or marker to determine efficiency.
Low titers (<10 ⁵ cfu/ml)	Serum in medium contains tetracycline contaminants, which can interfere with the expression of viral proteins, resulting in lower titers	Use Tet System Approved FBS (Cat. Nos. 631101 & 631106) in the 293T culture medium.
	Virus was harvested too early	Harvest virus 48–72 hr after the start of transfection.
	Vector is too large	The limit for efficient packaging function is 9.7 kb from the end of the 5'-LTR to the end of the 3'-LTR
	Polybrene is missing or at suboptimal concentration	Add Polybrene (4 µg/ml) during transduction or optimize the concentration (2–12 µg/ml)
	Virus was exposed to multiple freeze-thaw cycles	Each cycle reduces titer by approximately 2–4 fold. Limit the number of freeze-thaws.
	Suboptimal selection procedure during titration	Perform an antibiotic kill curve on the cell line prior to using it for titration.

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Problem	Possible Explanation	Solution	
D. Transduction of Target Cells			
Poor transduction efficiency	Low titer	See Section C or use the Lenti-X Concentrator (Section III.G) to increase your available titer up to 100-fold without ultracentrifugation.	
	Poor transfection efficiency	Follow the protocol in Section VII.B. Be sure to use 5 µg of transfection-grade plasmid.	
	Low viability of target cells during transduction	Optimize culture conditions for target cells prior to infection	
		Packaging cell line-conditioned media may affect cell growth; dilute viral supernatant or shorten exposure time to viral supernatant. Consider using RetroNectin Reagent and the RetroNectin-Bound Virus transduction protocol or purify your virus prior to transduction using the Lenti-X Maxi Purification Kit (Cat. Nos. 631233 & 631234).	
	Excessive exposure to Polybrene: optimize amount (titrate) or shorten exposure time to viral supernatant		
Viral supernatant contains transduction inhibitors	Use RetroNectin Reagent or RetroNectin-coated plates in the RetroNectin-Bound Virus transduction protocol, which allows virions to bind the RetroNectin substratum and be washed free of inhibitors prior to target cell infection; or, purify your virus prior to transduction using the Lenti-X Maxi Purification Kit (Cat. Nos. 631233 & 631234).		
E. Inducing Expression			
Low fold induction (ratio of maximal to basal expression of the GOI)	Cells were harvested and analyzed too soon or too late.	Harvest and analyze cells between 2–24 hr after addition of Tet-Express (2 hr may be too soon to detect some proteins). The detection time is also dependent on the assay sensitivity. Typical expression peaks at 24 hr and starts to decline after 48 hr.	
	Poor infection efficiency	<ul style="list-style-type: none"> • Confirm virus titers using a titration kit (Section III.F) • Increase amount of virus applied to target cells • Optimize density of cells when transducing 	
	Poor target cell viability	<ul style="list-style-type: none"> • Optimize passage number of target cells. • Optimize target cell culture conditions. • Optimize tissue culture plasticware 	
	The FBS used in the cell culture medium contains tetracycline derivatives.	Use our Tet System Approved FBS (Section III.A), which was functionally tested with our double-stable CHO-AA8-Luc Tet-Off Control Cell Line.	
Decrease in fold induction after several passages	The appropriate antibiotics are missing from the cell culture medium.	Maintain optimal antibiotic concentrations (Section III.B).	
or Loss of inducibility after passaging of a (previously frozen) double-stable cell line.	Mixed cell population	Reselect the current cell line through single colony selection using a selective concentration of puromycin (Section IX).	

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Problem	Possible Explanation	Solution
F. Establishment of Stable Cell Lines		
Untransduced cells do not die at the high antibiotic concentration established via titration in Section III.B	<ul style="list-style-type: none"> The cells have not been recently passaged, so they remain well-attached to the plate surface even when they are dead. You have achieved 100% transduction efficiency. 	To determine the appropriate antibiotic concentration, use cells that have been split within the last 2–3 days.
There are no surviving cells after transduction followed by selection	The antibiotic concentration which caused massive cell death when determining the appropriate dose via titration could be too high.	Use a lower antibiotic concentration for selection of stably transfected cell clones.
Poor cell viability	Cells were not properly frozen.	See Appendix C, Section A.
	Cells were not properly thawed.	See Appendix C, Section B.
G. Detection and Inhibition of Expression		
No detectable GOI expression by Western Blot.	Low sensitivity of detection method.	Check sensitivity of primary and secondary antibodies. Analyze GOI expression by qRT-PCR, using different sets of primers to ensure optimal detection of GOI expression.
Continuous GOI/Fluorescent Protein expression after the removal of Tet-Express	Depending on the stability of the protein, it may persist in the cell in the absence of gene induction and de novo synthesis of GOI mRNA. Fluorescent proteins tend to have long half-lives.	Upon degradation, GOI/Fluorescent Protein expression will not be detectable in cells in the absence of induction. For faster degradation of an inducible GOI, use pTRE-Cycle Vectors (see takarabio.com).

Appendix B: Lenti-X Tet-Express System Vector Information

The Lenti-X Tet-Express Inducible Expression System (Section II) contains two vectors (Figure 5), a response vector (pLVX-TRE3G) and a control response vector (pLVX-TRE3G-Luc). For complete descriptions of the vectors provided with each system, refer to the enclosed Certificate of Analysis, which is also available at takarabio.com

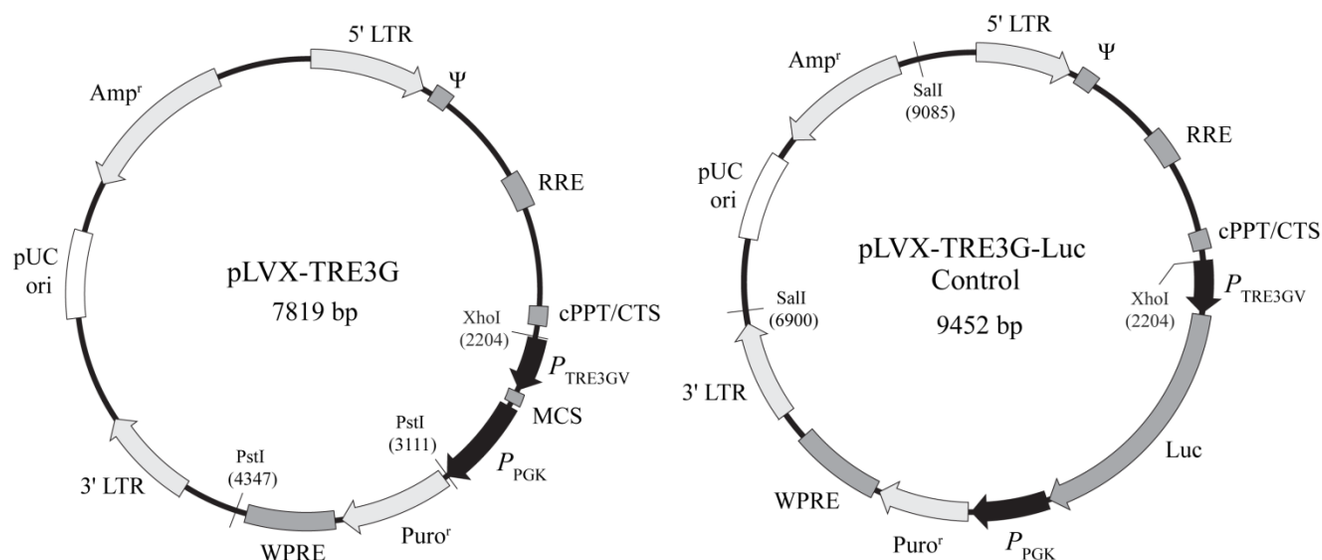


Figure 4. pLVX-TRE3G and pLVX-TRE3G-Luc Vector Maps.

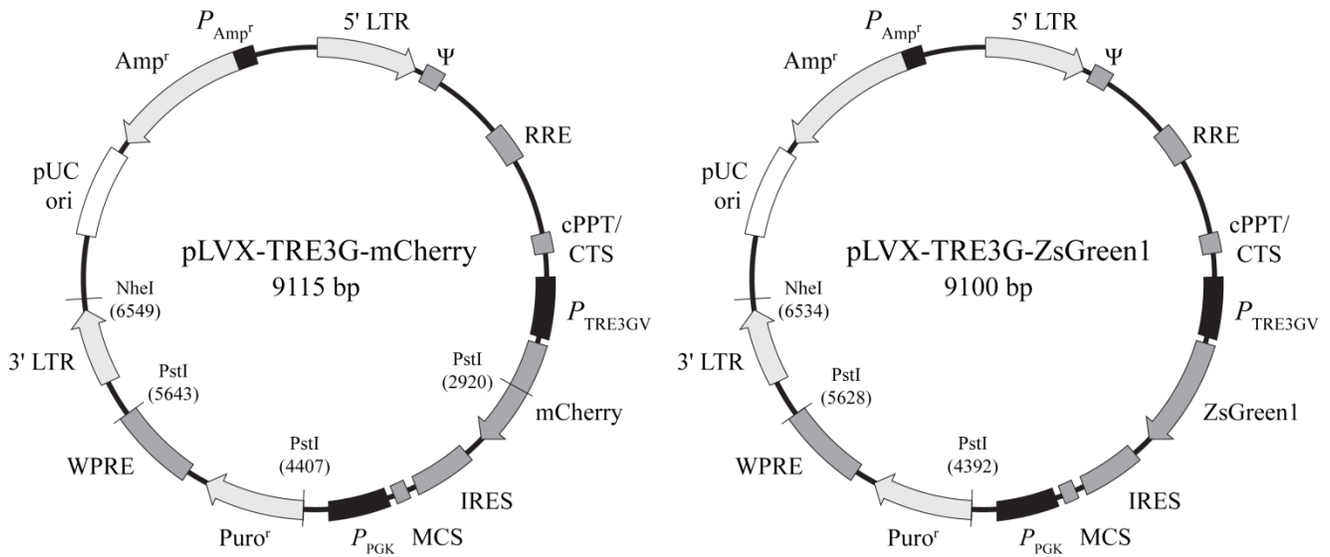


Figure 5. pLVX-TRE3G-mCherry Vector and pLVX-TRE3G-ZsGreen1 Vector Maps.

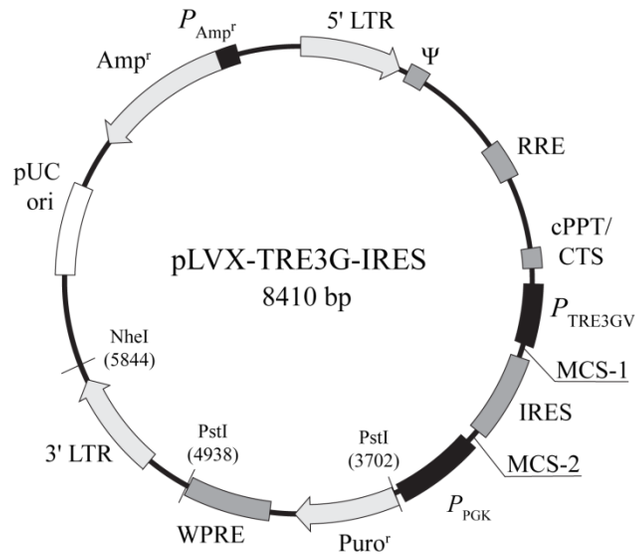


Figure 6. pLVX-TRE3G-IRES Vector Map.

Appendix C: Preparing and Handling Cell Line Stocks

A. Protocol: Freezing Cell Line Stocks

Once you have created and tested your TRE 3G cell line, you may choose to prepare multiple frozen aliquots to ensure a renewable source of cells, according to the following protocol:

1. Expand your cells to multiple 10 cm dishes or T75 flasks.
2. Trypsinize and pool all of the cells, then count the cells using a hemocytometer.
3. Centrifuge the cells at 100 x g for 5 min. Aspirate the supernatant.
4. Resuspend the pellet at a density of at least $1-2 \times 10^6$ cells/ml in freezing medium. Freezing medium can be purchased from Sigma (Cat. Nos. C6164 & C6039), or use 70–90% FBS, 0–20% medium (without selective antibiotics), and 10% DMSO.
5. Dispense 1 ml aliquots into sterile cryovials and freeze slowly (1°C per min). For this purpose, you can place the vials in Nalgene cryo-containers (Nalgene, Cat. No. 5100-001) and freeze at –80°C overnight. Alternatively, place vials in a thick-walled styrofoam container at –20°C for 1–2 hr. Transfer to –80°C and freeze overnight.
6. The next day, remove the vials from the cryo-containers or styrofoam containers, and place in liquid nitrogen storage or an ultra-low temperature freezer (–150°C) for storage.
7. Two or more weeks later, plate a vial of frozen cells to confirm viability.

B. Protocol: Thawing Cell Line Frozen Stocks

To prevent osmotic shock and maximize cell survival, use the following procedure to start a new culture from frozen cells:

1. Thaw the vial of cells rapidly in a 37°C water bath with gentle agitation. Immediately upon thawing, wipe the outside of the vial with 70% ethanol. All of the operations from this point on should be carried out in a laminar flow tissue culture hood under strict aseptic conditions.
2. Unscrew the top of the vial slowly and, using a pipet, transfer the contents of the vial to a 15 ml conical centrifuge tube containing 1 ml of prewarmed medium (without selective antibiotics such as G418). Mix gently.
3. Slowly add an additional 4 ml of fresh, prewarmed medium to the tube and mix gently.
4. Add an additional 5 ml of prewarmed medium to the tube and mix gently.
5. Centrifuge at 100 x g for 5 min, carefully aspirate the supernatant, and GENTLY resuspend the cells in complete medium without selective antibiotics. (This method removes the cryopreservative and can be beneficial when resuspending in small volumes. However, be sure to treat the cells gently to prevent damaging fragile cell membranes.)

- Mix the cell suspension thoroughly and add to a suitable culture vessel. Gently rock or swirl the dish/flask to distribute the cells evenly over the growth surface and place in a 37°C humidified incubator (5–10% CO₂ as appropriate) for 24 hr.

NOTE: For some loosely adherent cells (e.g. HEK 293-based cell lines), we recommend using collagen-coated plates to aid attachment after thawing. For suspension cultures, suspend cells at a density of no less than 2 x 10⁵ cells/ml.

- The next day, examine the cells under a microscope. If the cells are well-attached and confluent, they can be passaged for use. If the majority of cells are not well-attached, continue culturing for another 24 hr.

NOTE: Note: For some loosely adherent cell lines (e.g., HEK 293-based cell lines), complete attachment of newly thawed cultures may require up to 48 hr.

- Expand the culture as needed. The appropriate selective antibiotic(s) should be added to the medium after 48–72 hr in culture. Maintain Cell Lines in complete culture medium containing a maintenance concentration of puromycin, as appropriate (Section III.B).

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