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GelNestTM Matrix, for hESC Culture, LDEV-Free

Product overview

GelNestTM Matrix is prepared from basement membrane components extracted from mouse tumor tissues. The main components include laminin, type IV collagen, heparan sulfate proteoglycan, etc. These components can provide the support and signals required for cell adhesion, differentiation, and proliferation. They can also simulate the characteristics of the basement membrane in a physiological environment and improve the success rate and effect of cell culture.

In addition to basement membrane components, GelNestTM Matrix is also rich in a variety of growth factors. These growth factors can promote cell differentiation, proliferation, and migration, further mimicking cell signaling pathways and interactions in physiological environments. GelNestTM Matrix has a wide range of application prospects, especially in tissue engineering, cell culture and research. It can be used for research on organoid culture, stem cell differentiation, angiogenesis, migration or invasion, and *in vivo* tumorigenesis.

Product information

Product number	Product name	Packaging specifications
211272	GelNest TM Matrix, for hESC Culture,	Bag Package, 5 mL/bottle, 1
	LDEV-Free	bottle/bag

Product parameters

Source	Mouse tumor tissue basement membrane components	
Formulation*	With phenol red	
Protein concentration	Please refer to the COA/COC for batch-specific protein concentration.	
Appearance	GelNest™ Matrix is liquid at 4°C but forms a gel at 37°C.	
Applications	Validated with ESC culture experiment. Suitable for feeder-free hESC	



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	and iPSC culture.
Storage and shelf life	It is recommended to aliquot the melted product into single-use portions
Storage and shell life	and store it in a -80°C freezer. The product has a shelf life of 2 years.
Precautions	GelNest™ Matrix will start to solidify when the temperature is
Frecautions	higher than 10°C. Please operate on ice.

Experimental procedures

Please determine the specific experimental steps based on cell types, culture conditions, and application experience.

Feeder-free culture of human embryonic stem cells (hESCs) and induced pluripotent stem cells (iPSCs)

- 1. Take the GelNestTM Matrix from frozen storage and thaw in an ice bath at 4°C overnight. Use a pre-cooled pipette tip to slowly pipette the matrix gel 3 times to mix. Use pre-cooled pipette tips to aliquot the thawed matrix gel. If bubbles form, briefly centrifuge the aliquoted matrix gel using a handheld centrifuge to remove bubbles.
- 2. Place the cell culture plate in the 37°C incubator to preheat.
- 3. Dilute the matrix gel solution at a ratio of 1:100 with serum-free medium that is precooled at 4°C, and completely cover the culture plate with the matrix gel diluent. It is recommended to use 300µL/cm² of matrix gel diluent in a culture dish.
- **4.** Allow the culture plate containing the diluted matrix gel to sit at room temperature for 1 hour.
- 5. Remove the remaining matrix gel diluent and immediately seed the stem cells with premixed mTeSR solution onto the culture plate. Be careful not to let the modified culture plate surface dry out.





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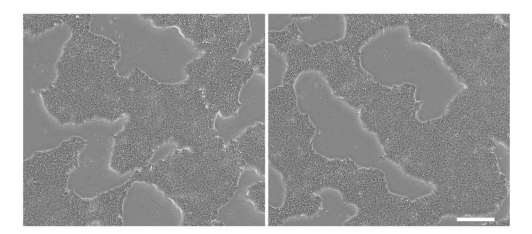


Figure 1. Results of hESC grown for 3 days on dishes coated with Brand C matrix gel (left) and GelNestTM Matrix gel (right). Scale bar is 300μm.

Safety recommendations and limitations

Please follow good laboratory safety practices.

For research use only. Not intended for diagnostic or therapeutic purposes. Contains ingredients of animal origin.

Technical support and contact information

For FAQ, GelNestTM Matrix Selection Guide, Quality Assurance COA/COC or other technical support and product issues, please refer to our website or use the following contact information.

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