# **Product Description**

TribioScience's Lentivirus (LV) qPCR Titer Kit incorporates TribioScience FAST technology paradigm and probe Taqman qPCR technology for an accurate, one-step, and time-saving qPCR titer determination of lentivirus. The probe is labeled with Fluorescence FAM. The entire procedure, which does not require an additional reverse transcription step, can be completed within 2 hours. With its time-saving 1-minute enzyme activation technology, specificity, and sensitivity, the performance of TribioScience's Lentivirus qPCR Titer Kit is unmatched by similar kits currently on the market.

# **Kit Content for 100RXN**

| Component         | Volume | Part Number |
|-------------------|--------|-------------|
| FAST Lysis Buffer | 1.0 mL | LV001       |
| Fast qPCR mix     | 1.0 mL | LV002       |
| Primer-Probe Mix  | 0.3 mL | LV003       |
| LV Control DNA    | 30 µL  | LV004       |
| PCR Buffer        | 1.0 mL | LV005       |

Note: The probe is labelled with Fam.

# Storage and Shelf-life

Store at -25°C to -15°C immediately upon arrival. Minimize the number of freeze-thaw cycles to ensure superior performance. The Kit is stable for one (1) year from the date of arrival.

## Protocol

- 1. Viral Sample Preparation
- **A. Purified viral particles:** Bring viral particles to 10<sup>7</sup> IU/ml with PBS.
- **B.** Virus producing cell cultures: centrifuge cell culture medium at 2000g for 5 minutes to sediment suspended cells and cell debris. Use supernatant for the next step.

## 2. Viral Lysis

Add 2 mL of the sample preparation to 18 mL of FAST Lysis Buffer (1:10 dilution). Pipette gently up and down to mix and incubate at room temperature (23°C) for 3 minutes. Use the lysed sample for the reaction set up in Step4. *Note: the viral sample has been diluted 1/10, thus take this dilution factor into consideration when calculating the final titer*.

3. Standard Control DNA Preparation: The LV Control DAN is  $1 \times 10^{10}$  IU/mL. Perform 5 serial dilutions of the Standard Control DNA at 10-fold manner by diluting 2 mL Standard DNA into 18 mL PCR buffer (Nuclease-Free water) in each concentration. Dilutions 1/10 to 1/100,000 will be used for generating the standard curve. The standard preparation is listed in the table 1:

#### **Table 1: Standard Preparation**

| Dilution | Virus Titer (IU/ml) |  |
|----------|---------------------|--|
| 1/10     | $2x10^{9}$          |  |
| 1/100    | $2x10^{8}$          |  |
| 1/1000   | $2x10^{7}$          |  |
| 1/10000  | 2x10 <sup>6</sup>   |  |
| 1/100000 | $2x10^{5}$          |  |

4. **qPCR Preparation:** All reactions are set up on ice in duplications. The reaction volume is 20 mL. The detail listed in Table 2.

#### Table 2: qPCR Reaction Setup

| Component                   | Volume (µL) |  |
|-----------------------------|-------------|--|
| Fast qPCR mix               | 7.0         |  |
| Primer-Probe mix            | 2.0         |  |
| DNA Sample or Standard, NTC | 2.0         |  |
| PCR Buffer                  | 9.0         |  |
| Final Volume                | 20          |  |

Note: PCR Buffer is used as NTC.

**5. qPCR Running Parameters:** Program the qPCR instrument as follows, and Color Channel is FAM:

#### **Table 3: PCR Conditions**

| Purpose           | Temperature (°C) | Time | Cycle(s) |
|-------------------|------------------|------|----------|
| Enzyme            | 95               | 60S  | 1        |
| Activation        |                  |      |          |
| Denaturation      | 95               | 10S  | 40       |
| Priming/Extension | 60               | 60S  |          |

## 6. Titer Calculation

Plot Ct value (Y-axis, Linear scale) vs. Virus titer (X-axis, Logarithmic scale). Generate a logarithmic regression using the five (5) Standard Control DNA dilutions to determine the unknown virus sample titer using y = mIn(x) + b for the trendline equation. The R<sup>2</sup> value should be >0.95 to justify the proper assay setup. Note to include the dilution factor in the final calculation (i.e. if you diluted your purified viral samples 1/100 in step 1 and 1/10/ in Step 2, then the titer of the unknown sample should be multiplied by a factor of 100x10).

Use the Ct values to calculate the viral titer of the viral sample with the following formula:

Titer of unknown sample  $(IU/ml) = e^{(Ctx-b)/m}$ , Where m is the slope of the line, and b is the y-intercept.

Example: Trendline equation is y = -1.349In(x) + 40.898; Ct if unknown sample = 16.98

Virus titer  $(IU/ml) = e^{(16.98-40.898)/-1.349} = 5.01 \times 10^7 IU/ml$ 

Note: Remember to include the dilution factor in the calculation if dilutions of viral particles had been performed.