

Technical Data Sheet

Adult Bovine Serum

Rev: 0

PRODUCT DESCRIPTION

Biowest USA Adult Bovine Serum is a high-quality, aseptically collected serum derived from clotted whole blood of healthy cattle. Sourced from the United States, this serum is meticulously processed and rigorously tested to ensure the highest standards of purity and consistency. With stringent quality control measures, including filtration, endotoxin testing, and virus screening, Biowest USA Adult Bovine Serum offers reliable performance for biological and biomedical studies.

COUNTRY OF ORIGIN

Adult Bovine Serum is sourced from the United States, where it is collected in accordance with stringent animal health and safety regulations.

Catalog Number	Description	Origin(s)
S0230	Adult Bovine Serum	United States

INTENDED USE

Adult Bovine Serum is intended for in vitro applications, including cell culture, research, and further manufacturing purposes. It provides essential nutrients, hormones, and growth factors necessary to support the growth and maintenance of cells. This product is not for human or animal consumption and is not intended for use as an Active Pharmaceutical Ingredient (API). It is strictly for research and laboratory use.

COLLECTION SOURCE

Biowest USA Adult Bovine Serum is derived from clotted whole blood that is aseptically collected. Every batch undergoes rigorous control from serum collection through all stages of treatment and production, ensuring traceability. The serum is processed in accordance with USDA requirements.

QUALITY CONTROL PARAMETERS

- 1.1 pH: monitored and reported for each batch, with specifications provided in the Certificate of Analysis (CoA).
- 1.2 Osmolality: determined by the lowering of the freezing temperature, calibrated against standard solutions. Osmolality specifications are provided in the Certificate of Analysis.
- 1.3 Endotoxin: tested using USP 85 Photometric Quantitative Techniques. Test results are reported in the Certificate of Analysis.
- 1.4 Hemoglobin: measured by spectrophotometry. Results are provided in the Certificate of Analysis.
- 1.5 Total Protein: determined using a colorimetric assay. Results are reported in the Certificate of Analysis.
- 1.6 Sterility Testing: tested for the absence of aerobic and anaerobic bacteria, fungi, and yeast in compliance with 9CFR regulations.
- 1.7 Mycoplasma: tested via culture-based methods or PCR (Polymerase Chain Reaction), depending on customer requirements. Tested for absence of Mycoplasma.
- 1.8 Virus Testing: Testing is performed by inoculating permissive cell cultures and following the 9CFR 113.53c, 113.46, and 113.47 requirements. Each batch undergoes virus testing for the following pathogens:
 - Bovine Viral Diarrhea (BVD)
 - Infectious Bovine Rhinotracheitis (IBR) / BHV-1
 - Parainfluenza Type 3 (PI3)
 - Bluetongue virus
 - Bovine Respiratory Syncytial Virus (BRSV)
 - Reovirus

biowest
DIUWCSL
The Serum Specialist

Technical Data Sheet

	Rabies virus		
1.9	Other Testing: Additional testing may be available upon request.		
	'RATION		
	l Filter Size: 0.2µm		
-	ATMENT PROCESS		
	Applicable		
	RAGE CONDITIONS		
	e at \leq -10°C, protected from light to maintain the serum's integrity.		
	LF LIFE		
	ars from the date of manufacture when stored under the recommended conditions.		
	DLING INSTRUCTIONS		
2.1	Thawing: Thaw the serum in a refrigerator (2°C to 8°C) or at room temperature. Avoid rapid		
	thawing methods to preserve protein integrity.		
2.2	Aliquoting: For optimal preservation, aliquot the serum after thawing using aseptic techniques.		
2.3	Storage after Thawing: If not all serum is used, thawed serum may be stored at 2°C to 8°C for		
	up to 26 weeks without significant loss in quality, provided sterility is maintained.		
2.4	Repeated Freeze/Thaw Cycles: To maintain serum quality, avoid repeated freeze/thaw cycles.		
	Always refreeze aliquots, not the entire bottle.		
PRECAUTIONS AND SAFETY			
3.1	For Research Use Only: Not for human or animal consumption.		
3.2	Protective Equipment: Always wear appropriate PPE, such as gloves, lab coats, and face		
	protection, when handling the serum.		
3.3	Aseptic Handling: Ensure that serum is handled under aseptic conditions (e.g., laminar flow		
	hood) to prevent contamination.		
	ULATORY INFORMATION		
The serum is treated in accordance with strict animal health and safety standards to ensure the			
highest quality and traceability. Biowest USA adheres to all relevant regulatory guidelines and			
	res that the product is safe, traceable, and compliant for research and laboratory applications.		
_	POSAL INSTRUCTIONS		
Dispose of unused serum and packaging in accordance with local regulations for biological materials and hazardous waste.			
	TIFICATE OF ANALYSIS (CoA) rtificate of Analysis is provided with each batch and includes detailed specifications, including		
test results for sterility, endotoxin levels, virus testing, and other quality control parameters.			
DISCLAIMER			
This product is not intended as an Active Pharmaceutical Ingredient (API). It is intended for research,			
	diagnostics, and medical device manufacturing only. Before use, users should refer to the Certificate		
_	of Analysis (CoA) for specific lot details.		
01711			

For further information or inquiries, please contact Biowest USA Customer Service.