

Membrane fusion is a highly efficient method for transfecting various molecules and particles into mammalian cells, even into sensitive and primary cells. The Fuse-It reagents are cargo-specific liposomal carriers that attach and rapidly fuse with plasma membranes in a physiochemical-driven manner, thus releasing their cargo into the cytoplasm of the target cell. Membrane fusion is an effective tool for functional studies, which works independently of biological processes such as endocytosis, pinocytosis, or specific receptor binding.

Overview

Fuse-It-mRNA is a proprietary formulation reagent created for the transfection of messenger RNA (mRNA) into the cytoplasm of mammalian cells. Requiring only short incubation times of up to twenty minutes, the transfer of mRNA is highly biocompatible and non-toxic, which results in high transfer efficiency without influencing the cell viability. Due to mRNA transfer directly into the cytoplasm, lysosomal degradation and the use of lysosome function-blocking molecules are completely omitted, leading to rapid and immediate translation of the mRNA.

Shipping and Storage

Packaging and Storage	
Shipping conditions	Ambient temperature *
Storage conditions	-20°C
Shelf life	Under proper storage conditions as indicated on vial 3 months after opening

*Shipped with additional ice pack to assure temperature below 25°C.

Specifications

Specifications of Fuse-It-mRNA	
Formulation	Proprietary lipids
Concentration	6 mM
Fluorescence properties	
Ex _{max} /Em _{max}	750/780 nm (see Figure 1)
Quality Control	See Certificate of Analysis (on request)

Kit Contents

Fuse-It-mRNA easy comes as a kit containing the following components:

- Fusogenic Solution (FS): 2 vials, ready-to-use
- Neutralization Buffer (NB): 2 vials, ready-to-use

Note:

FS is solubilized in a low osmotic buffer (20 mM HEPES, pH7.4). Aliquoting of FS or NB is not necessary.

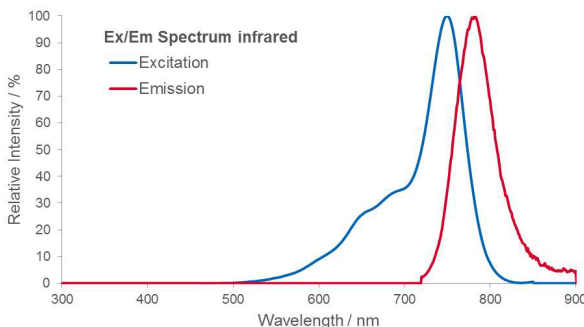


Figure 1: Spectrum of Fuse-It-mRNA fusion control dye.

Important Guidelines

- Fuse-It-mRNA easy is optimized for the transfer of RNA > 100 nt, e.g. mRNA. It is not recommended using Fuse-It-mRNA for the transfer of RNA < 100 nt, e.g. siRNA. In this case, use Fuse-It-siRNA (Cat. No. 60510, 60511) instead.
- Due to the infrared (IR) control dye, the fusion process can be visualized directly after fusion. Make sure to use the appropriate IR filter sets and an IR-sensitive camera. The IR fluorescence signal of the

liposomal carriers can also be applied for cell sorting using flow cytometry.

- Use high-quality, thin bottom cell culture materials to achieve the best imaging results (e.g., ibidi's μ -Slides and μ -Dishes).

Protocol: Transfection of mRNA Into Mammalian Cells

Material Required but Not Supplied

- mRNA (0.5–2 $\mu\text{g}/\mu\text{l}$) in RNase-free buffer. Find the necessary amount for various culture formats in Table 1.
- RNase-free micro reaction tubes
- 1 \times PBS (pH 7.0–8.0; osmolarity 250–320 mOs/kg)
Note: Divalent ions do not influence the fusion process.

Day 0

Preparation of Adherent Cells

- One day before transfection, seed the adherent cells and incubate them under standard conditions (typically 37°C and 5% CO₂). The cells should be 70% –90% optically confluent at time of transfection.

Preparation of Suspension Cells

- Prepare enough cells for the experiment. A cell number of 1–3 $\times 10^5$ cells is recommended for each mRNA transfection reaction.
Note: Higher cell numbers should be used for sensitive cells to achieve a less harsh fusion reaction.

Day 1

Important Notes

- All steps should be performed on ice, unless indicated otherwise.
- The cells should be in a healthy condition at the time of transfection.
- The volumes in this protocol (**marked in red**) refer to one single mRNA transfection of adherent cells in a μ -Dish^{35 mm, high} or 1–3 $\times 10^5$ suspension cells. Please calculate the respective volumes according to the experimental setup and culture format as indicated in Table 1.

Culture Format	Growth Area [†]	mRNA Amount	Neutralization Buffer (NB)	Fusogenic Solution (FS)	Transfection Mix	Culture Medium
μ-Dish 35 mm, high (ibidi)	3.5 cm ²	1.0 μg	2.0 μl	2.5 μl	250 μl	2 ml
μ-Slide 8 Well (ibidi)	1.0 cm ²	0.6 μg	1.2 μl	1.5 μl	150 μl	300 μl
μ-Plate 96 Well (ibidi)	0.55 cm ²	0.4 μg	0.8 μl	1.0 μl	100 μl	300 μl
96-well	0.3 cm ²	0.28 μg	0.56 μl	0.7 μl	70 μl	300 μl
48-well	0.75 cm ²	0.6 μg	1.2 μl	1.5 μl	150 μl	750 μl
24-well	1.9 cm ²	1.2 μg	2.4 μl	3.0 μl	300 μl	1 ml
12-well	3.8 cm ²	2.2 μg	4.4 μl	5.5 μl	550 μl	2 ml
6-well	9.5 cm ²	4.0 μg	8.0 μl	10.0 μl	1000 μl	3 ml
μ-Slide VI ^{0.4} (ibidi)	0.6 cm ²	0.2 μg	0.4 μl	0.5 μl	60 μl*	150 μl

[†]Growth area varies depending on the manufacturer.

*Please note that for each channel 60 μl are prepared and used for pipetting, while only 30 μl will remain in the channel.

Table 1: Volumes for mRNA transfection of adherent cells in different culture vessels. The transfection conditions must be adjusted in proportion to the relative surface area. Volumes and amounts are given on a per-well-basis.

Before Starting

- Thaw Neutralization Buffer (NB) and Fusogenic Solution (FS) at room temperature. Then place the vials on ice.
- Heat **2 ml** culture medium to 37°C.
- Cool 1× PBS to 4°C.

Preparation of Fusogenic Solution

1. Dilute the mRNA to a final concentration of **1 μg/μl** in RNase-free water. Mix by pipetting up and down several times.
Note: The mRNA concentration can range from 0.5 to 2 μg/μl, according to the experimental setup and the required mRNA amount.
2. Briefly vortex the Neutralization Buffer (NB). Into one tube, add **2 μl** of NB and **1 μg** of mRNA (e.g. 1 μl of a 1 μg/μl mRNA solution) and mix by pipetting up and down several times. This is the neutralized mRNA solution.
Note: Always keep the same ratio: 2 μl NB : 1 μg mRNA. Adjust according to the experimental setup and the required mRNA amount, as indicated in Table 1.
3. Incubate neutralized mRNA for 10 minutes at **room temperature**.
4. Vortex the Fusogenic Solution (FS), which contains the liposomal carriers, until the solution is homogeneous. If necessary, mix by pipetting up and down several times.
Note: The solution needs to be homogeneous at every step. Aggregates can affect the fusion efficiency!
5. Transfer 2.5 μl FS to the tube containing the neutralized mRNA solution and mix by pipetting up and down several times. This is the non-diluted transfection solution.
6. Add cooled 1× PBS to the non-diluted transfection solution to a final volume of **250 μl** and vortex briefly.
7. For adherent cells, continue with "A) Transfection of Adherent Cells", for suspension cells, continue with "B) Transfection of Cells in Suspension".



A) Transfection of Adherent Cells

1. Remove the culture medium and add the transfection solution drop-wise onto the cells.
2. Incubate for 5–20 minutes at 37°C until the cells start to detach.
Note: Reaching 37°C is critical for optimal fusion efficiency! Observe the cell morphology every 5 minutes to determine the optimal reaction time for the cells. The reaction should be stopped before signs of cell detachment are observed.
3. Remove the transfection solution and add fresh culture medium (equilibrated to 37°C) to the cells to stop the fusion process.
4. Incubate the cells under standard cell culture conditions (typically 37°C and 5% CO₂) before analysis of gene expression. The timepoint of analysis depends on the experimental setup, typically 2–96 hours after transfection.

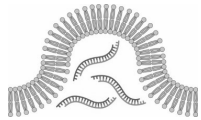
B) Transfection of Cells in Suspension

1. Centrifuge 1–3×10⁵ cells at 200 ×g for 5 minutes and discard the supernatant.
Note: For certain cell types, a higher centrifugation speed might be necessary for pelleting.
2. Resuspend the cell pellet in 50 µl of fresh cell culture medium containing 10% serum.
Note: The cell culture medium must contain 10% serum! The serum proteins protect the cells in suspension to achieve a less harsh fusion reaction.
3. Add the transfection solution to the resuspended cells and vortex briefly.
4. Incubate for 1–5 minutes at room temperature.
Note: Fusion is very efficient when using cells in suspension. Therefore, only a short incubation time is needed. The optimal incubation time has to be determined for each cell type.
5. Pellet the cells at 600–800 ×g for 3 minutes.
Note: 600–800 ×g is required for pelleting, depending on the cell type. At lower speed, the cells largely remain in the supernatant due to liposomal fusion.
6. Remove the transfection solution and resuspend the cells in fresh culture medium.
7. Incubate the cells under standard cell culture conditions (typically 37°C and 5% CO₂) before analysis of gene expression. The timepoint of analysis depends on the experimental setup, typically 2–96 hours after transfection.

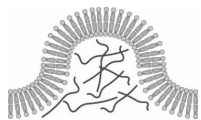
Troubleshooting Guide

Problem	Possible Cause	Recommendation
Cell viability is reduced after fusion and/or cells detach from the surface during or after fusion.	Incubation of sensitive cells in 1× PBS caused cell stress.	To reduce cell stress, cell culture medium can be used for the dilution of the transfection solution instead of 1× PBS. Note: For adherent cells, the use of a serum-containing medium may reduce the fusion efficiency.
	Cells were stressed or generally showed a weak adherence to the surface.	Reduce the incubation time with the transfection solution. A substrate coating may be helpful for sensitive cells.
	Cell density was too low.	Increase cell number. Optical confluence should range from 70% to 90%. The number of cells per transfection in suspension can be increased up to 1×10 ⁶ cells.
	Standard mRNA : FS ratio was not optimal for the cell type.	Optimize the mRNA : FS ratio for the used cell type. Increase the initial FS volume. Note: A more than twofold increase of FS can affect cell viability.
	mRNA was of poor quality or contained endotoxins.	Ensure that the stock solution of the mRNA is of high quality and reconstituted in RNase-free and endotoxin-free buffer, according to the manufacturer's recommendations.
	Transfection solution was not homogeneously added, resulting in accumulation of the transfection solution.	Add the transfection solution dropwise onto the adherent cells for a homogenous distribution.
Fusion efficiency, as indicated by the included infrared control dye, was low.	Remaining serum proteins from the culture medium reduced the fusion efficiency.	Wash the cells with 1× PBS prior to adding the transfection solution to the cells.
	Incubation time with transfection solution was too short.	Increase the incubation time to a maximum of 20 minutes for the fusion of adherent cells or to a maximum of 5 minutes for the fusion of suspension cells, respectively.
	Transfection solution was not homogeneously distributed during incubation.	Gentle motion during incubation will improve fusion efficiency.
	Cell density was not optimal.	For optimal fusion efficiency, optical confluence should range from 70% to 90%. If a lower cell density is required, adherent cells can also be transfected in suspension. This might enhance the fusion efficiency.

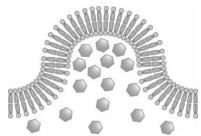
Problem	Possible Cause	Recommendation
	Fuse-It reagents were stored improperly.	Store Fuse-It reagents properly according to the section Shipping and Storage.
Despite high fusion efficiency, mRNA transfection efficiency was too low.	Low amount of mRNA was transferred.	Increase the initial mRNA amount, but keep the same ratio 10 pmol mRNA : 2 µl NB. Note: A more than fivefold increase of mRNA can affect cell viability.
	Incubation period of mRNA and NB was too short.	We recommend a neutralizing incubation time of 10 minutes. To optimize this process, incubate for up to 20 minutes.
	Degradation or poor quality of mRNA.	Ensure that the stock solution of the mRNA is of high quality and reconstituted in RNase-free and endotoxin-free buffer, according to the manufacturer's recommendations.
Transfection results are not reproducible.	Transfection was conducted using different cell confluencies.	Keep cell confluency constant throughout all experiments.
	Cells changed their morphology or behavior over time.	Transfection efficiency is dependent on passage number. Use similar passage numbers for each transfection.
	Wrong amount of mRNA, NB, or FS were used, due to pipetting errors.	When pipetting low volumes of mRNA, NB, or FS, it is recommended to prepare 1:10 dilutions in RNase-free water.
Protein expression was too low.	mRNA construct worked not efficient enough.	Make sure to use an mRNA-construct with a 5' cap and a 3' polyA-tail for efficient protein expression. If protein expression is still low, use a different promoter for higher levels of protein expression.

Ordering Information
Fuse-It-siRNA: Transfer of siRNA


Cat. No.	Description	Amount
60510	Fuse-It-siRNA, infrared fluorescent: ready to use, 6 mM	2 × 150 µl
60511	Fuse-It-siRNA, infrared fluorescent: ready to use, 6 mM	2 × 300 µl

Fuse-It-mRNA: Transfer of mRNA


Cat. No.	Description	Amount
60505	Fuse-It-mRNA infrared fluorescent: ready to use, 6 mM	2 × 150 µl
60506	Fuse-It-mRNA infrared fluorescent: ready to use, 6 mM	2 × 300 µl

Fuse-It-P: Transport of Proteins


Cat. No.	Description	Amount
60220	Fuse-It-P, infrared fluorescent: lyophilized, 3 mM	for 100 µl solution
60221	Fuse-It-P, infrared fluorescent: lyophilized, 3 mM	for 4 × 25 µl solution
60222	Fuse-It-P, infrared fluorescent: lyophilized, 3 mM	for 400 µl solution
60223	Fuse-It-P, infrared fluorescent: lyophilized, 3 mM	for 4 × 100 µl solution

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