

CERTIFICATE OF ANALYSIS

GemCell™ Human Serum AB

Heat Inactivated*

(Catalog Number 100-512-100)

Lot Number: H425007

Date of Manufacture: Jun2021

Product Expiry: Jun2026

Origin: United States

Storage Temperature: ≤ -10°C

For Cell Culture, further Manufacturing or Research use only. Not for direct Therapeutic use.

Product Description: GemCell™ Human Serum AB is collected from healthy male donors of the AB serotype at FDA-licensed facilities located in the United States. Donor units are tested for infectious disease markers prior to processing and found to be non-reactive. GemCell™ Human Serum AB is converted to serum from human plasma using bovine thrombin and sterile-filtered through a 0.1 µm filter prior to freeze.

| Test | Methodology | Specification | Analysis |
|-------------------------|---------------------------|------------------------|---------------|
| Biological Testing | | | |
| Endotoxin | USP<85>, EP 2.6.14 | <10.0 EU/mL | < 0.893 EU/mL |
| Hemoglobin | Fleming, AF and Woolf, AJ | <20.0 mg/dL | 5.8 mg/dL |
| | (1965) | | |
| Microbiological Testing | | | |
| Sterility | USP<71>, EP 2.6.1 | | |
| Bacteria | | No Growth | No Growth |
| Fungi | | No Growth | No Growth |
| Mycoplasma | USP <63> | Not Detected | Not Detected |
| Viral Testing | 21 CFR 610.40 | | |
| HBsAg | ABBOTT ChLIA | Non-Reactive | Non-Reactive |
| Anti-HCV | ABBOTT ChLIA | Non-Reactive | Non-Reactive |
| Anti-HIV-1/HIV-2 | ABBOTT ChLIA | Non-Reactive | Non-Reactive |
| Syphilis | ASI RPR | Negative | Negative |
| HBV-NAT | ROCHE NAT/ PCR | Not Detected | Not Detected |
| HIV-NAT | ROCHE NAT/ PCR | Not Detected | Not Detected |
| HCV-NAT | ROCHE NAT/ PCR | Not Detected | Not Detected |
| Physical Testing | | | |
| Osmolality | USP<785>, EP 2.2.35 | 260 - 350 mOsm/kg | 330 mOsm/kg |
| pH | USP<791> | Test and Report | 7.98 |
| Biochemistry Testing | | • | • |
| Albumin | | Test and Report | 3.4 g/dL |
| ALT (SGPT) | | Test and Report | 14 U/L |
| AST (SGOT) | | Test and Report | 14 U/L |
| Bilirubin, Total | | Test and Report | 0.3 mg/dL |



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| Test | Methodology | Specification | Analysis |
|----------------|-------------|-----------------|-------------|
| BUN | | Test and Report | 13 mg/dL |
| Calcium | | Test and Report | >15.0 mg/dL |
| Chloride | | Test and Report | 117 mmol/L |
| Cholesterol | | Test and Report | 147 mg/dL |
| Creatinine | | Test and Report | 0.83 mg/dL |
| Glucose | | Test and Report | 97 mg/dL |
| Phosphorus | | Test and Report | 3.0 mg/dL |
| Potassium | | Test and Report | 4.0 mmol/L |
| Protein, Total | | Test and Report | 5.3 g/dL |
| Sodium | | Test and Report | >165 mmol/L |
| riglycerides | | Test and Report | 114 mg/dL |
| Uric Acid | | Test and Report | 3.1 mg/dL |

All blood products are collected from stringently screened male donors at FDA-licensed collection centers located in the United States. Viral testing is performed on the individual donor units. All other testing is performed on the final product pool prior to release. The bovine thrombin that is used as part of the conversion to serum is sourced from controlled herds located in the United States and the source cattle are ante and post-mortem inspected by a U.S. Veterinary Service Inspector where they were deemed free of infectious and contagious diseases. All animals used in the production of the thrombin were from a natural beef program in accordance with FDA regulations. Material is derived from human blood and should be considered biohazardous. Universal precautions should be used when handling this material.

The testing that has been performed as part of this lot release has been reviewed by Quality Assurance personnel and has confirmed that the testing meets the specifications presented on this Certificate of Analysis. *Results shown were obtained by carefully performed methods believed to be reliable prior to heat inactivation.

Name Ouline Wilson

QA Specialist

Title

18MAR2025