

#### This product is for laboratory research ONLY and not for diagnostic use

#### Introduction:

PepMute<sup>™</sup> Plus Reagent is an upgraded version from PepMute<sup>™</sup> siRNA transfection reagent. With addition of several pre-screened hydrophobic groups to its peptide backbone, PepMute<sup>™</sup> Plus Reagent gains self-assembly capacity when binding nucleic acids, making PepMute<sup>™</sup> Plus Reagent a versatile and most powerful gene delivery tool. PepMute<sup>™</sup> Plus Reagent have been validated to effectively and reproducibly transfect single siRNA, And co-transfect DNA/siRNA to variety of mammalian cells.

#### **Important Guidelines for Transfection:**

- PepMute<sup>™</sup> Plus reagent was formulated as a powerful siRNA delivery tool. For most adherent cell lines and primary cells, siRNA at ~5.0 nM is basically sufficient to obtain up to 90% gene silencing. For hard-to-transfect cells, we recommend using a final siRNA concentration of 50 nM.
- While the standard protocols for siRNA transfection and siRNA/DNA co-transfection are being given below, optimization is often needed for maximal gene silencing.

#### PART I. Standard siRNA Transfection of Adherent Cells

### Step I. Preparation of Working Solution of PepMute™ Transfection Buffer:

PepMute<sup>™</sup> Transfection Buffer (5x) is provided as 5x concentrated stock solution. To make working solution, dilute one part of the stock solution with 4 parts of ddH<sub>2</sub>O into a sterile bottle. The working solution is table at 4 °C~RT for 12 months.

#### Step II. Cell Seeding:

Cells should be plated 18 to 24 hours prior to transfection so that the monolayer cell density reaches to the optimal  $\sim$ 50% confluency at the time of transfection. Complete culture medium with serum and antibiotics is freshly added to each well 30 $\sim$ 60 minutes before transfection.

**Note:** PepMute<sup>™</sup> Plus reagent is NOT interfered by serum and antibiotics, therefore serum and antibiotic containing medium can be used during the entire experiment.

## Table 1. A Guideline for siRNA transfection per cell culture vessel

Culture Dish	Growth Medium (ml)	Transfection Buffer (µL)	siRNA (pmoles) Final 5.0 or <u>50</u> nM	PepMute™ Plus (µL)
24-well	0.5	50	2.5 / <u><b>25</b></u>	1.2 ~ <u><b>2.0</b></u>
12-well	0.75	75	3.75 / <u><b>38</b></u>	2.0 ~ <u>3.3</u>
6-well	1.0	100	5.0 / <u><b>50</b></u>	2.4 ~ <u>4.0</u>
60 mm	3.0	300	15 / <b>150</b>	7.2 ~ <u>12</u>
10 cm / Flask 75	8.0	800	40 / <b><u>400</u></b>	20 ~ <u>33</u>

#### Step III. siRNA Transfection Protocol:

For optimal siRNA-mediated silencing, we recommend using  $1 \sim 100$  nM siRNA. As a starting point, we recommend using 5.0 nM siRNA which usually gives satisfactory silencing result for most adherent cell lines or primary cells. For hard-to-transfection cells, we recommend using a final siRNA concentration of 50 nM. (bold & underlined in **Table 1**).

The following conditions are given per well in a 6 well plate. For other culture format, please refer to **Table 1**.

- For each well, add 1.0 ml of complete medium with serum and antibiotics freshly 30~60 minutes before transfection.
- Dilute 5.0 or <u>50</u> pmoles siRNA (final concentration of 5.0 or <u>50</u> nM respectively per well) into 100 µl of working solution of PepMute<sup>™</sup> Transfection Buffer prepared in <u>Step I</u>. Pipette up and down to mix.
- Note: For maximum gene silencing, dilute siRNA and PepMute<sup>™</sup> Plus reagent with PepMute<sup>™</sup> Transfection Buffer (1x). We strongly suggest reconstituting siRNA stock solution at 10.0 µM, so add 0.5 or 5.0 µl siRNA stock solution per well of 6well plate to make final 5.0 and 50 nM siRNA respectively.
- Add 2.4 µl or 4.0 µl (for hard-to-transfect cells, bold and underlined in <u>Table 1</u>) PepMute<sup>™</sup> Plus reagent, mix by pipetting up and down.
- Incubate for  ${\sim}15$  minutes at RT to let transfection complex form.

# Note: Never keep the complex longer than 30 minutes.

- Add the transfection mix to the cells drop wise. Gently rock the plate back and forth and return the plate to CO<sub>2</sub> incubator.
- Gene silencing is usually measured 24~78 hours post transfection.

### PART II. A Standard Protocol for DNA/siRNA Cotransfection

# Step I. Preparation of Working Solution of PepMute<sup>™</sup> Transfection Buffer:

PepMute<sup>™</sup> Transfection Buffer (5x ) is provided as 5x concentrated stock solution. To make working solution, dilute one part of the stock solution with 4 parts of ddH<sub>2</sub>O into a sterile bottle. The working solution is table at 4 °C~RT for 12 months.



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#### Step II. Cell Seeding:

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

Note: PepMute<sup>™</sup> Plus reagent is NOT interfered by serum and antibiotics, therefore serum and antibiotic containing medium can be used during the entire experiment.

# Table 2. A Guideline for DNA & siRNA Co-transfection Per Cell Culture Vessel

Culture Dish	Growth Medium (ml)	Transfection Buffer (µL)	Plasmid DNA (µg)	siRNA (pmoles) Final 5.0 nM	PepMute™ Plus (μL)
24-well	0.5	50	0.25	2.5	1.5
12-well	0.75	75	0.375	3.25	2.25
6-well	1.0	100	0.5	5.0	3
60 mm	3.0	300	1.5	15	9
10 cm / flask 75	8.0	800	4.0	40	24

## Step III. DNA & siRNA co-transfection protocol:

For DNA/siRNA co-transfection experiment, we recommend using 0.25~0.5  $\mu$ g DNA and 1~20 nM siRNA per well in a 6-well plate. As a starting point, we recommend using 0.5  $\mu$ g DNA and 5.0 pmoles siRNA (final concentration 5.0 nM) per well of a 6-well plate which usually give satisfactory knockdown effect. The following conditions are given per well of a 6 well plate. For other culture format, please refer to **Table 2**.

- For each well, add 1.0 ml of complete medium with serum and antibiotics freshly 30~60 minutes before transfection.
- Dilute 0.5  $\mu g$  DNA and 5.0 pmoles siRNA (final 5.0 nM) into 100  $\mu l$  of working solution of PepMute^ $^{\rm TM}$  Transfection Buffer. Vortex to mix followed by brief spin to bring drops to the bottom of the tube.

Note: For optimal transfection efficiency and maximum gene silencing, PepMute<sup>™</sup> Transfection Buffer is a must for diluting siRNA/DNA and PepMute<sup>™</sup> reagent. We strongly suggest preparing siRNA stock solution at 5.0 µM, so add 1.0 µl siRNA stock solution per well of 6-well plate to make final 5.0 nM of siRNA.

- Add 3  $\mu I$  PepMute  $^{\rm TM}$  Plus reagent immediately, mix by pipetting up and down.
- Incubate for ~15 min at RT to let transfection complex form. **Note**: Never keep the complex longer than 30 minutes.
- Add the transfection complex to the cells drop wise.

- Gently rock the plate back and forth and return the plate to the incubator.
- Replace transfection medium by cell growth medium
   ~5 hours after transfection when necessary.
- Gene silencing is usually measured 24~72 hours post transfection.

**Storage:** PepMute<sup>TM</sup> Plus siRNA Transfection Reagent is stable for up to 12 months at 4 °C. This item shipped at ambient temperature