CORNING

Certificate of Analysis

Heat Inactivated **Product:** Fetal Bovine Serum **Description:**

Premium,

United States Origin

Catalog Number: Lot Number: 35-016-CV 03822002

Date of Heat **Heat Inactivation** Individual Bottle 10 FEB 2022 Inactivation: Inactivation Type:

Product Prior to Heat Heat Inactivation 56°C (±2°C) for 30

Testing Performed on: Temperature: Inactivation minutes

Parent Catalog Number: Parent Lot Number: 35-015-CV 22821001

Date of Parent Lot 16 AUG 2021 Date of Expiration: AUG 2026

Manufacture:

Country of Raw Material Country of Final United States United States Origin: **Processing:**

Sterility Assurance 10⁻³ (Triple 0.1 μm Storage: -40°C to -10°C Level: Filtered)

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201 1007 4 21 210111	U/mL
Hemoglobin USP <90> ≤ 25 8.69 mg Barile and Kern & N. D. J.	g/dL
Mycoplasma Hoechst Fluorochrome Stain Not Detected Not Detected N/	/A
Osmolality USP <785> & EP 2.2.35 260 – 350 282 mC	Osm/kg H ₂ O
pH USP <791> & EP 2.2.3 6.5 – 8.5 7.02 N//	/A
Sterility (Bacteria & Fungi) USP <71> & EP 2.6.1 No Growth No Growth N/	/A
Total Protein USP <1057> & EP 2.5.33 3.0 – 4.5 3.5 g/c	dL
Gamma-Glutamyl Transferase (GGT) Chemistry Analyzer ≤ 10 4 IU/	J/L
Cell Growth Performance Testing MTT Assay Passed Passed N/A	/A
Identification (Species ID) EP 2.7.1 Bovine Bovine N/A	/A
Virus Testing 9 CFR Part 113.53c	
Bluetongue Virus Not Detected Not Detected N/	/A
Bovine Adenovirus 1 & 5 Not Detected Not Detected N/A	/A
Bovine Parvovirus Not Detected Not Detected N/A	/A
Bovine Respiratory Syncytial Virus Not Detected Not Detected N/	/A
Bovine Viral Diarrhea Virus As Reported Not Detected N/A	/A
Rabies Virus Not Detected Not Detected N/	/A
Reovirus Not Detected Not Detected N/	/A
Cytopathogenic Agents (IBR) Not Detected Not Detected Not Detected	/A
Hemadsorbing Agents (PI3) Not Detected Not Detected Not Detected	/A
BVDV Serum Neutralization Alpha Antibody Test	
BVDV Antibody Type 1 As Reported 10 ^{2.95} TC	CID ₅₀ /mL
BVDV Antibody Type 2 As Reported No Neutralization N/	/A
Biochemical Testing - Proteins & Organic Compounds	
Alkaline Phosphatase As Reported 207 IU/	J/L
Blood Urea Nitrogen (BUN) As Reported 14 mg	g/dL
	g/dL
Creatinine As Reported 2.6 mg	g/dL

Glucose		As Reported	93	mg/dL
Glutamic Oxaloacetic Transaminase (AST)		As Reported	44	IU/L
Glutamic Pyruvic Transaminase (ALT)		As Reported	6	IU/L
High Density Lipoprotein Cholesterol (HDL)		As Reported	8	mg/dL
Immunoglobulin G (IgG)	ELISA	As Reported	59.3	μg/mL
Low Density Lipoprotein Cholesterol (LDL)		As Reported	6	mg/dL
Total Bilirubin		As Reported	0.3	mg/dL
Triglyceride		As Reported	71	mg/dL
Uric Acid		As Reported	2.5	mg/dL
Biochemical Testing - Electrophoretic Profile	Cellulose Acetate Electrophoresis	Normal	Normal	N/A
Albumin		As Reported	2.0	g/dL
Alpha		As Reported	1.2	g/dL
Beta		As Reported	0.3	g/dL
Gamma		As Reported	0	g/dL
Biochemical Testing - Trace Metals				
Calcium		As Reported	13.5	mg/dL
Chloride		As Reported	97	mmol/L
Iron		As Reported	192	μg/dL
Magnesium		As Reported	3.2	mg/dL
Phosphorus		As Reported	10.2	mg/dL
Potassium		As Reported	>10.0	mmol/L
Sodium		As Reported	135	mmol/L
Biochemical Testing - Hormones				
Cortisol		As Reported	<0.20	μg/dL
Insulin		As Reported	4.39	μIU/mL
Progesterone		As Reported	<0.1	ng/mL
Testosterone		As Reported	<0.01	ng/mL
T3		As Reported	118.0	ng/dL
T4		As Reported	18.5	μg/dL

Notes:

For additional information on this Mediatech product, please contact our Technical Services department at (800) 492-1110 or via email at scientificsupport@corning.com.

This product was tested in accordance with Mediatech, Inc. specifications and procedures current as of date of manufacture. Testing procedures are maintained in compliance with the current versions of the USP and/or EP, where applicable.

Cell lines used in MTT Assay: 3T3, L929, MDCK and Vero. Cell lines used in 9CFR Part 113.53c virus testing: MDBK and Vero.

For research or further manufacturing uses only. Not for use as an excipient. Not for therapeutic or diagnostic uses. Not for human or animal consumption. Utilization of this product apart from the labeled intended use may be a violation of local and/or Federal Law.

Product meets European Union requirements for the production of technical blood products.

This product has been granted a Certificate of Suitability, R0-CEP 2018-271, by the European Directorate for the Quality of Medicines (EDQM) certifying that the product meets the criteria described in the current version of the monograph no. 1483 of the European Pharmacopoeia "Product with risk of transmitting agent of animal spongiform encephalopathies"

Origin and BSE/TSE Statements:

The aforementioned lot was manufactured from fetal bovine blood collected from government approved harvest facilities located in the United States, a country recognized by the United States Department of Agriculture - Animal and Plant Health Inspection Service (USDA-APHIS) and the World Organization for Animal Health (OIE) as having a negligible risk of Bovine Spongiform Encephalopathy (BSE), a bovine-specific Transmissible Spongiform Encephalopathy (TSE). All fetal bovine serum is derived from fetuses collected from healthy bovine cows and heifers that have passed ante and post mortem inspection and were found free of contagious diseases.

Lot Number: 03822002

The following signatures indicate the above material has met all quality specifications and has been reviewed by a Quality representative.

Written By/Date:

DocuSigned by:

Signer Name: La Her
Signing Reason: I am the author of this document
Signing Time: 04-Mar-2022 | 9:43:18 AM EST

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DocuSigned by:

January 04-Mar-2022

Signer Name: Daniel Hu
Signing Reason: I have reviewed this document
Signing Time: 04-Mar-2022 | 12:16:49 PM EST

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Lot Number: 03822002