



CLIENT & SAMPLE INFORMATION

Client Mile High Compounds

Analysis Date 04/10/2026

Product Name BPC-157

Strength 10mg

Lot / Batch BPC-010-MH-04

Condition Sterile Injectable

This Certificate of Analysis certifies that the sample listed herein was tested for sterility using membrane filtration methodology in accordance with USP <71> Sterility Tests. The test is designed to detect the presence of viable bacteria and fungi in pharmaceutical products.

TEST METHODOLOGY

Test Performed USP <71>

Compendial Ref USP <71>

Filter Type 0.45 um Membrane

Sample Volume Entire Contents

Incubation Temp (Bacteria) 30-35C

Incubation Temp (Fungi) 20-25C

Incubation Period 14 days

Test Type USP <71>

TEST RESULTS

Parameter	Result
Bacteria (Aerobic)	No Growth
Bacteria (Anaerobic)	No Growth
Fungi / Yeast	No Growth
Acceptance Criteria	No Growth
Final Determination	PASS

CULTURE MEDIA

Medium	Target Organisms	Result
Fluid Thioglycollate Medium (FTM)	Aerobic & Anaerobic Bacteria	No Growth
Soybean Casein Digest Medium (SCDM)	Fungi & Aerobic Bacteria	No Growth

Incubation Duration 14 days

INTERPRETATION

The sample was tested for sterility using membrane filtration in accordance with USP <71>. After the required incubation period, no microbial growth was observed in any of the culture media. The sample meets the acceptance criteria for sterility and is considered to pass the sterility test.

AUTHORIZATION

REVIEWED BY
Lemar Arghandiwal
Lab Director

