

# jetMESSENGER™

## *in vitro* mRNA transfection reagent

# PROTOCOL

### Description

jetMESSENGER™ is a novel powerful transfection reagent manufactured at Polyplus-transfection®. jetMESSENGER™ has been specifically designed for high mRNA transfection efficiency in usually difficult to transfect cells such as primary cells, cancer cell lines, neurons and stem cells. jetMESSENGER™ can also be used on a wide variety of easy to transfect cells. Transfection with jetMESSENGER™ leads to very low cytotoxicity as it requires low amounts of mRNA and low volumes of reagent.

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# 1 Transient mRNA transfection protocol

## 1.1 Cell seeding

For optimal mRNA transfection conditions, we recommend using cells which are 60 to 80% confluent at the time of transfection. Typically, for experiments in 24-well plates, 50 000 adherent cells or 100 000 suspension cells are seeded per well in 0.5 mL of cell growth medium 24 h prior to transfection. For other culture formats, refer to Table 1. For more details about seeding various cell lines, refer to Table 2.

jetMESSENGER™ is compatible with the presence of serum and antibiotics therefore you may use serum and antibiotic containing medium during the entire experiment.

**Table 1. Recommended seeding density the day before transfection.**

Culture vessel	Adherent cell number	Suspension cell number	Surface area per well (cm <sup>2</sup> )	Volume of medium per well to seed the cells (mL)
96-well	2 000 – 12 500	25 000	0.3	0.125
24-well	7 000 – 50 000	100 000	1.9	0.5
12-well	14 000 – 100 000	200 000	3.8	1
6-well/35 mm	80 000 – 200 000	400 000	9.4	2
60 mm / flask 25 cm <sup>2</sup>	200 000 – 800 000	700 000	20 - 25	5
100 mm / flask 75 cm <sup>2</sup>	1 x 10 <sup>6</sup> – 2 x 10 <sup>6</sup>	1.2 x 10 <sup>6</sup>	60 - 75	10
150 mm / flask 175 cm <sup>2</sup>	2 x 10 <sup>6</sup> – 5 x 10 <sup>6</sup>	1.7 x 10 <sup>6</sup>	150 - 175	20

**Table 2. Recommended seeding density for various cell lines**

Cell type	Cells	Number of cells to seed per well of a 24-well plate	Number of plating days before transfection (day)
Epithelial	Caco-2	40 000	1
	HeLa	50 000	1
	MCF10-A	80 000	1
	MCF-7	50 000	1
	MDCK	40 000	1
	U-87 MG	50 000	1
Fibroblast	BJ	20 000	1
	MEF	12 000	3
	IMR-90	40 000	1
Hepatocyte	Hep G2	100 000	1
Human stem cells	hMSC	12 000	3
Lymphocyte	Jurkat	100 000	1
	K-562	100 000	1
Monocyte	THP-1	100 000	1
Mouse stem cells	mES	50 000	3
Primary cells	Monocytes	400 000	1
	Dendritic cells *	400 000	10
	Macrophages *	400 000	10
	Fibroblasts	7 000	3

\*Obtained from differentiation and maturation of monocytes

## 1.2 mRNA Transfection Protocol

The following conditions are given per well of a 24-well plate. For other culture formats, please refer to Table 3.

1. Dilute 0.5 µg mRNA into 50 µL jetMESSENGER™ mRNA buffer (supplied). Mix by vortexing.
2. Add 1 µL jetMESSENGER™, mix by vortexing, spin down briefly.
3. Incubate for 15 min at RT.
4. Add 50 µL of transfection mix per well dropwise onto the cells in growth medium (containing serum or not) and/or additives (standard culture medium), and distribute evenly.
5. Gently rock the plate back and forth and from side to side.
6. Analyze at least 24 - 48 h later.

**Table 3. mRNA transfection guidelines per well according to the cell culture vessel**

Culture vessel	Volume of mRNA buffer (µL)	Amount of mRNA (µg)	Volume of jetMESSENGER™ Reagent (µL)
96-well	12.5	0.1 ± 0.05	0.25 ± 0.05
24-well	50	0.5 ± 0.1	1 ± 0.2
12-well	100	1 ± 0.2	2 ± 0.4
6-well/35 mm	200	2 ± 0.5	4 ± 0.8
60 mm / flask 25 cm <sup>2</sup>	625	4 ± 1	8 ± 1.6
100 mm / flask 75 cm <sup>2</sup>	1 875	10 ± 2.5	20 ± 4

**NOTE:** the provided mRNA buffer should be used for successful transfection with jetMESSENGER™.

Prepare a master mix of minimum 50 µL to allow accurate pipetting and homogenous preparation of the complexes.

## 1.3 Guidelines

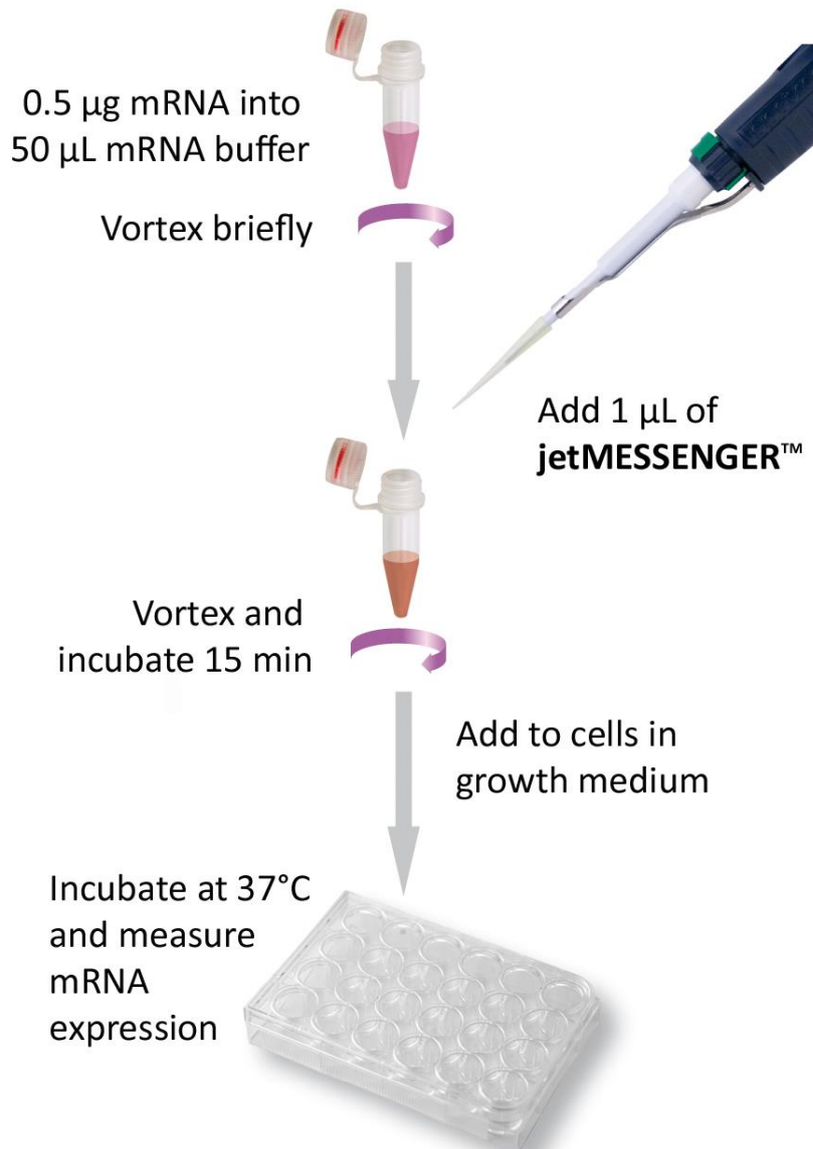
For optimal mRNA transfection conditions, we recommend using chemically modified mRNA. Transfection should be performed in a RNase-free working-environment and mRNA should be diluted and aliquoted in RNase-free water.

**Note:** Performing media change 4 h post-transfection may improve cell viability.

**Browse our cell transfection database to find the optimized conditions according to your cell line:**

<http://www.polyplus-transfection.com/resources/cell-transfection-database/>



**jetMESSENGER™ Transfection in 24-well Plate**

## 1.4 Optimization guidelines

Transfection conditions should be optimized for each cell line. You may refer to the optimised conditions for various cell lines detailed in **Table 3**, and on our online cell transfection database (<http://www.polyplus-transfection.com/resources/cell-transfection-database/>).

You may adjust the volume of reagent and/or the amount of mRNA. The volume of jetMESSENGER™ may range between **1.6 - 2.4 µL per µg of mRNA** depending on the transfected cell line. The amount of mRNA may range between 0.5 X and 2 X, X being the amount indicated in Table 3.

## 2 Transfection of CRISPR/CAS9

jetMESSENGER™ is suited for genome editing applications using Cas-9 encoding mRNA co-transfected with guide RNA into mammalian cells.

For co-transfection of multiple nucleic acids, the total mRNA amount added per well (or plate) should correspond to the mRNA amounts indicated in **Table 3**.

The following conditions are given per well of a 24-well plate. For other culture format, please refer to **Table 3**.

1. Dilute 0.5 µg mRNA into 50 µL mRNA Buffer (supplied). Mix by vortexing.
2. Add 1 µL jetMESSENGER™, mix by vortexing, spin down briefly.
3. Incubate for 15 min at RT.
4. Add 50 µL of transfection mix per well dropwise onto the cells in medium containing serum, and distribute evenly.
5. Gently rock the plates back and forth and from side to side.
6. Analyze after 48 - 72 h or later.

If you would like to perform CRISPR/Cas9 transfection experiments using another type of nucleic acids such as plasmid DNA, please contact our technical support at [support@polyplus-transfection.com](mailto:support@polyplus-transfection.com).

### 3 Troubleshooting

Observations	Actions
<p><b>Low mRNA transfection efficiency</b></p>	<p>Optimize the volume of jetMESSENGER™ reagent and the amount of mRNA added per well. Increase the volume of jetMESSENGER™ reagent first; if insufficient, increase the amount of mRNA according to <b>Table 3</b>.</p> <p>To adjust the volume of reagent and/or the amount of mRNA:</p> <ul style="list-style-type: none"> <li>- the volume of jetMESSENGER™ may range between <b>1.6 - 2.4 µL per µg of mRNA</b> depending on the transfected cell line.</li> <li>- the amount of mRNA may range between 0.5X and 2X, X being the amount indicated in <b>Table 3</b>.</li> </ul>
	<p>Replace medium containing serum with serum-free medium (OptiMEM®) during transfection.</p>
	<p>Ensure the medium is permissive to the transfection.</p>
	<p>Ensure that the mRNA is diluted in the provided mRNA buffer by Polyplus-transfection®.</p>
	<p>Use a common reporter gene-encoding mRNA as a positive control (ex: Luciferase or GFP).</p>
	<p>Ensure that the quality of your mRNA is optimal. Preferably use mRNA purchased from an oligo supplier, instead of homemade transcribed mRNA.</p>
	<p>The use of chemically modified mRNA (Pseudouridine, 5' Methylcytosine, etc...) could improve the transfection efficiency.</p>
	<p>Ensure that all reagents are RNase-free.</p>
<p><b>Cellular toxicity</b></p>	<p>Analyze transfection at an earlier time point (e.g. at 24 h instead of 48 h).</p>
	<p>Wash cells 4 h after transfection.</p>
	<p>Decrease the amount of mRNA added per well.</p>
	<p>Ensure that the mRNA is diluted in the provided mRNA buffer.</p>
	<p>Decrease the volume of jetMESSENGER™ reagent.</p>
	<p>Ensure that the mRNA used is chemically modified.</p>
	<p>Check if the expressed protein may cause toxicity. If the expressed protein is toxic for the cells, reduce the amount of mRNA.</p>

## 4 Product Information

### 4.1 Ordering information

Cat. N°	jetMESSENGER™ Reagent	mRNA Buffer
150-01	0.1 mL	10 mL
150-07	0.75 mL	60 mL
150-15	1.5 mL	2 x 60 mL

### 4.2 Additional Buffer

jetMESSENGER™ reagent is provided with an optimized sterile buffer (mRNA buffer). This buffer **must** be used to ensure successful transfection experiments.

### 4.3 Content

1.5 mL of jetMESSENGER™ transfection reagent is sufficient to perform up to 1500 transfections in a 24-well plate and 375 transfections in 6-well plate format.

### 4.4 Reagent use and Limitations

For research use only. Not for use in humans.

### 4.5 Quality control

Every batch of jetMESSENGER™ mRNA transfection reagent is tested in-house by mRNA transfection of CaCo-2 cells with a GFP-expressing mRNA. Each vial of reagent is provided with Certificate of Analysis.

### 4.6 Formulation and Storage

jetMESSENGER™ and its buffer are shipped at room temperature but should be stored at 4°C upon arrival to ensure long term stability. jetMESSENGER™, as guaranteed and indicated in the Certificate of Analysis, is stable for 6 months (150-01) to at least one year (other packaging sizes) when stored appropriately.

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.

## 4.7 Trademarks

Polyplus-transfection and jetMESSENGER are registered trademarks of Polyplus-transfection.

## 4.8 Technical Assistance and Scientific Advice

**Contact the friendly Polyplus technical support *via*:**

- The Polyplus website: [www.polyplus-transfection.com](http://www.polyplus-transfection.com)
- Email: [support@polyplus-transfection.com](mailto:support@polyplus-transfection.com)
- Phone: +33 3 90 40 61 87