

10X TeSR[™] cloning and single-cell survival supplement

Catalog #05888	
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10 mL 5 x 10 mL



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Product Description

CloneR[™] is a defined, serum-free TeSR[™] supplement designed to increase the cloning efficiency and single-cell survival of human embryonic stem (ES) cells and induced pluripotent stem (iPS) cells. CloneR[™] enables the robust generation of clonal cell lines without single-cell adaptation, thus minimizing the risk of acquiring genetic abnormalities.

Properties

Storage:Store at -20°C.Shelf Life:Stable for 12 months from date of manufacture (MFG) on label.

Materials Required But Not Included

PRODUCT NAME	CATALOG #
Vitronectin XF™	07180
OR	OR
CellAdhere™ Laminin-521	77003
OR	OR
Corning® Matrigel® hESC-Qualified Matrix	Corning 354277
CellAdhere™ Dilution Buffer	07183
OR	OR
DMEM/F-12 with 15 mM HEPES	36254
Enzymatic dissociation reagent (e.g. ACCUTASE™)	e.g. 07920
mTeSR™1	05850
OR	OR
TeSR™-E8™	05940

Preparation of Reagents and Materials

Coating Cultureware with Cloning Matrix

Coat cultureware with cloning matrix; use Vitronectin XF[™], CellAdhere[™] Laminin-521, or Corning[®] Matrigel[®] hESC-Qualified Matrix. NOTE: For Vitronectin XF[™], use non-tissue culture-treated cultureware; for the other matrices, use tissue culture-treated cultureware. NOTE: If using CellAdhere[™] Laminin-521, coat cultureware the day before cloning, as overnight incubation is required.

- 1. Thaw cloning matrix according to the applicable Product Information Sheet.
- 2. Dilute cloning matrix according to Table 1.



Table 1. Dilution and Incubation of Cloning Matrix

CLONING MATRIX	DILUENT	FINAL CONCENTRATION OF CLONING MATRIX	INCUBATION CONDITIONS
Vitronectin XF™	CellAdhere™ Dilution Buffer	10 µg/mL	Room temperature (15 - 25°C) for 1 hour
CellAdhere™ Laminin-521	CellAdhere™ Dilution Buffer	10 µg/mL	2 - 8°C overnight
Corning® Matrigel®	DMEM/F-12 with 15 mM HEPES	1 in 100 dilution (following manufacturer's reconstitution instructions)	Room temperature (15 - 25°C) for 1 hour

- 3. Add diluted cloning matrix to cultureware as follows:
 - 10 cm dish: 6 mL
 - 96-well plate: 50 µL/well

NOTE: For other cultureware, use 100 $\mu L/cm^2.$

- 4. Tilt plate back and forth to distribute matrix solution evenly.
- 5. Incubate cultureware as indicated in Table 1.

Cloning Medium

The following example is for preparing 25 mL of cloning medium. If preparing other volumes, adjust accordingly.

- Thaw CloneR[™] at room temperature (15 25°C).
 NOTE: If not used immediately, aliquot and store at -20°C. Once thawed, use immediately. Do not re-freeze.
 - Prepare complete mTeSR™1 or TeSR™-E8™.
 NOTE: For complete instructions, refer to the applicable Product Information Sheet (Document #29917 [mTeSR™1] or #28021 [TeSR™-E8™]).
- 3. Add 2.5 mL of CloneR[™] to 22.5 mL of complete mTeSR[™]1 or TeSR[™]-E8[™]. Mix thoroughly. NOTE: If not used immediately, store cloning medium at 2 8°C for up to 1 week.

Directions for Use

Use ES or iPS cells cultured in mTeSR™1 or TeSR™-E8™ at day 6 - 8.

Please read the entire protocol before proceeding.

Use sterile techniques when performing the protocols below.

- A. PREPARING CULTUREWARE FOR CLONING
- 1. Aspirate cloning matrix from coated plates (see Preparation of Reagents and Materials).
- 2. Add cloning medium to cultureware as follows (initial seed volume):
 - 10 cm dish: 8 mL
 - 96-well plate: 100 µL/well

NOTE: For other cultureware, use 150 $\mu L/cm^2.$

- 3. Incubate at 37°C for 1 hour prior to seeding cells.
- B. PREPARING A SINGLE-CELL SUSPENSION
- 1. Remove ES/iPS cell culture from incubator.
- 2. Under the microscope, mark regions of differentiation using a marker pen.
- 3. Remove regions of differentiation by aspiration. Rinse with PBS and aspirate.
- Add enzymatic dissociation reagent (e.g. ACCUTASE[™]) at room temperature (15 25°C) at 1 mL per 10 cm² surface area. Incubate at 37°C for 5 - 8 minutes.

NOTE: Dissociation time may be cell line-dependent.

CloneR™



- 5. Rinse cells from cultureware by pipetting the dissociation reagent up and down, dispensing onto the colonies and breaking them up.
- 6. Dilute cell suspension 1 in 4 by adding to a conical tube (e.g. Falcon® Conical Tube, 15 mL, Catalog #38009) containing complete mTeSR™1 or TeSR™-E8™.
- 7. Create a single-cell suspension by flicking the tube 3 5 times. Count cells using a hemocytometer or other cell counting method. Proceed to section C.

C. PLATING CELLS (Day 0)

Plating at clonal density

- 1. Centrifuge the single-cell suspension from section B at 300 x *g* for 5 minutes. Aspirate supernatant and resuspend cells in cloning medium.
- 2. Add desired number of cells to the cultureware containing warm cloning medium from section A.
- 3. Swirl the plate and rock back and forth 4 5 times to distribute cells evenly. Incubate at 37°C for 2 days. Proceed to section D.

OR

Single-cell sorting (96-well plates)

- 1. Centrifuge the single-cell suspension from section B at 300 x *g* for 5 minutes. Aspirate supernatant and resuspend cells at 1 x 10^6 cells/mL in cloning medium.
- 2. Pass the single-cell suspension from section B through a 40 µm Cell Strainer (Catalog #27305) to remove any large clumps.
- Sort cells into individual wells of the plate containing warm cloning medium (from section A) using a fluorescence-activated cell sorter (FACS) (as low as 1 cell/well).
- 4. Incubate at 37°C for 2 days. Proceed to section D.

D. FEEDING CELLS

- 1. Day 2: Perform a full medium change with cloning medium. Incubate at 37°C for 24 hours.
- 2. Day 3: Add cloning medium (25% of initial seed volume). Incubate at 37°C for 24 hours.
- 3. Day 4 10/14: Perform a full medium change with mTeSR[™]1 or TeSR[™]-E8[™] (without CloneR[™]) daily until colonies are ready to be picked.

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