

jetPRIME[®] transfection reagent

Short protocol – DNA transfection



Day 0: Cell seeding

→ Seed cells in **V** mL of cell growth medium according to the table below

Quantities per well, dish or flask

Culture vessel	Number of cells*	V = volume of medium during transfection
96-well	7500 - 10 000	0.1 mL
24-well	50 000 - 80 000	0.5 mL
12-well	80 000 - 150 000	1 mL
6-well / 35 mm	150 000 - 250 000	2 mL
100 mm / flask 75 cm ²	1 x 10 ⁶ - 2 x 10 ⁶	10 mL

*For specific cell type or suspension cells, please refer to the complete protocol.

Day 1: Transfection

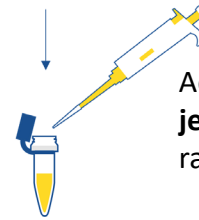
→ Perform transfection **in the presence of serum**

→ Use **jetPRIME[®] buffer only**

→ Transfect cells at **60-80% confluency**



Dilute **X** µg of DNA in **W** µL of **jetPRIME[®] buffer**. Vortex 10 s and spin down

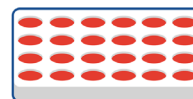


Add **Y** µL of **jetPRIME[®] reagent** (starting ratio 1:1)



Vortex 1 s, spin down and incubate 10 min at RT

Add transfection mix to the cells in serum containing medium



Incubate 24 to 48 h

Quantities per well, dish or flask

Culture vessel	W = volume of jetPRIME [®] buffer	X = amount of DNA added	Y = volume of jetPRIME [®] reagent
96-well	10 µL	0.1 µg	0.2 µL
24-well	50 µL	0.5 µg	1 µL
12-well	75 µL	0.8 µg	1.6 µL
6-well / 35 mm	200 µL	2 µg	4 µL
100 mm / flask 75 cm ²	500 µL	10 µg	20 µL

Day 2-3: Measure gene expression

See back page for optimization tips

Download complete protocol on <https://myaccount.polyplus-transfection.com/>

+33 (0)3 90 40 61 80

support@polyplus-transfection.com

www.polyplus-transfection.com

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jetPRIME[®] transfection reagent

Short protocol – Optimization tips (DNA)



+ Protocol Optimization

- + Test different DNA amounts: X, 0.5X and 1.5X.
- + Test different DNA/jetPRIME[®] ratios, 1:2 to 1:3.
- + For cell specific protocols, check our online Cell Transfection Database:
<http://www.polyplus-transfection.com/resources/cell-transfection-database/>

Quantities per well, dish or flask

Culture vessel	W = volume of jetPRIME [®] buffer	X = amount of DNA added	Y = volume of jetPRIME [®] reagent
96-well	10 µL	0.05 – 0.20 µg	0.10 – 0.60 µL
24-well	50 µL	0.25 – 0.75 µg	0.50 – 2.25 µL
12-well	75 µL	0.4 – 1.2 µg	0.8 – 3.6 µL
6-well / 35 mm	200 µL	1 – 3 µg	2 – 9 µL
100 mm / flask 75 cm ²	500 µL	5 – 15 µg	10 – 45 µL

For HEK-293 and HeLa cells, you may decrease the DNA amount to 0.5X and use the 1:2 DNA/jetPRIME[®] ratio.

+ Tips to increase cell viability of sensitive cells

- + Replace medium 4 h after transfection.
- + Decrease DNA amount to 0.5 X while maintaining the DNA/jetPRIME[®] ratio previously used.
- + Analyze transfection at an earlier time point (24 h after transfection instead of 48 h for instance).
- + Check that the target gene does not affect cell viability.

+ Good mRNA Transfection Practices

- + Store appropriately jetPRIME[®] (5 ± 3°C).
- + Ensure that cells have been passaged more than twice and less than 20 times prior to transfection.
- + Discard overconfluent cells.
- + Regularly check for mycoplasma contaminations.
- + Use a reporter gene to set up and optimize transfection conditions.
- + Serum quality may drastically affect transfection efficiency. When purchasing a new batch of serum or trypsin, check cell viability as well as transfection efficiency.

Note: jetPRIME[®] is also recommended for DNA/siRNA co-transfection. Please refer to the complete protocol available when creating your account online at: <https://myaccount.polyplus-transfection.com/>.

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