



Certificate of Manufacture / Analysis

Certificate ID: 10855

Drug Product: Cefazolin 2 Gram/20 mL, Syringe for Injection

Batch/Lot ID: 20211217CEF-1

NDC: 71139-7087-1

Strength: 2 gram Container: 30 mL syringe Volume (Amount): 20 mL

Production Date: 12/17/2021

Expiration Date: 01/31/2022

Storage: Refrigerated Temperature, 2°C - 8°C

Testing Results

Test/Parameter	Specification	Result	Date of Analysis
Cefazolin Potency / ID	90.0 % - 115.0 % / Positive	98.9 % / Positive	12/28/2021
Endotoxin (USP <85>)	≤ 15.0 EU/mL	< 0.50 EU/mL	12/27/2021
Sterility (USP <71>)	Negative Growth (at 14 Days)	Negative	01/06/2022
Visual Inspection	NMT 0 Units Non-Conforming	Pass	12/17/2021
Filter Integrity	Pressure > 55 psi	NA	NA

*Comments: NA

Statement of Conformity:

IntegraDose Compounding Services, LLC certifies that the above product has successfully passed all testing as per the applicable test specifications. All products manufactured under IntegraDose's Quality System meet the regulatory requirements for 503B Outsourcing Facilities (21 USC 353b: Outsourcing facilities and the FDA Current Good Manufacturing Practice – Guidance for Human Drug Compounding Outsourcing Facilities Under Section 503B of the FD&C Act) and all other applicable internal Quality requirements. All products are manufactured with validated processes to current specifications and are inspected to ensure they meet quality requirements.

Certificate Preparation Date: 01/06/2022

By/Date: _____

Quality Review/Date: _____

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