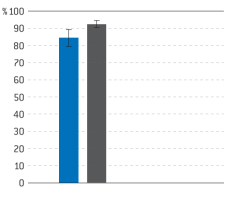
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Amaxa[™] 4D-Nucleofector[™] Protocol for HeLa S3 [ATCC[®]] For 4D-Nucleofector[™] X Unit—Transfection in suspension

Human cervix adenocarcinoma; adherent epithelial cell line; [ATCC[®] CCL-2.2[™], cryopreserved]

Example for Nucleofection™ of HeLa S3 cells

Transfection efficiency of HeLa S3 cells 24 hours post Nucleofection[™]. HeLa S3 cells [ATCC[®] CCL-2.2[™]] were transfected using Nucleofection[™] Program DS-150 and 0.4 µg pmaxGFP[™] Vector in 20 µl Nucleovette[™] Strips. 24 hours post Nucleofection[™] cells were analyzed on a FACSCalibur[™] [Becton Dickinson]. Cell viability was determined with CellTiter-Glo[®] Viability Assay [Promega, Cat. No. G7570].



Transfection efficiency
Viability

Product Description

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

Cat. No.	V4XC-1012	V4XC-1024	V4XC-1032
Transfection volume	100 µl	100 µl	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 µl)	12	24	
16-well Nucleocuvette™ Strips (20 µI)	-		2

Storage and stability

Note

Store Nucleofector[™] Solution, Supplement and pmaxGFP[™] Vector at 4 °C. For long-term storage, pmaxGFP[™] Vector is ideally stored at -20 °C. The expiration date is printed on the solution box. Once the Nucleofector[™] Supplement is added to the Nucleofector[™] Solution, it is stable for three months at 4 °C. 4D-Nucleofector[™] Solutions can only be used with Nucleocuvettes[™] (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[>] Solution. The ratio of Nucleofector[>] 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^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 µl for 20 µl reactions; 10 µl for 100 µl reactions).

- Substrate of interest, highly purified, preferably by using endotoxinfree kits; A260:A280 ratio should be at least 1.8
- Cell culture plates of your choice
- Culture medium: Ham's F12K media with 2 mM L-glutamine adjusted to contain 1.5 g/L sodium bicarbonate, 90 % [ATCC[®] 30-2004]; fetal bovine serum, 10 % [ATCC[®] 30-2020]
- Prewarm appropriate volume of culture medium to 37 °C (see table 2)
- Appropriate number of cells/sample (see table 2)

1. Pre Nucleofection™

Cell culture recommendations

- 1.1 Replace media and passage the cells 3 times a week
- 1.2 Seed out $2.4-3 \times 10^6$ cells per T162 flask
- 1.3 Cells should be passaged 2–3 days before Nucleofection™
- 1.4 A subcultivation ratio of 1:4–1:6 is recommended

Trypsinization

- 1.5 Remove media from the cultured cells and wash cells once with PBS; use at least same volume PBS as culture media
- 1.6 For harvesting, incubate the cells at 37 °C with e.g. 0.5mg/ml trypsin; 0.2mg/ml EDTA in PBS
- 1.7 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 table3)
- 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 % CO₂ 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 1.5–1.7)
- 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 °C/5 % CO₂ 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

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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 GmbH.

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Table 1: Volumes required for a single reaction

	100 µl Single Nucleocuvette™	20 µl Nucleocuvette™ Strip
Volume of Nucleofector™ Solution	82 μl	16.4 μl
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)	2 ml	360 µl
Cell number per Nucleofection™ Sample	1×10^6 (Lower or higher cell numbers may influence transfection results)	2 x 10 ⁵ (Lower or higher cell numbers may influence transfection results)

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

		_100 μl Single Nucleocuvette™	20 µl Nucleocuvette™ Strip
Cells		1 x 10 ⁶	2 x 10 ⁵
Substrate*	pmaxGFP™ Vector	2 µg	0.4 µg
or	plasmid DNA (in H_20 or TE)	1-2 µg	0.2-1 µg
or	siRNA	30–300nM siRNA (3–30 pmol/sample)	30–300nM siRNA (0.6–6 pmol/sample)
SE Cell Line 4D-Nucleo	ofector™ X Solution	100 µl	20 µl
Program		DS-150 for high transfection efficiency or CA-123 for high viability	DS-150 for high transfection efficiency or CA-123 for high viability
* Volume of substrate should	comprise maximum 10 % of total reaction v	olume	

Table 4: Culture volumes required for post Nucleofection™ Steps

	100 µl Single Nucleocuvette™	20 µl Nucleocuvette™ Strip*
12-well culture plate	1.5 ml	· .
96-well culture plate	·	180 µl
Culture medium to be added to the sample post Nucleofection™	500 µl	180 µl
* 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	180 µl
Volume of sample transferred to culture plate	complete sample (use supplied pipettes)	20 µl
* Maximum cuvette volume 200 µl		