

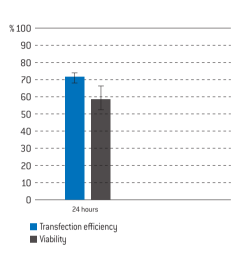
# Amaxa™ 4D-Nucleofector™ Protocol for MCF7

# For 4D-Nucleofector™ X Unit-Transfection in suspension

Human mammary gland adenocarcinoma; adherent epithelial cell line

# Example for Nucleofection™ of MCF7

Transfection efficiency and viability of MCF7 cells 24 hours post Nucleofection™. MCF7 cells were transfected with program EN-130 and 0.4 µg of pmaxGFP™ Vector in 20 µl Nucleovette™ Strips. 24 hours post Nucleofection™ cells were analyzed on a FACSCalibur™ [Becton Dickinson]. Cell viability was determined as % PI negative cells.



# **Product Description**

#### Recommended Kit(s)-SE Cell Line 4D-Nucleofector™ X Kit

Cat. No.	V4XC-1012	V4XC-1024	V4XC-1032
Transfection volume	100 μΙ	100 μΙ	20 µl
Size [reaction]	2 x 6	24	2 x 16
Nucleofector™ Solution	2 x 0.675 ml (0.492 ml + 27% overfill)	2.25 ml [1.968 ml + 13% overfill]	0.675 ml (0.525 ml + 22% overfill)
Supplement	2 x 0.15 ml (0.108 ml + 27% overfill)	0.5 ml (0.432 ml + 13% overfill)	0.15 ml (0.115 ml + 22% overfill)
pmaxGFP™ Vector (1 µg/µl in 10 mM Tris pH 8.0)	50 μg	50 μg	50 µg
Single Nucleocuvette™ (100 µI)	12	24	<u>-</u>
16-well Nucleocuvette™ Strips (20 µI)	-	-	2

## Storage and stability

Store Nucleofector<sup>™</sup> Solution, Supplement and pmaxGFP<sup>™</sup> Vector at  $4^{\circ}$ C. For long-term storage, pmaxGFP<sup>™</sup> Vector is ideally stored at -20°C. The expiration date is printed on the solution box. Once the Nucleofector<sup>™</sup> Supplement is added to the Nucleofector<sup>™</sup> Solution, it is stable for three months at  $4^{\circ}$ C.

### Note

4D-Nucleofector™ Solutions can only be used with Nucleovettes™ (conductive polymer cuvettes), i.e. in the 4D-Nucleofector™ System and the 96-well Shuttle™ Device. They are not compatible with the Nucleofector™ II/2b Device.

# Required Material

#### Note

Please make sure that the entire supplement is added to the Nucleofector $^{\text{\tiny{M}}}$  Solution. The ratio of Nucleofector $^{\text{\tiny{M}}}$  Solution to supplement is 4.5:1 (see table 1)

- 4D-Nucleofector™ System (4D-Nucleofector™ Core Unit and 4D-Nucleofector™ X Unit)
- Supplemented 4D-Nucleofector™ Solution at room temperature
- Supplied 100 µl single Nucleocuvette™ or 20 µl 16-well Nucleocuvette™
  Strips
- Compatible tips for 20 µl Nucleocuvette™ Strips: epT.I.P.S. [US/CDN: Eppendorf North America, Cat. No. 2491.431, Rest of World: Eppendorf AG, Cat. No. 0030073.266], Matrix TallTips® [Matrix Technologies Corp., Cat. No. 7281] or LTS Tips [Rainin Instruments, LLC, Cat. No. SR-L10F, SR/SS-L250S, SR/SS-L300S]. Before using other types of pipette tips, please ensure they reach the bottom of the Nucleocuvette™ Wells without getting stuck
- Supplied pmaxGFP™ Vector, stock solution 1µg/µl

#### Note

For positive control using pmaxGFP $^{\text{\tiny M}}$ , dilute the stock solution to an appropriate working concentration. Further details are provided in table 3 of this Optimized Protocol. The volume of substrate solution added to each sample should not exceed 10% of the total reaction volume (2  $\mu$ l for 20  $\mu$ l reactions; 10  $\mu$ l for 100  $\mu$ l reactions).

# Pre Nucleofection™

#### Cell culture recommendations

- 1.1 Replace media 2 times a week (30 ml per 162 cm² flask)
- 1.2 Cells should be passaged at 75-80% confluency
- 1.3 Seed out 2 x 10<sup>5</sup> cells/cm<sup>2</sup>
- 1.4 Subculture 3–4 days before Nucleofection™
- 1.5 For Nucleofection™ cells should be 75–80% confluent

#### **Trypsinization**

- 1.6 Remove media from the cultured cells and wash cells once with PBS; use at least same volume of PBS as culture media
- 1.7 For harvesting, incubate the cells at 37°C with e.g. 0.05% trypsin/0.02% EDTA
- 1.8 Inactivate trypsinization reaction with supplemented culture media or PBS/0.5% BSA

# 2. Nucleofection™

For Nucleofection™ Sample contents and recommended Nucleofector™ Program, please refer to Table 3.

- 2.1 Please make sure that the entire supplement is added to the Nucleofector™ Solution
- 2.2 Start 4D-Nucleofector™ System and create or upload experimental parameter file (for details see device manual)
- 2.3 Select/Check for the appropriate Nucleofector™ Program (see table 3)
- 2.4 Prepare cell culture plates by filling appropriate number of wells with desired volume of recommended culture media (see table 4) and pre-incubate/equilibrate plates in a humidified 37°C/5% CO<sub>2</sub> incubator
- 2.5 Pre-warm an aliquot of culture medium to 37°C (see table 4)
- 2.6 Prepare plasmid DNA or pmaxGFP™ Vector or siRNA (see table 3)
- 2.7 Harvest the cells by trypsinization (please see 1.6–1.8)
- 2.8 Count an aliquot of the cells and determine cell density
- 2.9 Centrifuge the required number of cells (see table 3) at 90xg for 10 minutes at room temperature. Remove supernatant completely
- 2.10 Resuspend the cell pellet carefully in room temperature 4D-Nucleofector™ Solution (see table 3)
- 2.11 Prepare mastermixes by dividing cell suspension according to number of substrates
- 2.12 Add required amount of substrates to each aliquot (max. 10% of final sample volume)
- 2.13 Transfer mastermixes into the Nucleocuvette™ Vessels

#### Note

As leaving cells in Nucleofector™ Solution for extended periods of time may lead to reduced transfection efficiency and viability it is important to work as quickly as possible. Avoid air bubbles while pipetting.

- 2.14 Gently tap the Nucleocuvette™ Vessels to make sure the sample covers the bottom of the cuvette
- 2.15 Place Nucleocuvette™ Vessel with closed lid into the retainer of the 4D-Nucleofector™ X Unit. Check for proper orientation of the Nucleocuvette™ Vessel
- 2.16 Start Nucleofection™ Process by pressing the "Start" on the display of the 4D-Nucleofector™ Core Unit (for details, please refer to the device manual)
- 2.17 After run completion, carefully remove the Nucleocuvette™ Vessel from the retainer
- 2.18 Resuspend cells with pre-warmed medium (for recommended volumes see table 5). Mix cells by gently pipetting up and down two to three times. When working with the 100 µl Nucleocuvette™ use the supplied pipettes and avoid repeated aspiration of the sample
- 2.19 Plate desired amount of cells in culture system of your choice (for recommended volumes see table 5)

# 3. Post Nucleofection™

3.1 Incubate the cells in humidified  $37^{\circ}\text{C}/5\%$  CO<sub>2</sub> 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:

#### USA /Canada

Phone: 800 521 0390 (toll-free)

Fax: 301 845 8338

E-mail: scientific.support@lonza.com

#### Europe and Rest of World

Phone: +49 221 99199 400 Fax: +49 221 99199 499

E-mail: scientific.support.eu@lonza.com

#### Lonza Cologne GmbH 50829 Cologne, Germany

Please note that the Amaxa™ Nucleofector™ Technology is not intended to be used for diagnostic purposes or for testing or treatment in humans.

The Nucleofector" Technology, comprising Nucleofection" Process, Nucleofector" Device, Nucleofector" Solutions, Nucleofector" 96-well Shuttle" System and 96-well Nucleocuvette" plates and modules is covered by patent and/or patent-pending rights owned by Lonza Cologne AG.

A maxa, Nucleo fector, Nucleo fection, 4D-Nucleo fector, Nucleo cuvette and max GFP are registered trademarks of the Lonza Cologne AG in Germany and/or U.S. and/or other countries.

Other product and company names mentioned herein are the trademarks of their respective owners.

This kit contains a proprietary nucleic acid coding for a proprietary copepod fluorescent protein intended to be used as a positive control with this Lonza product only. Any use of the proprietary nucleic acid or protein other than as a positive control with this Lonza product is strictly prohibited. USE IN ANY OTHER APPLICATION REQUIRES A LICENSE FROM EVROGEN. To obtain such a license, please contact Evrogen at license@evrogen.com.

The CMV promoter is covered under U.S. Patents 5,168,062 and 5,385,839 and its use is permitted for research purposes only. Any other use of the CMV promoter requires a license from the University of lowa Research Foundation, 214 Technology Innovation Center, lowa City, IA 52242.

The use of this product in conjunction with materials or methods of third parties may require a license by a third party. User shall be fully responsible for determining whether and from which third party it requires such license and for the obtainment of such license.

No statement is intended or should be construed as a recommendation to infringe any existing patent.

© Copyright 2010, Lonza Cologne GmbH. All rights reserved.

# Table 1: Volumes required for a single reaction

	100 µl Single Nucleocuvette™	20 µl Nucleocuvette™ Strip
Volume of Nucleofector™ Solution	82 µl	16.4 μΙ
Volume of Supplement	18 µl	3.6 µl

## Table 2: Required amounts of cells and media for Nucleofection™

	100 μl Single Nucleocuvette™	20 µl Nucleocuvette™ Strip
Culture medium per sample post Nucleofection™ (for transfer and culture)	1.5 ml	255 μl
Cell number per Nucleofection™ Sample	$2\times10^6$ (Minimal cell number: $8\times10^5$ cells, a lower cell number may lead to a major increase in cell mortality; maximum cell number: $4\times10^6$ )	$4 \times 10^5$ (Lower or higher cell numbers may influence transfection results)

# Table 3: Contents of one Nucleofection™ Sample and recommended program

2 x 10 <sup>6</sup>	4 x 10 <sup>5</sup>
2 µg	0.4 μg
1-2 µg	0.2-1 µg
30-300nM siRNA (3-30 pmol/sample)	30-300nM siRNA $(0.6-6$ pmol/sample)
100 µl	20 µl
EN-130	EN-130
	30–300nM siRNA (3–30 pmol/sample) 100 µl

# Table 4: Culture volumes required for post Nucleofection™ Steps

	100 µl Single Nucleocuvette™	20 µl Nucleocuvette™ Strip*
6-well culture plate	1 ml	<u> </u>
96-well culture plate	<u> </u>	_175 µl
Culture medium to be added to the sample post Nucleofection™	500 μΙ	80 μΙ
* Maximum cuvette volume 200 µl		

# Table 5: Recommended volumes for sample transfer into culture plate

	100 µl Single Nucleocuvette™	20 μl Nucleocuvette™ Strip*
Culture medium to be added to the sample post Nucleofection™	500 μl	80 µl
Volume of sample transferred to culture plate	complete sample (use supplied pipettes)	25 µl
* Maximum cuvette volume 200 µl		