

Chemi - Trans™ Polyfectin DNA Transfection Reagent

(Cat #: T001)

Introduction:

The Chemi-Trans™ Polyfectin DNA Transfection Reagent is formulated as a biodegradable Polymer-based DNA delivery tool that performs well for DNA transfecting in a variety of commonly used and hard-to-transfect mammalian cell lines. A remarkable feature of Chemi-Trans™ Polyfectin Reagent is the rapid and complete degradation of polymer after transfection complex by endocytosis, leading to much less cytotoxicity.

Advantages:

- ❖ Bio-degradable and low cytotoxicity.
- ❖ Excellent for long DNA transfection.
- ❖ Easy to use, forms a DNA complex in just five minutes.
- ❖ Compatible with serum and antibiotics in culture medium.

Table 1: Product Package & Storage

Cat #	Product Name	Volume	Storage
T001	Chemi -Trans™ Polyfectin DNA Transfection Reagent	1.0 mL	2 ~ 8 °C, stable for up to 12 months when stored appropriately. (DO NOT FREEZE.)

Important Guidelines for Transfection:

- (1) For maximum transfection efficiency, using serum-free medium (such as serum-free DMEM with High Glucose. DO NOT use Opti-MEM Medium, it contains serum and will disrupt transfection complex) to dilute Chemi - Trans™ Polyfectin Reagent and DNA is a must.
- (2) While the standard protocols for DNA transfection being given below, optimization is often needed for maximal transfection efficiency.

Standard Protocol for DNA Transfection of Adherent Cells

Step I . Cell Seeding:

Cells should be plated 18 to 24 hours prior to transfection so that the monolayer cell density reaches to 70 ~ 80% confluency at the time of transfection. Complete culture medium with serum and antibiotics is freshly added to each well 30 ~ 60 min before transfection.

Step II . DNA Transfection Protocol

Use the following procedure to transfect DNA into mammalian cells in a 6-well format. For other formats, please refer to **A Guideline for DNA transfection (Table 2)**. All amounts and volumes are given on a per well basis. For each transfection sample, prepare complexes as follows:

- a. Dilute 2.0 µg Endotoxin-free plasmid DNA in 100 µL of serum-free DMEM with High Glucose. Vortex to mix gently but thoroughly.

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- b.** Mix Chemi - Trans™ Polyfectin Reagent gently before use, then dilute 6.0 µL of Chemi - Trans™ Polyfectin in 100 µL of serum-free DMEM with High Glucose.
- c.** Add the diluted Chemi - Trans™ Reagent immediately to the diluted DNA mixture (prepared in Step a, **do not mix the solutions in reverse.**) all **(1:1 ratio)** after the diluted formulation be incubated for 5 minutes, and mix by pipetting up and down. Incubate for 10 ~ 15 minutes at room temperature to let transfection complex form well.
- d.** Add the 200 µL of complexes to each well containing cells and medium. Mix gently by rocking the plate back and forth. Incubate cells at 37°C in a CO₂ incubator.
- e.** Remove the Chemi - Trans™ Reagent/DNA complex-containing medium and replace with pre-warmed fresh complete serum/antibiotics containing medium 4 ~ 6 hours post transfection when it is necessary. Then, to measure the gene silencing by qRT-PCR and Western Blotting respectively 24 ~ 72 hours post transfection.

Table 2: A Guideline for DNA transfection per cell culture vessel

Culture Vessel	Growth Medium (mL)	Serum - Free Medium (µL)	DNA (µg)	Chemi - Trans™ Polyfectin Reagent (µL)
	Volume used per well			
96-well	0.1	2 × 5	0.1	0.15~0.4
24-well	0.5	2 × 25	0.5	0.75~2.0
12-well	1.0	2 × 50	1.0	1.5~4.0
6-well	2.0	2 × 100	2.0	3.0~8.0

*[1] We strongly suggest that keep the concentration of plasmid DNA be 0.5 ~ 2.0 µg/µL, and the Endotoxin-free plasmid is extremely important for a successful transfection.

[2] To obtain a robust transfection efficiency and low cytotoxicity, optimize transfection conditions by varying cell density as well as DNA and Chemi - Trans™ Polyfectin Reagent concentrations. Ensure that cells are greater than 90% confluent and vary DNA (µg): Chemi - Trans™ Polyfectin Reagent (µl) ratios from 1:1 to 1:4

[3] You may perform a rapid 96-well plate transfections by plating cells directly into the transfection complexes. Prepare complexes in the plate and directly add cells drop wise at twice the cell density as in the basic protocol in the complete growth medium. Cells will adhere as usual in the presence of complexes.