CULTREX[®] Product Data

For Research Use Only. Not For Use In Diagnostic Procedures

Cultrex[®] Bovine Collagen I

Catalog: 3442-050-01

Size: 50 mg

Description: Type I collagen is the major structural component of extracellular matrices found in connective tissue and internal organs, but is most prevalent in the dermis, tendons, and bone. It is a 300 kDa molecule composed of two alpha₁(I) chains and one alpha₂(I) chain that spontaneously forms a triple helix scaffold at a neutral pH and 37 °C. This phenomenon can be exploited to promote cell attachment, growth, differentiation, migration, and tissue morphogenesis during development.

Specifications:

Concentration:	Type I Collagen is provided at 5 mg/mL (Sircol Assay).
Source:	Fetal Bovine Extensor Tendons
Storage Buffer:	20 mM Acetic Acid
Storage/Stability:	Product is stable for a minimum of 3 months if stored at 4 °C. Do Not
	Freeze.

Materials Qualification:

Gelling:

• Type I Collagen forms a firm gel at neutral pH and 37 °C when diluted to 0.4 mg/ml.

Functional Assavs:

 Tested for the ability to promote cell attachment and spreading of HT-1080 human fibrosarcoma cells.

Sterility Testing:

- No bacterial or fungal growth detected after incubation at 37 °C for 14 days following USP XXIV Chapter 71 sterility test.
- No mycoplasma contamination detected by PCR.
- Endotoxin concentrations ≤ 20 EU/ml by LAL assay.

Gelling Procedures:

Note: To prevent contamination maintain aseptic techniques in a laminar flow biological hood throughout this procedure. Working with solutions that are pre-chilled at 4°C, and keeping solutions on ice extends the time that Collagen I will remain in solution after neutralization.

- 1. Place the following on ice:
 - a. Type I Collagen (5 mg/ml)
 - b. Sterile 10X PBS
 - c. Sterile, distilled water (dH₂O)
 - d. Sterile 1N NaOH (fresh)
- 2. Determine the concentration and final volume of Collagen I needed for experimentation.

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Gelling Procedures (cont.):

- 3. Determine the amount of reagents needed so that Collagen I is at the desired concentration in 1X phosphate buffered saline (PBS), neutralized by 1N NaOH.
 - a. Volume of Collagen needed= (Final conc. of Collagen) x (Total Volume)
 - Initial conc. of Collagen
 - b. Volume of 10X PBS needed= <u>Total Volume</u>

10

- c. Volume of 1N NaOH needed= (Volume of Collagen) x 0.023 ml
- d. Volume of dH₂O needed = Total Volume (sum of volumes from steps A+B+C)
- 4. In a sterile tube mix the 10X PBS, 1N NaOH and dH_2O .
- 5. Add the Collagen I to the tube and pipet up and down to mix (do not vortex).
- Place the Collagen solution into the desired plates or dishes. This solution is stable for up to 1 hour on ice. Plates may be centrifuged 300 x g for 10 minutes at 4°C to prevent bubbles from forming in the gel.
- 7. Incubate the plate at 37 °C for 1 hour to promote gel formation.

For your cell type, a gelling procedure using 7.5% (w/v) Sodium Bicarbonate for neutralization may be preferred:

- 1. Place the following on ice:
 - a. Type I Collagen (5 mg/ml)
 - b. Sterile 10X PBS
 - c. Sterile, distilled water (dH₂O)
 - d. 7.5% Sodium Bicarbonate, sterile
- 2. Determine the concentration and final volume of Collagen needed for experimentation.
- Determine the amount of reagents needed so that Collagen I is at the desired concentration in 1X phosphate buffered saline (PBS), neutralized by 7.5% sodium bicarbonate.
 - a. Volume of Collagen needed= (Final conc. of Collagen) x (Total Volume)

(Initial conc. of Collagen I)

b. Volume of 10X PBS needed= <u>Total Volume</u>

10

- volume of 7.5% sodium bicarbonate needed = (Volume of Collagen I, step a) x 0.0125 ml
- d. Volume of dH_2O needed = Total Volume (sum of volumes from steps A+B+C)
- 4. In a sterile tube mix the 10X PBS, and dH₂O and 7.5% sodium bicarbonate.
- 5. Add the Collagen I to the tube and pipette up and down to mix (do not vortex).
- Place the neutralized Collagen I solution into the desired plates or dishes. This solution is stable for up to 1 hour on ice. Plates may be centrifuged 300 x g for 10 minutes at 4°C to prevent bubbles from forming in the gel.
- 7. Incubate the plate at 37 °C for 1 hour to promote gel formation.

High Concentration Collagen gel method:

- 1. Place Collagen I (5 mg/ml), 7.5% sodium bicarbonate solution, sterile tube and cell culture plate on ice.
- 2. Add necessary amount of Collagen I into sterile tube.
- 3. Add 5 µl of 7.5% sodium bicarbonate per 0.1 ml of Collagen I (5 mg/ml)
- 4. Pipette Collagen I up and down to mix. (Do not vortex.)
- Place neutralized collagen into a cell culture plate. Plate may be centrifuged for 300 x g for 10 minutes at 4 °C to prevent bubbles from forming in the gel.
- 6. Incubate the plate at 37 °C for 1 hour to promote gel formation.

Thin Coating Procedure:

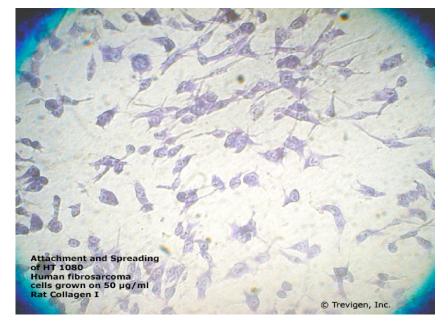
Optimization for desired protein concentration may be required. A starting concentration of

5 μ g per cm² is recommended. Increasing the temperature of acidic Collagen I will decrease viscosity. It is recommended that collagen is separated into aliquots prior to warming to maximize shelf life. Aliquots may be warmed to 37°C for up to 5 minutes or 25°C for up to 30 minutes prior to diluting.

- 1. Determine the volume needed for experimentation.
- 2. Dilute the Collagen to 50 µg/ml in 0.02 M acetic acid at the final volume needed.
 - a. Volume of Collagen= <u>(50 µq/ml of Collagen) x (Final Volume)</u> (Initial Concentration of Collagen)

b. Volume of 0.02 M acetic acid= Final Volume - Volume of Collagen (Step A)

- Add solution to plates or dishes at 5 μg per cm² (e.g. 50 μg, or 1 ml of 50 μg/ml, of Collagen is required for coating a 35 mm dish, which has a surface area of approximately 10 cm²).
- 4. Incubate at 37°C for 1 hour.
- 5. Carefully aspirate solution from the well or dish.
- 6. Rinse dish three times with equal volumes of PBS or media to remove the acid.
- 7. Plates may be used immediately or air dried for future use.



References:

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Related Products:

Itelatea I Ioaa			
Catalog#	Description	Size	
3415-001-02	Cultrex [®] Human BME, PathClear [®]	1 ml	
3432-005-02	Cultrex [®] BME, PathClear [®]	5 ml	
3432-005-01	Cultrex [®] BME without Phenol Red	5 ml	
3431-005-01	Cultrex [®] BME with Phenol Red; Reduced Growth Factors	5 ml	
3433-005-01	Cultrex [®] BME; no Phenol Red; Reduced Growth Factors	5 ml	
3430-005-02	Cultrex [®] BME with Phenol Red, PathClear [®]	5 ml	
3431-005-02	Cultrex [®] BME with Phenol Red, Reduced Growth Factor PathClear [®]	5 ml	
3400-010-01	Cultrex [®] Mouse Laminin I	1 mg	
3440-100-01	Cultrex [®] Rat Collagen I	100 mg	
3410-010-01	Cultrex [®] Mouse Collagen IV	1 mg	
3420-001-01	Cultrex [®] Human Fibronectin, PathClear [®]	1 mg	
3416-001-01	Cultrex [®] Bovine Fibronectin, NZHD*	1 mg	
3421-001-01	Cultrex [®] Human Vitronectin, PathClear [®]	50 µg	
3417-001-01	Cultrex [®] Bovine Vitronectin, NZHD	50 µg	
3438-100-01	Cultrex [®] Poly-L-Lysine	100 ml	
3439-100-01	Cultrex [®] Ploy-D-Lysine	100 ml	
3445-048-01	Cultrex [®] 3-D Culture Matrix [™] BME	15 ml	
3446-005-01	Cultrex [®] 3-D Culture Matrix [™] Laminin I	5 ml	
3447-020-01	Cultrex [®] 3-D Culture Matrix [™] Collagen I	100 mg	
*New Zealand Herd-Derived			



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