

CHOgro® Transfection and Titer Enhancer Kit

Quick Reference Protocol

Instructions for MIR 6225

Full protocol, SDS and Certificate of Analysis available at mirusbio.com/6225



SPECIFICATIONS

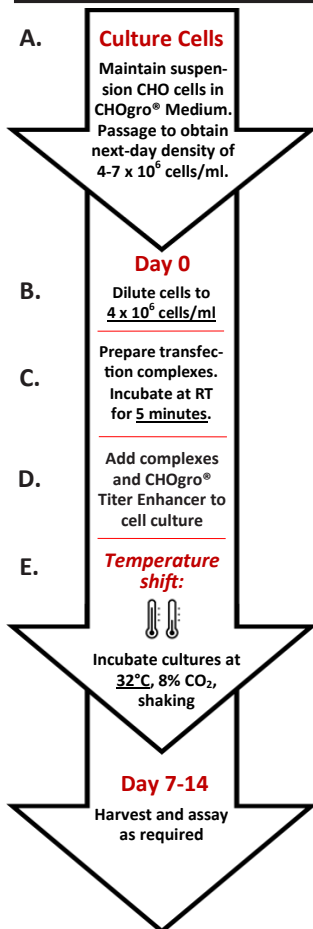
Storage	Store <i>TransIT-PRO</i> ® Transfection Reagent (MIR 5740) tightly capped at -20°C. Store CHOgro® Titer Enhancer (MIR 6220) at 2-10°C, protected from light. Before each use , warm to room temperature and vortex gently.
Product Guarantee	1 year from the date of purchase, when properly stored and handled.
Intended Usage	Designed for use with CHOgro® High Yield Expression System (MIR 6270)

► PLASMID DNA TRANSFECTION PROTOCOL



Full protocol and additional documentation available at mirusbio.com/6225

CHOgro® High Yield Expression System Flow Chart



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

A. Maintenance of cells

1. Passage cells 18–24 hours prior to transfection to obtain a next day density of 4-7 x 10⁶ cells/ml. DO NOT proceed with transfection if cells are not doubling every 24 hours or are < 98% viable.
2. Incubate cells overnight at 37°C in 8% CO₂, shaking.

B. Prepare/dilute suspension CHO cells for transfection

1. Seed cells at 4 x 10⁶ cells/ml immediately before transfection. DO NOT proceed if cells are not doubling normally or < 98% viable.

C. Prepare *TransIT-PRO*® Reagent:DNA complexes

1. Warm *TransIT-PRO*® Reagent to room temperature. Vortex gently.
2. Place ___ ml CHOgro® Complex Formation Solution in sterile tube.
3. Add ___ µg plasmid DNA. Mix gently by pipetting.
4. Add ___ µl of *TransIT-PRO*® Reagent. Mix gently by pipetting.
5. Incubate at room temperature for 5 minutes to allow transfection complexes to form.

D. Add transfection complexes and CHOgro® Titer Enhancer to cells

1. Add transfection complexes to cell culture. Mix gently by swirling.
2. Add ___ µl of CHOgro® Titer Enhancer to cell culture + transfection complexes. Mix gently by swirling.

E. Place culture at 32°C after transfection and enhancer addition

1. Incubate cultures at 32°C (8% CO₂, shaking) for 2-14 days.
NOTE: Optimal culture time will depend on cell type, protein of interest, culture temperature, and detection method.
2. Harvest cells and/or supernatant and assay as required.

Table 1. Scaling worksheet for CHOgro® Transfection and Titer Enhancer Kit.

Starting transfection conditions per milliliter of CHOgro® Expression Medium:			
	Per 1 ml	Total culture volume	Reagent quantities
Complex Formation Solution	0.1 ml ×	___ ml	= ___ ml
Plasmid DNA (1 µg/µl stock)	1 µl ×	___ ml	= ___ µl
<i>TransIT-PRO</i> ® Reagent	1 µl ×	___ ml	= ___ µl
Enhancer addition (add to culture after transfection complex addition):			
CHOgro® Titer Enhancer	20 µl ×	___ ml	= ___ µl

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► Critical Parameters for Success with CHOgro® High Yield Expression System

- **Cell adaptation and maintenance.** Cells grown in alternate media formulations should be fully adapted to CHOgro® Expression Medium supplemented with 4mM L-Glutamine and 0.3% Poloxamer 188 prior to transfection with the CHOgro® High Yield Expression System. Cells are fully adapted when they are ≥98% viable and doubling normally.
- **Cell density at transfection.** Cells should be passaged 18–24 hours prior to transfection to obtain a next day density of 4-7 x 10⁶ cells/ml. This allows for a minimal cell dilution for a final density of 4 x 10⁶ cells/ml at the time of transfection. Cultures should be placed at 37°C in 8% CO₂ prior to transfection. DO NOT proceed with transfection if cells are not doubling daily and at least 98% viable by trypan blue exclusion.
- **DNA concentration.** Start with 1 µg of DNA per 1 ml of culture. Vary the DNA concentration from 1–2 µg/ml to find the best working DNA concentration. To maintain the *TransIT-PRO*® Reagent:DNA ratio, adjust the reagent volume accordingly. NOTE: Use only high quality, endotoxin-free DNA for transfections. Contaminants such as protein, carbohydrate and lipids may affect transfection efficiency and gene expression levels. Ensure that the plasmid preparation exhibits an A260/A280 ratio of > 1.8.
- **Ratio of *TransIT-PRO*® Reagent to DNA.** Start with 1 µl of *TransIT-PRO*® Reagent per 1 µg of DNA. Vary the concentration of *TransIT-PRO*® Reagent from 1–2 µl per 1 µg of DNA to find the optimal ratio.
- **Transfection complex formation.** Prepare *TransIT-PRO*® Reagent:DNA complexes in CHOgro® Complex Formation Solution (MIR 6210). Incubate complexes at room temperature for 5 minutes to allow sufficient time for complexes to form. Following the 5 minute incubation, add transfection complexes directly to the flask of cells and swirl gently to mix thoroughly.
- **CHOgro® Titer Enhancer addition.** CHOgro® Titer Enhancer should be added to the culture immediately after transfection complex addition. Add 20 µl of CHOgro® Titer Enhancer per 1 ml cell culture (see Table 1 on front page for scaling chart). Cultures should then be placed at 32°C, 8% CO₂ (shaking) for the remainder of the culture.
- **Temperature shift to 32°C post-transfection.** Placing flasks at 32°C immediately post-transfection will increase overall protein titers and decrease protein degradation. Typically, greater than 2-fold higher antibody titers are achieved if incorporating the temperature shift into the production workflow.
- **Post-transfection incubation time.** The optimal post-transfection incubation time may vary depending on the experiment goal and the nature of the plasmid used. For secreted antibody constructs, optimal titers are obtained at 32°C at 7-14 days post-transfection in batch culture.



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