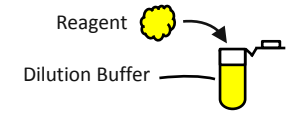
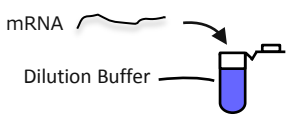


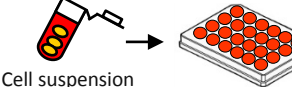
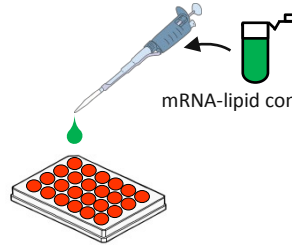
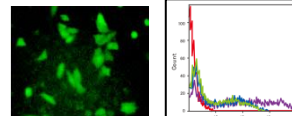


## ScreenFect™ A plus Transfection Protocol

The optimal condition for successful transfection varies. Start any new transfection by testing the recommended two concentrations of ScreenFect™ A plus Reagent to determine an optimum amount.

### 1-Step method (Reverse transfection method)

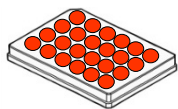
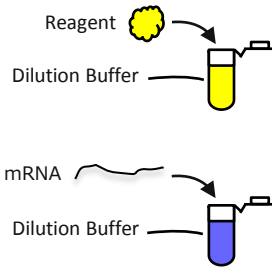

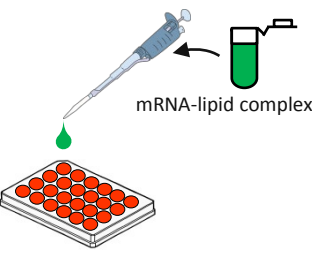
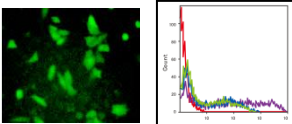
Timeline	Steps
<p><b>1</b></p>  <p>Reagent Dilution Buffer</p>	<p>Dilute ScreenFect™ A plus Reagent*1 in Dilution Buffer, and then mix well *1 Vortex the reagent before use</p>
	<p>Dilute mRNA in Dilution Buffer, and then mix well</p>  <p>mRNA Dilution Buffer</p>
<p><b>2</b></p>  <p>mRNA-lipid complex</p>	<p>Add diluted mRNA to diluted ScreenFect™ A plus Reagent, and then incubate for 5 minutes ~ at room temperature*2 *2 Incubation is available until the step 4 has been completed</p>
<p><b>3</b></p>  <p>Cultured cells</p>	<p>Prepare required cells for transfection</p>
<p><b>4</b></p>  <p>Cell suspension</p>	<p>Detach cells and prepare the cell suspension, and then transfer the required numbers of cell suspension to cell culture plate</p>
<p><b>5</b></p>  <p>mRNA-lipid complex</p>	<p>Add mRNA-lipid complex from step 2 to well of cell culture plate from step 4</p>
<p><b>6</b></p> 	<p>Visualize/analyze transfected cells</p>

Procedure Details		96-well		24-well		12-well		6-well	
Component									
Dilution Buffer for ScreenFect™ A plus		5 µL		25 µL		50 µL		125 µL	
mRNA : Transfection Reagent ratio		1 : 3	1 : 4	1 : 3	1 : 4	1 : 3	1 : 4	1 : 3	1 : 4
ScreenFect™ A plus Transfection Reagent		0.15 µL	0.2 µL	0.75 µL	1.0 µL	1.5 µL	2.0 µL	3.75 µL	5.0 µL
Dilution Buffer for ScreenFect™ A plus		5 µL		25 µL		50 µL		125 µL	
mRNA (0.1-2.5 µg / µL)		50 ng		250 ng		500 ng		1250 ng	
Diluted mRNA		5 µL		25 µL		50 µL		125 µL	
Diluted ScreenFect™ A plus Transfection Reagent		5 µL		25 µL		50 µL		125 µL	
Adherent cells or suspension cells		1.0-4.0 × 10 <sup>4</sup>		0.5-2.0 × 10 <sup>5</sup>		1.0-4.0 × 10 <sup>5</sup>		0.25-1.0 × 10 <sup>6</sup>	
Cell Detachment (Trypsin or Accutase®)									
Final composition [per well]		96-well		24-well		12-well		6-well	
mRNA-lipid complex		10 µL		50 µL		100 µL		250 µL	
mRNA amount		50 ng		250 ng		500 ng		1250 ng	
ScreenFect™ A plus Transfection Reagent used		0.15 or 0.2 µL		0.75 or 1.0 µL		1.5 or 2.0 µL		3.75 or 5.0 µL	
Medium volume		100 µL		500 µL		1000 µL		2000 µL	
Incubate cells for 1 day ~ at 37°C. Then, analyze transfected cells.									
For support, please visit the <a href="http://screenfect.jp">http://screenfect.jp</a>									

## ScreenFect™ A plus Transfection Protocol

The optimal condition for successful transfection varies. Start any new transfection by testing the recommended two concentrations of ScreenFect™ A plus Reagent to determine an optimum amount.

### 2-Step method (Forward transfection method)

Timeline		Steps	Procedure Details										
1	Day 0	 <p>Pre-Cultured cells</p>	<p>Seed cells to be 70-90% confluent at transfection</p>	Component	96-well		24-well		12-well		6-well		
				Adherent cells or suspension cells	1.0-4.0 × 10 <sup>4</sup>		0.5-2.0 × 10 <sup>5</sup>		1.0-4.0 × 10 <sup>5</sup>		0.25-1.0 × 10 <sup>6</sup>		
Seed cells to be 70-90% confluent at transfection. The medium replacement may improve a transfection efficiency in a 2-Step method.													
2		 <p>Reagent Dilution Buffer</p> <p>mRNA Dilution Buffer</p>	<p>Dilute ScreenFect™ A plus Reagent<sup>※1</sup> in Dilution Buffer, and then mix well  <sup>※1</sup> Vortex the reagent before use</p> <p>Dilute mRNA in Dilution Buffer, and then mix well</p>	Dilution Buffer for ScreenFect™ A plus		5 μL		25 μL		50 μL		125 μL	
				mRNA : Transfection Reagent ratio		1 : 3	1 : 4	1 : 3	1 : 4	1 : 3	1 : 4	1 : 3	1 : 4
				ScreenFect™ A plus Transfection Reagent		0.15 μL	0.2 μL	0.75 μL	1.0 μL	1.5 μL	2.0 μL	3.75 μL	5.0 μL
				Dilution Buffer for ScreenFect™ A plus		5 μL		25 μL		50 μL		125 μL	
				mRNA (0.1-2.5 μg / μL)		50 ng		250 ng		500 ng		1250 ng	
3	Day 1	 <p>mRNA-lipid complex</p>	<p>Add diluted mRNA to diluted ScreenFect™ A plus Reagent, and then incubate for 5 minutes ~ at room temperature<sup>※2</sup>  <sup>※2</sup> Incubation is available until the step 4 has been completed</p>	Diluted mRNA		5 μL		25 μL		50 μL		125 μL	
				Diluted ScreenFect™ A plus Transfection Reagent		5 μL		25 μL		50 μL		125 μL	
4		 <p>mRNA-lipid complex</p>	<p>Add mRNA-lipid complex from step 3 to well of cell culture plate from step 1</p>	Final composition [per well]		96-well		24-well		12-well		6-well	
				mRNA-lipid complex		10 μL		50 μL		100 μL		250 μL	
				mRNA amount		50 ng		250 ng		500 ng		1250 ng	
				ScreenFect™ A plus Transfection Reagent used		0.15 or 0.2 μL		0.75 or 1.0 μL		1.5 or 2.0 μL		3.75 or 5.0 μL	
				Medium volume		100 μL		500 μL		1000 μL		2000 μL	
5	Day 2 ~		<p>Visualize/analyze transfected cells</p>	Incubate cells for 1 day ~ at 37°C. Then, analyze transfected cells.									
				For support, please visit the <a href="http://screenfect.jp">http://screenfect.jp</a>									