Electroporation protocol for SaOS2 cells

GTporator®

Transfection protocol Protocol No. 09/2008-010

Cell line SaOS2

Washing solutions Phosphate buffered saline (PBS), pH 7.4, Gtporator®-M

Cell count1-3 x 106Electroporation solutionGtporator®-MCuvette2 mm gap width

Volume 80 μl

Temperature Room temperature

DNA 5 μg in water

Instrument settings

Single amplitude

Protocol Square-wave

 $\begin{array}{ll} \mbox{Voltage} & 200 \mbox{ V} \\ \mbox{(Capacitance} & 960 \mbox{ } \mu\mbox{F)} \\ \mbox{Pulse time} & 20 \mbox{ ms} \end{array}$

Pulse number 1

Radio-frequency

Voltage 200 V
Frequency 40 kHz
Pulses 5 x 4 ms
Interval 1 s

Modulation 100%

- 1. Trypsinize subconfluent cells
- 2. Wash the cells once in 1 ml PBS, and once in 300 µl of Gtporator® solution
- 3. Resuspend the cells in 80 µl Gtporator® solution with 5 µg of plasmid DNA
- 4. Transfer into a 2 mm electroporation cuvette
- 5. Electroporate
- 6. Immediately add pre-warmed culture medium with serum, transfer the cells into cell culture flask and place in incubator
- 7. Test the gene expression 24 h after electroporation

Notes:

- The overall incubation time in the Gtporator® solution must not exceed 20 minutes to guarantee a successful electroporation!
- The best results are obtained with DNA purified by anion-exchange chromatography.
- DNA should be dissolved in sterile water, using the TE buffer slightly lowers the survival rate.

Typical results:

Survival rate in control cells 70%

Transfection rate: 60-75% with variations depending on the quality of DNA.

Results were measured 24 hours after transfection.

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