DCN: BSI-COA-0097 v.8.1

BISPECTRA

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

Effective Date:	1-Aug-2022	1-Aug-2025	: Date of Next Review
Prepared By:	Wendy Santay	BSI-COA-0097 v.8.0	: Supersedes
QA/QC Approval:	Carissa McCollian	Amy Yencho	: Management Approval
Reason for Revision	See Revision History in MasterControl		

CERTIFICATE OF ANALYSIS

TREHALOSE, DIHYDRATE

BIO EXCIPIENT GRADE / NEW CODE TRED-3250-25

(HISTORICAL CODE TE3250-K025)

LOT: TRED-0124-00004

C₁₂H₂₂O₁₁ · 2H₂O F.W. 378.33 g/mol. CAS# 6138-23-4

Manufacturing Date: 08/18/23 Retest Date: 08/31/25 Manufacturing Site: 100 Majestic Way, Bangor PA, 18013

Packaging Date: 04/05/24 Packaging Site: 100 Majestic Way, Bangor PA, 18013

Meets or Exceeds USP/NF, EP and JP Specifications

		NF COMPENDIA	
Anal	YSIS	SPECIFICATION	TEST RESULT
¹ Assay		³ 98.0% - 101.0%	98.7%
Chloride and Sulfate,	, Chloride	≤ 0.0125%	<0.0125%
Color and Clarity	A720	≤ 0.050	< 0.003
of Solution	A420 - A720	\leq 0.100	0.013
² Endotoxins		$^3 \le 2.4 \text{ EU/g}$	<0.2 EU/g
² Identification A		Conforms to Standard	Conforms to standard
² Identification B		Passes Test	Passes Test
² Identification C		Passes Test	Passes Test
	Escherichia coli	Absent/g	Absent/g
² Microbial	Salmonella species	Absent/10g	Absent/10g
Content	TAMC	≤ 100 CFU/g	<10 CFU/g
	TYMC	$\leq 100 \text{ CFU/g}$	<10 CFU/g
² Nitrogen Determination		$\leq 0.005\%$	<0.005 %
² Optical Rotation, Sp 20°C	pecific Rotation @	+197° to +201°	+200°
²pH @ 25°C		4.5 - 6.5	5.8
Related Substances	Total Impurities with RRT <1.0	≤ 0.5%	0.18%
Related Substances	Total Impurities with RRT >1.0	≤ 0.5%	<0.01%
² Residue on Ignition		≤ 0.1%	<0.1%

DCN: BSI-COA-0097 v.8.1

Analysis	SPECIFICATION	TEST RESULT
² Soluble Starch	Passes Test	Passes Test
Chloride and Sulfate, Sulfate	$\leq 0.0200\%$	<0.0200%
² Water Determination	9.0% to 11.0%	9.8%

		EP COMPENDIA	
	Analysis	SPECIFICATION	TEST RESULT
¹ Assay		³ 98.0 - 101.0%	98.7%
Appearance of S	Solution	Clear, colorless	Clear, colorless
Chlorides		≤ 0.0125%	<0.0125%
² Endotoxins		$^3 \le 2.4 \text{ EU/g}$	<0.2 EU/g
² Identification A	A	Conforms to Standard	Conforms to standard
² Identification I	3	Passes Test	Passes Test
² Identification (C	Passes Test	Passes Test
	Impurity A	≤ 0.5%	<0.10%
¹ Related	Impurity B	≤ 0.5%	<0.10%
Substances	Unspecified Impurities	≤ 0.2%	0.18%
	Total Impurities	≤ 1.0%	0.18%
	Escherichia coli	Absent/g	Absent/g
² Microbial	Salmonella species	Absent/10g	Absent/10g
Content	TAMC	≤ 100 CFU/g	<10 CFU/g
	TYMC	$\leq 100 \text{ CFU/g}$	<10 CFU/g
²pH @ 25°C		4.5 - 6.5	5.8
² Soluble Starch		Passes Test	Passes Test
² Specific Optica	al Rotation @ 20°C	+197° to +201°	+200°
Sulfated Ash		≤ 0.1%	<0.1%
Sulfates		≤ 0.0200%	<0.0200%
² Water		9.0% to 11.0%	9.8%

JP COMPENDIA			
Analysis	SPECIFICATION	TEST RESULT	
¹ Assay	98.0% - 101.0%	98.7%	
Chloride	≤ 0.018%	<0.018%	
² Dextrin, Soluble Starch, Sulfite	Passes Test	Passes Test	
Heavy Metals (as Pb)	≤ 5 ppm	<5 ppm	
² Identification 1	Passes Test	Passes Test	

1	Analysis	SPECIFICATION	Test Result
² Identification 2	2	Passes Test	Passes Test
² Identification 3	3	Conforms to Standard	Conforms to Standard
² Nitrogen		≤ 0.005%	<0.005%
² Optical Rotatio	on @ 20°C	+197° to +201°	+200°
²pH @ 25°C		4.5 - 6.5	5.8
² Residue on Ignition		≤ 0.1%	<0.1%
¹ Related	Total Impurities with RRT < 1.0	≤ 0.5%	0.18%
Substances	Total Impurities with RRT >1.0	≤ 0.5%	<0.01%
Sulfate		≤ 0.024%	<0.024%
² Water		9.0% to 11.0%	9.8%

	Non-Compendial Analyses	
Analysis	SPECIFICATION	TEST RESULT
Appearance and Color	White to Almost White Crystalline Powder	White to Almost White Crystalline Powder
¹ Residual Ethanol	≤ 200 ppm	<95 ppm
¹ Residual Isopropyl Alcohol	≤ 250 ppm	<135 ppm
¹ Residual Methanol	≤ 50 ppm	<25 ppm

¹Alternate Validated Method

COUNTRY OF ORIGIN: U.S.A.

TEST METHOD REFERENCE: DCN: BSI-ATM-0027

<u>INTENDED USE:</u> Material represented by this Certificate of Analysis is suitable for use as an excipient. It is 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 an Active Pharmaceutical Ingredient, Drug Product or Household Item.

<u>RESIDUAL SOLVENTS STATEMENT:</u> 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. Ethanol and Methanol are not used in the manufacturing process.

Prepared by Aud McCall	Date: 4/8/24	Job Title: QA Tech I
Reviewed by: Jan Augh	Date: 4/9/24	Job Title: QA Supervisor

²Analyses are Harmonized

³Specifications is more stringent than Compendia Monograph

•		•	