DCN: 16-001195 v6.0



100 Majestic Way, Bangor, PA 18013 / www.biospectra.us

Effective Date:	26-Mar-2021		26-Mar-2024	: Date of Next Review
Prepared By:	Jared L Lobb		16-001195 v.5.0	: Supersedes
QA/QC Approval:	Jaron Hughes		Wendy Santay	: Management Approval
Reason for Revision:	See Revision History in ensur.			

CERTIFICATE OF ANALYSIS

POTASSIUM BROMIDE

BIO ACTIVE GRADE / NEW CODE KBRO-2220-25

(HISTORICAL CODE PB2220-K025)

LOT#: KBRO-0123-00004

KBr - F.W. 119.00 g/mol - CAS#: 7758-02-3

Manufacturing Date: 12/1/22 Retest Date: 12/31/24 Manufacturing Site: 100 Majestic Way, Bangor PA, 18013

Packaging Date: 2/5/23 Packaging Site: 100 Majestic Way, Bangor PA, 18013

Meets or exceeds USP Specifications

TES	ST	SPECIFICATION	TEST RESULT
Acidity or Alkalinity	, 1	Passes Test	Passes Test
Appearance of Solution		Clear and Colorless	Clear and Colorless
Assay		98.0 - 100.5%	100.1%
Bromates		Passes Test	Passes Test
Heavy Metals		10 ppm max.	< 10 ppm
Identification	A	Passes Test	Passes Test
	В	Passes Test	Passes Test
Iodides		Passes Test	Passes Test
Limit of Chlorine		0.6% max.	<0.6%
Limit of Iron		20 ppm max.	< 20 ppm
Loss on Drying		1.0% max.	0.1%
Magnesium and Alka	line Earth-Metals	0.02% max.	<0.02%
Sulfates		0.01% max.	<0.01%
	Arsenic (As)	5 ppm max.	< 5 ppm
Trace Metals	Copper (Cu)	5 ppm max.	< 5 ppm
	Iron (Fe)	5 ppm max.	< 5 ppm
	Lead (Pb)	5 ppm max.	< 5 ppm

COUNTRY OF ORIGIN: U.S.A.

TEST METHOD REFERENCE: DCN: 16-001310

DCN: 16-001195 v6.0

CAUTION STATEMENT: For manufacturing, processing, or repacking.

CAUTION STATEMENT: Rx only.

<u>INTENDED USE:</u> Material represented by this Certificate of Analysis is suitable for use as a non-Sterile Active Pharmaceutical Ingredient manufactured in accordance with the ICH Q7 Good Manufacturing Practice Guide. The material represented by this Certificate of Analysis is not suitable to be used as a Sterile or Injectable Active Pharmaceutical Ingredient, Drug Product or Household Item.

STATEMENT: Meets the chemical testing specifications of the current edition of the European Pharmacopoeia.

OVI STATEMENT: Based on the manufacturing process and the controlled handling, storage and analysis of this product, this product complies with the requirements and specifications listed in the current USP method <467> Tables 1, 2, 3, or 4.