

jetOPTIMUS[®] *in vitro* DNA transfection reagent PROTOCOL

DESCRIPTION

jetOPTIMUS[®] is an innovative cationic nanotechnology developed to improve cellular uptake and endosomal escape of DNA in adherent cells resulting in higher transfection efficiency, even in hard-totransfect cells. In order to work in relevant physiological conditions, transfection with jetOPTIMUS[®] requires a minimum DNA quantity and reagent volume which preserves cell viability and morphology.

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1 DNA TRANSFECTION PROTOCOL

1.1 CELL SEEDING

For optimal transfection of adherent cells using jetOPTIMUS[®] reagent, cells should be seeded the day before transfection to reach <u>60 to 80 % confluency</u> at the time of transfection. Typically, for experiments in 24-well plates, 50 000 adherent cells are seeded per well in 500 μ L complete growth medium 24 h prior to transfection. For other formats, please refer to Table 1:

Culture vessel	Number of cells to prepare <u>per well</u>	Surface area per well (cm²)	Volume of growth medium to seed the cells (mL)
96-well	7500 - 25 000	0.3	0.125
24-well	40 000 - 100 000	1.9	0.5
6-well / 35 mm	150 000 - 400 000	9.4	2
60 mm / flask 25 cm ²	200 000 - 850 000	25 - 28	5
100 mm / flask 75 cm ²	1 x 10 ⁶ - 4 x 10 ⁶	75 - 78.5	10

1.2 DNA TRANSFECTION PROTOCOL

The following protocol is given for transfection of DNA <u>at 0.5 μ g</u> per well in a <u>24-well</u> plate. For other culture formats, please refer to Table 2.

- 1. Dilute 0.5 μg DNA into 50 μL jetOPTIMUS[®] buffer. Vortex for 1 second and spin down briefly.
- 2. Vortex jetOPTIMUS[®] reagent for 5 seconds and spin down before use.
- Add 0.5 μL jetOPTIMUS[®] onto the DNA solution (ratio 1:1 corresponding to μg_{DNA}:μL_{reagent}), <u>vortex</u> <u>for 1 second</u> and spin down briefly.
- 4. Incubate for 10 minutes at room temperature.
- 5. Add 50 μ L of transfection mix per well dropwise onto the cells in serum containing medium and distribute evenly.
- 6. Gently rock the plates back and forth and from side to side, incubate the plate at 37 °C.
- 7. If needed, replace transfection medium after 4 hours by cell growth medium and return the plates to the incubator.
- 8. Analyze transgene expression after 24 hours or later.





Culture vessel	Volume of jetOPTIMUS® buffer (μL)	Amount of DNA (μg)	Volume of jetOPTIMUS® reagent (µL)	Volume of growth medium (mL)
96-well	12.5	0.13	0.13 - 0.19	0.125
24-well	50	0.5	0.5 - 0.75	0.5
6-well / 35 mm	200	2	2 - 3	2
60 mm / flask 25 cm ²	500	4	4 - 6	5
100 mm / flask 75 cm ²	1000	10	10 - 15	10

Table 2. DNA transfection guidelines according to the cell culture vessel used

NOTE: jetOPTIMUS[®] buffer must be used for successful transfection.



jetOPTIMUS® transfection protocol for 24-well plates

1.3 OPTIMIZATION GUIDELINES AND CONDITIONS FOR SPECIFIC CELL TYPES

You may adjust the volume of reagent and/or the amount of DNA. We recommend testing different ratios of DNA / jetOPTIMUS[®] reagent from 1:1 to 1:1.5 (μg_{DNA} : $\mu L_{reagent}$) (see Table 3).

Culture vessel	Volume of jetOPTIMUS® buffer (μL)	Amount of DNA (μg)	Volume of jetOPTIMUS® reagent (µL)	Volume of growth medium (mL)
96-well	12.5	0.10 - 0.20	0.10 - 0.30	0.125
24-well	50	0.25 - 0.75	0.25 - 1	0.5
6-well / 35 mm	200	1 - 3	1 - 4.5	2
60 mm / flask 25 cm ²	500	2 - 6	2 - 9	5
100 mm / flask 75 cm ²	1000	5 - 15	5 - 22	10

Table 3. Optimization guidelines according to the cell culture vessel used

Specific conditions for many cell types can be found in our cell transfection database, following this link: <u>www.polyplus-transfection.com/resources/cell-transfection-database/</u>.

1.4 OTHER APPLICATIONS

For cotransfection of multiple plasmids, the total DNA amount per well/plate should not exceed the DNA amount indicated in Table 2. The ratio to use for each plasmid depends on the size of the plasmids, the plasmid constructs and the desired expression level of each plasmid. Please adjust the ratios according to your application, each plasmid representing at least 10 % of the total DNA amount per well/plate.

For other applications, such as stable transfection and plasmid-based CRISPR/Cas9 genome editing, please contact our Scientific Support team for specific recommendations at support@polyplus-transfection.com.





2 TROUBLESHOOTING

Observations	Actions			
	Ensure that the nucleic acid is diluted in the provided jetOPTIMUS [®] buffer.			
	Ensure that cells have been passaged more than twice and less than 20 times prior to transfection. Discard overconfluent cells.			
	Ensure that the medium is permissive to the transfection.			
Low DNA transfection	Use high-quality plasmid preparation, free of proteins.			
efficiency	Optimize the volume of jetOPTIMUS [®] reagent and the amount of plasmid DNA added per well. Increase the volume of jetOPTIMUS [®] reagent first; if insufficient, increase the amount of DNA (both according to Table 2).			
	Serum quality may drastically affect transfection efficiency. When purchasing a net batch of serum or trypsin, check cell viability as well as transfection efficiency.			
	Use a plasmid containing a common reporter gene such as Luciferase or GFP as a positive control.			
	Ensure that the nucleic acid is diluted in the provided jetOPTIMUS [®] buffer.			
	Ensure that the plasmid preparation is endotoxin-free.			
	Replace complete medium (containing serum or supplements) with serum-free medium (OptiMEM [®]) at the time of transfection and during 4 hours.			
Cellular toxicity	Analyze transfection at an earlier time point (e.g. at 24 h instead of 48 h).			
	Decrease the volume of jetOPTIMUS [®] reagent.			
	Decrease the amount of plasmid DNA added per well.			
	Verify the toxicity of the expressed protein. If the expressed protein is toxic for the cells, reduce the amount of plasmid DNA.			

3 PRODUCT INFORMATION

3.1 ORDERING INFORMATION

Ref. N°	jetOPTIMUS® Reagent	jetOPTIMUS [®] Buffer	
117-01	0.1 mL	10 mL	
117-07	0.75 mL	2 x 60 mL	
117-15	1.5 mL	4 x 60 mL	

3.2 PROVIDED BUFFER

jetOPTIMUS[®] reagent is provided with an optimized sterile buffer (jetOPTIMUS[®] buffer). This buffer <u>must</u> be used to ensure successful transfection experiments.

3.3 CONTENT

1.5 mL of jetOPTIMUS[®] transfection reagent is sufficient to perform 3 000 transfections in 24-well plates or 750 transfections in 6-well plates following the standard protocol (DNA:reagent ratio = 1:1).

3.4 REAGENT USE AND LIMITATIONS

For research use only. Not for use in humans.

3.5 QUALITY CONTROL

Every batch of jetOPTIMUS[®] reagent is tested by DNA transfection of HeLa cells with a GFP-expressing plasmid.

3.6 FORMULATION AND STORAGE

jetOPTIMUS[®] and its buffer are shipped at room temperature but should be stored at 5 ± 3 °C upon arrival to ensure long term stability. jetOPTIMUS[®], as guaranteed and indicated on the Certificate of Analysis, is stable at least for 6 months (Ref. N° 117-01) to at least one year (Ref. N° 117-07, 117-15) when stored appropriately.

jetOPTIMUS[®] is chemically-defined and guaranteed free of animal origin products.

Polyplus-transfection[®] has been awarded ISO 9001 Quality Management System Certification since 2002, which ensures that the company has established reliable and effective processes for manufacturing, quality control, distribution and customer support.





3.7 TRADEMARKS

Polyplus-transfection and jetOPTIMUS are registered trademarks of Polyplus-transfection S.A.

3.8 TECHNICAL ASSISTANCE AND SCIENTIFIC ADVICE

Contact the friendly Polyplus technical support via:

- The Polyplus website: www.polyplus-transfection.com
- <u>Email</u>: support@polyplus-transfection.com
- Phone: +33 3 90 40 61 87