293 Cells, SFM adapted

Catalog Numbers 11625019, 11631017

Pub. No. MAN0007367  Rev. B.0

WARNING! Read the Safety Data Sheets (SDSs) and follow the handling instructions. Wear appropriate protective eyewear, clothing, and gloves. Safety Data Sheets (SDSs) are available from thermofisher.com/support.

CAUTION! Human origin materials are non-reactive (donor level) for anti-HIV 1 & 2, anti-HCV, and HBsAg. Handle in accordance with established bio-safety practices.

Product description
293-F and 293-H Cells are clonal isolates selected for their superior serum-free growth and transfection efficiency derived from adenovirus type 5 transformed embryonal human kidney cells expressing the E1A adenovirus gene. The 293-F strain is a fast-growing variant of the 293 cell line. The 293-H strain is a variant, which when grown in serum supplemented medium, demonstrates better adherence in monolayer culture and ease of use for plaque assays and other anchorage dependent applications. Both 293-F and 293-H cell lines can be adapted to serum-free suspension culture in 293 SFM II or CD 293 Medium supplemented with L-glutamine for recombinant protein production and adenovirus propagation. 293-F and 293-H cells as provided are adapted to CD 293 Medium.

Contents and storage

<table>
<thead>
<tr>
<th>Contents</th>
<th>Cat. No.</th>
<th>Amount</th>
<th>Storage</th>
</tr>
</thead>
<tbody>
<tr>
<td>293-F Cells, SFM adapted</td>
<td>11625019</td>
<td>1 vial; contains ≥7.5 × 10^6 cells</td>
<td>Liquid nitrogen vapor-phase</td>
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<tr>
<td>293-H Cells, SFM adapted</td>
<td>11631017</td>
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</tbody>
</table>

Important guidelines for thawing and storing cells
- Upon receipt, immediately thaw cells or place into vapor-phase liquid nitrogen storage until ready to use. Do not store the cells at >80°C.
- Avoid short-term extreme temperature changes. When storing cells in liquid nitrogen after shipping on dry ice, allow the cells to remain in liquid nitrogen for 3–4 days before thawing.
- Prior to starting experiments, ensure you have established cells and have frozen stocks on hand. Upon receipt, grow and freeze multiple vials of cells to ensure that you have an adequate supply of early-passage cells.

Important information
Cells are stable when maintained at −200°C to −125°C in liquid nitrogen vapor phase.

Culture conditions
CD 293 Medium and 293 SFM II require aseptic supplementation with L-glutamine or GlutaMAX™-I prior to use. Add 20 mL/L GlutaMAX™-I (200mM) or L-glutamine, 4 mM final concentration, to the medium before use.

Media: Complete 293 SFM II or CD 293 Medium, supplemented with 4 mM GlutaMAX™-I or L-glutamine.

Cell line: 293 human embryonic kidney cells.

Culture type: Stationary- adherent monolayer; or Suspension- orbital shaker platform rotating at 120–130 rpm.

Culture vessels T-flasks or shake flasks.

Temperature range: 36°C to 38°C.

Incubator atmosphere: Humidified atmosphere of 7–9% CO2 in air. Ensure proper gas exchange and minimize exposure of cultures to light.

Note: Antibiotics are not recommended; however 2.5–5 mL/L of 5000U/5000 μg Penicillin-Streptomycin may be used when required.

Recovery
1. Rapidly thaw (<1 minute) frozen vial of cells in a 37°C water bath.
2. Transfer the entire contents of the cryovial into a 125 mL shake flask containing 30 mL pre-warmed complete 293 SFM II or CD 293 Medium for suspension culture.
3. Incubate at 37°C in a humidified atmosphere of 7–9% CO2 in air. Loosen flask caps to allow for gas exchange.
4. Subculture cells 3–5 days post thaw when viable cell density reaches 2 × 10^6–3 × 10^6 viable cells/mL. Cell viability should be ≥90% and growth rate in mid-logarithmic phase.

Note: We recommend subculturing cells for a minimum of 3 passages before use in other applications or transfer into serum-supplemented media.

Subculture cells
293 cells may be subcultured either in suspension culture or as adherent monolayers depending upon the application.
1. Subculture cells 3–5 days post thaw or when cultures reach 60–80% confluence (adherent monolayer), or a viable cell density of 2 × 10^6–3 × 10^6 viable cells/mL (suspension). Ensure that cell viability is ≥90% and growth rate is in mid-logarithmic phase. For suspension culture proceed to step 3.
2. Displace cells from the flask surface by rapping sharply against your hand.
3. Transfer cell suspension to a sterile conical tube. Ensure cells are dispersed in a monocellular suspension by gently pipetting up and down or vortexing for up to 40 seconds.
4. Determine total viable cell density using a Countess™ Automated Cell Counter (alternate automated or manual methods may be used).
5. Seed cells into fresh prewarmed complete medium at the appropriate density for your chosen culture method.
   a. Adherent: 2 × 10^4–5 × 10^4 viable cells/cm^2 in 2–3 mL fresh complete pre-warmed medium per 10 cm^2.

For Research Use Only. Not for use in diagnostic procedures.
You may scale up the 293 cultures in spinner flasks or bioreactors using the following guidelines.

**Note:** 293 suspension cultures may grow as 2–10 cell clusters. Vigorous vortexing for up to 40 seconds may be required at each subculture for a number of passages until the cultures grow as a single cell suspension.

### Scaling-up 293 cell cultures

You may scale up the 293 cultures in spinner flasks or bioreactors using the following guidelines.

- We recommend using 293 SFM II for high-density suspension culture of 293 cells.
- Determine optimal seeding density for each system. We recommend seeding densities of $3 \times 10^5$–$5 \times 10^5$ viable cells/mL.

**Note:** If the split ratio of cells to fresh media is $<1:2$, centrifuge cell suspension at 200 x g for 5 minutes and resuspend the cell pellet in fresh 293 SFM II prior to inoculating the spinner or bioreactor culture.
- Optimize the spinner or impeller speed for your bioreactor depending on your needs.
- Higher stirring speeds and/or impeller design may necessitate decrease per minute).

Prepare the desired quantity of 293 cells in suspension culture, depending on your needs.

#### Suspension: $3 \times 10^5$ viable cells/mL fresh complete pre-warmed medium.

#### Preparation of 293 cells

1. Prepare the desired quantity of 293 cells in suspension culture, harvesting in mid-log phase of growth, viable cell density of $0.5 \times 10^6$–$1 \times 10^6$ viable cells/mL, and viability >90%. Retain the conditioned medium to prepare cryopreservation medium.
2. Determine the viable cell density and calculate the required volume of cryopreservation medium to give a final cell density of $\geq 5 \times 10^5$ cells/mL.
3. Prepare the required volume of cryopreservation medium of 92.5% growth media (50:50 ratio of fresh to conditioned media) +7.5% DMSO and store at 4°C until use.

**IMPORTANT!** Prepare cryopreservation medium on the day of intended use.

4. Harvest cells by centrifugation at 200 x g for 5 minutes. Discard the supernatant and resuspend the cell pellet in the pre-determined volume of 4°C cryopreservation medium.
5. Dispense aliquots of this suspension into cryovials according to the manufacturer’s specifications (i.e., 1 mL in a 2-mL cryovial).
6. Achieve cryopreservation in an automated or manual controlled rate freezing apparatus following standard procedures (1°C decrease per minute).
7. Transfer frozen cells to liquid nitrogen (vapor phase); we recommend storage at $-200°C$ to $-125°C$.

#### Cryopreservation

**Limited product warranty**


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**Transfection**

Several days prior to transfection, we recommend transferring cells to adherent (monolayer) culture in serum-supplemented media (i.e., Dulbecco’s Modified Eagle Medium supplemented with 0.1 mM MEM Non-Essential Amino Acids and 10% Fetal Bovine Serum). Do not add antibiotics to the media during transfection. Cells should adapt directly into the serum-supplemented media without any trouble. After transfection and selection, 293-F or 293-H cells can be expanded and re-adapted back into serum-free suspension culture in 293 SFM II or CD 293 Medium supplemented with 4 mM L-glutamine or GlutaMAX™-I and the appropriate selective antibiotic. For optimal transfection results, we recommend using Lipofectamine™ 2000 or 293Fectin™ Transfection Reagent. Refer to the accompanying product manuals for instructions. Other transfection reagents may be used if desired.

**Note:** CD 293 and 293 SFM II can inhibit complex formation of DNA with some cationic lipid transfection reagents (e.g., Lipofectamine™ 2000, 293Fectin™, Lipofectamine™, and Lipofectin™). If you are using one of these transfection reagents, culture cells in another medium immediately prior to and during the transfection.

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**Related products**

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<td>293 SFM II</td>
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<tr>
<td>CD 293 Medium</td>
<td>11913</td>
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<tr>
<td>L-Glutamine-200 mM (100X), Liquid</td>
<td>25030</td>
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<tr>
<td>GlutaMAX™</td>
<td>35050</td>
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<tr>
<td>Pluronic™ F-68, 10% (100X)</td>
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<td>Penicillin-Streptomycin, Liquid</td>
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<td>Gentamicin</td>
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<td>DMEM, high glucose</td>
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<td>MEM Non-Essential Amino Acids (100X), Liquid</td>
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<td>Qualified FBS, US</td>
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<td>293Fectin™ Transfection Reagent</td>
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<td>Lipofectamine™ 2000 Transfection Reagent</td>
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<td>Trypan Blue Stain</td>
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<tr>
<td>Countess™ Automated Cell Counter</td>
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**Corporate entity:** Life Technologies Corporation | Carlsbad, CA 92008 USA | Toll Free in USA 1 800 955 6288

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