

# TransIT-PRO® Transfection Reagent

## Quick Reference Protocol

Instructions for MIR 5720, 5730, 5740 and 5750

Full protocol, SDS and Certificate of Analysis available at [mirusbio.com/5740](http://mirusbio.com/5740)



### SPECIFICATIONS

|                   |   |
|-------------------|---|
| Storage           | Store TransIT-PRO® Reagent tightly capped at -20°C.<br><b>Before each use</b> , warm to room temperature and vortex gently. |
| Product Guarantee | 1 year from the date of purchase, when properly stored and handled.   |

### ► PLASMID DNA TRANSFECTION PROTOCOL



Full protocol and additional documentation available at [mirusbio.com/5740](http://mirusbio.com/5740)

#### Fill in volumes below based on total culture volume (Table 1).

##### A. Maintenance of cells

1. Split suspension CHO or 293cells 18–24 hours prior to transfection to ensure that cells are actively dividing at the time of transfection.
2. Culture overnight.

##### B. Prepare TransIT-PRO® Reagent:DNA complexes

1. Seed cells at a density of  $2 \times 10^6$  cells/ml immediately before transfection.
2. Warm TransIT-PRO® Reagent to room temperature and vortex gently.
3. Place \_\_\_ ml of serum-free medium (e.g. Opti-MEM® or Opti-PRO™ SFM)\* in a sterile tube.
4. Add \_\_\_ µg plasmid DNA. Mix gently by pipetting.
5. Add \_\_\_ µl of TransIT-PRO® Reagent. Mix gently by pipetting.
6. Incubate complexes at room temperature for recommended amount of time:  
For complexes that will be added to suspension 293 cells, incubate for 15-20 minutes.  
For complexes that will be added to suspension CHO cells, incubate for 5-10 minutes.

##### C. Distribute complexes to cells

1. Add TransIT-PRO® Reagent:DNA complexes to cultured cells.
2. Incubate cells for 2-14 days depending on cell type, culture temperature, nature of the protein, and detection method. For further optimization information, please see the full protocol.
3. Harvest cells and/or supernatant and assay as required.

\*When using TransIT-PRO® Reagent with the CHOgro® Expression System, form transfection complexes in CHOgro® Complex Formation Solution.

**Table 1.** Volume scaling worksheet for DNA transfections with TransIT-PRO® Transfection Reagent.

| Starting conditions per milliliter of complete growth medium |          |                      |                    |
|--|----------|----------------------|--------------------|
|  | Per 1 ml | Total culture volume | Reagent quantities |
| Serum-free Complex Medium                                    | 0.1 ml   | × _____ ml           | = _____ ml         |
| Plasmid DNA (1 µg/µl stock)                                  | 1 µl     | × _____ ml           | = _____ µl         |
| TransIT-PRO® Reagent   | 1 µl     | × _____ ml           | = _____ µl         |

### ► Transfection Optimization

Determine the best TransIT-PRO® Reagent:DNA ratio for each cell type. Start with 1 µl of TransIT-PRO® Reagent per 1 µg of DNA. Vary the concentration of TransIT-PRO® Reagent from 0.5–2 µl per 1 µg DNA to find the optimal ratio.

TransIT-PRO® Transfection Reagent is a key component of the CHOgro® Expression System (MIR 6260), which is an optimized platform for transient, high titer protein production in suspension CHO derived cells.

For additional optimization tips, see [full protocol](#).

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## Reagent Agent<sup>®</sup>

Reagent Agent<sup>®</sup> is an online tool designed to help determine the best solution for nucleic acid delivery based on in-house data, customer feedback and citations.

Learn more at: [mirusbio.com/ra](https://www.mirusbio.com/ra)

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