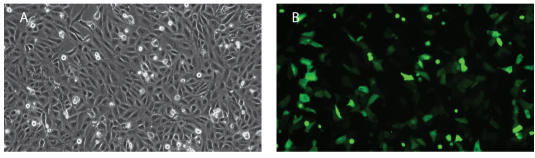
# Amaxa<sup>®</sup> Cell Line Nucleofector<sup>®</sup> Kit V

## For Vero

African Green Monkey kidney cells; epithelial cells

Example for Nucleofection® of Vero cells



Vero cells were transfected with the Cell Line Nucleofector® Kit V, Program V-001 and 2 µg of pmaxGFP® Vector. Cells were analyzed 24 hours post Nucleofection® using light (A) and fluorescence microscopy (B).



Average transfection efficiency of Vero cells. Vero cells were transfected with program V-001 and 2  $\mu$ g of pmaxGFP® Vector. Cells were analyzed 24 hours post Nucleofection® by flow cytometry. Cell viability (% PI negative cells) is around 97% 24 hours post Nucleofection®.

## **Product Description**

Cat. No.		VCA-1003
Size (reactions)		25
Cell Line Nucleofector <sup>®</sup> Solution V		2.25 ml (2.05 ml + 10% overfill)
Supplement		0.5 ml (0.45 ml + 10% overfill)
pmaxGFP® Vector (0.5 µg/µl in 10 mM Tris pH 8.0)		30 µg
Certified cuvettes		25
Plastic pipettes		25
Storage and stability	Store Nucleofector <sup>®</sup> Sol	ution, Supplement and pmaxGFP $^{\circ}$ Vector at 4°C. For long-term storage,
	pmaxGFP® Vector is idea	lly stored at -20°C. The expiration date is printed on the solution box. Once the
	Nucleofector <sup>®</sup> Suppleme	nt is added to the Nucleofector® Solution it is stable for three months at 4°C.

#### **Required Material**

#### Note

- Please make sure that the entire supplement is added to the Nucleofector<sup>®</sup> Solution. The ratio of Nucleofector<sup>®</sup> Solution to supplement is 4.5:1. For a single reaction use 82 µl of Nucleofector<sup>®</sup> Solution plus 18 µl of supplement to make 100 µl of total reaction volume.
- Nucleofector<sup>®</sup> Device; Software requirements: version V2.1 or higher for Nucleofector<sup>®</sup> | Device; version S3-1 or higher for Nucleofector<sup>®</sup> || Device
- Supplemented Nucleofector® Solution at room temperature
- Supplied certified cuvettes
- Supplied plastic pipettes
- Supplied pmaxGFP<sup>®</sup> Vector
- Substrate of interest, highly purified, preferably by using endotoxin-free kits; A260 : A280 ratio should be at least 1.8
- 6-well culture dish or culture system of your choice
- For detaching cells: 2.5 mg/mlTrypsin and 1.0 mg/mlEDTA in PBS (5 x concentrated) and supplemented culture media or PBS/0.5% BSA
- Culture medium: Minimum Essential Medium (Eagle) with 2 mM L-glutamine and Earle's BSS adjusted to contain 1.5 g/L sodium bicarbonate, 0.1 mM non-essential amino acids and 1.0 mM sodium pyruvate, 90%; fetal bovine serum, 10%
- Prewarm appropriate volume of culture medium to 37°C (1.5 ml per sample)
- Appropriate number of cells (1 x 10<sup>6</sup> cells per sample; lower or higher cell numbers may influence transfection results)

## 1. Pre Nucleofection®

#### Cell culture recommendations

- 1.1 Replace media 2-3 times a week
- 1.2 Passage cells 3 times a week. A subcultivation ratio of 1:4 to 1:8 is recommended. Use low spin centrifugation (90xg)
- 1.3 Seed out  $1 2 \times 10^6$  cells/T162 flask
- 1.4 Subculture 2 days before Nucleofection®

#### **Trypsinization**

- 1.5 Remove media from the cultured cells and wash cells once with PBS; use at least same volume of PBS as culture media
- 1.6 For harvesting, incubate the cells ~5 minutes at 37°C with indicated trypsinization reagent (please see required material)
- 1.7 Neutralize trypsinization reaction with supplemented culture medium or PBS/0.5% BSA once the majority of the cells (>90%) have been detached

## 2. Nucleofection®

#### One Nucleofection® Sample contains

1 x 10<sup>6</sup> cells 2 μg plasmid DNA (in 1 – 5 μl H<sub>2</sub>0 or TE) or 2 μg pmaxGFP® Vector or 30 – 300nM siRNA (3 – 30 pmol/sample) 100 μl Cell Line Nucleofector® Solution V

- 2.1 Please make sure that the entire supplement is added to the Nucleofector® Solution
- 2.2 Prepare 6-well plates by filling appropriate number of wells with 1 ml of supplemented culture media and pre-incubate/equilibrate plates in a humidified 37°C/5% CO<sub>2</sub> incubator
- 2.3 Harvest the cells by trypsinization (please see 1.5 1.7)
- 2.4 Count an aliquot of the cells and determine cell density
- 2.5 Centrifuge the required number of cells (1 x 10<sup>6</sup> cells per sample) at 90xg for 10 minutes at room temperature. Remove supernatant completely
- 2.6 Resuspend the cell pellet carefully in 100 µl room-temperature Nucleofector® Solution per sample

## Note Avoid leaving the cells in Nucleofector<sup>®</sup> Solution for extended periods of time (longer than 15 minutes), as this may reduce cell viability and gene transfer efficiency.

- 2.7 Combine 100 μl of cell suspension with 2 μg DNA, 2 μg pmaxGFP<sup>®</sup> Vector or 30 nM 300 nM siRNA (3 30 pmol/sample) or other substrates
- 2.8 Transfer cell/DNA suspension into certified cuvette (sample must cover the bottom of the cuvette without air bubbles). Close the cuvette with the cap
- 2.9 Select the appropriate Nucleofector® Program V-001 (V-01 for Nucleofector® | Device)
- 2.10 Insert the cuvette with cell/DNA suspension into the Nucleofector<sup>®</sup> Cuvette Holder and apply the selected program by pressing the X-button
- 2.11 Take the cuvette out of the holder once the program is finished
- 2.12 Immediately add ~500 µl of the pre-equilibrated culture medium to the cuvette and gently transfer the sample into the prepared 6-well plate (final volume 1.5 ml media per well). Use the supplied pipettes and avoid repeated aspiration of the sample

### 3. Post Nucleofection®

3.1 Incubate the cells in humidified 37°C/5%  $CO_2$  incubator until analysis. Gene expression or down regulation, respectively, is often detectable after only 4 – 8 hours

## Additional Information

For an up-to-date list of all Nucleofector® References, please refer to: www.lonza.com/nucleofection-citations

#### For more technical assistance, contact our Scientific Support Team:

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