BIOSPECTRA

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

Effective Date:	12-Dec-2019	12-Dec-2022	: Date of Next Review
Prepared By:	Hannah Bernier	18-002600 v.3.1	: Supersedes
QA/QC Approval:	Carissa McCollian	Dora Meissner	: Management Approval
Reason for Revision:	See Revision History in ensur.		

CERTIFICATE OF ANALYSIS TREHALOSE, DIHYDRATE BIO EXCIPIENT GRADE / TE3250 – G100 LOT: TE3250-009-0320

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

Manufacture Date: 2/9/2019 Retest Date: 2/28/2021

Manufacturing Site: 100 Majestic Way, Bangor PA, 18013

Packaging Date: 3/20/2020

Packaging Site: 100 Majestic Way, Bangor PA, 18013

Trehalose, Dihydrate is currently undergoing a stability shelf life study in accordance with BioSpectra's Stability Program. The proposed retest period is 24 months based on information obtained from development, industry review and raw material supply chain. This retest period may be used for material represented by this CoA unless otherwise

notified by BioSpectra.

ANALY	/SIS	SPECIFICATION	TEST RESULT
Appearance and Color		White to Off-White Crystalline Powder	White to Off-White Crystalline Powder
Appearance of Solution	on (EP)	Clear, Colorless	Clear, Colorless
Assay % w/w		98.0% - 101.0%	99.5 %
	(NF)	$\leq 0.0125\%$	\leq 0.0125 %
Chloride	(EP)	$\leq 0.0125\%$	\leq 0.0125 %
	(JP)	< 0.018%	< 0.018 %
Color and Clarity of Solution	A720	≤ 0.050	0.002 a.u.
	A420 - A720	≤ 0.100	0.016 a.u.
Dextrin, Soluble Starc	h, Sulfite	Passes Test	Passes Test
Endotoxins		\leq 2.4 EU/g	< 1.2 EU/g
Heavy Metals (as Pb)		\leq 5 ppm	\leq 5 ppm
Identification A		Conforms to Standard	Conforms to standard
Identification B		Passes Test	Passes Test
Identification C		Passes Test	Passes Test
Identification 1	(JP)	Passes Test	Passes Test
Identification 2	(JP)	Passes Test	Passes Test
Identification 3	(JP)	Passes Test	Passes Test

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DCN:	18-002600	v.4.0

			DCN: 18-002000 V.4.0	
Analysis		SPECIFICATION	TEST RESULT	
	Maltotriose (Impurity B)	$\leq 0.5\%$	\leq 0.5 %	
Te Impurities	Total Impurities with RRT < 1.0	$\leq 0.5\%$	\leq 0.5 %	
	Total Impurities with RRT > 1.0	$\leq 0.5\%$	≤ 0.5 %	
	Glucose (Impurity A)	$\leq 0.5\%$	\leq 0.5 %	
	Any Other Impurities	$\leq 0.2\%$	\leq 0.2 %	
	Sum of Glucose, Maltotriose, and Other Impurities	$\leq 1.0\%$	\leq 1.0 %	
	Escherichia coli	Absent	Absent	
Microbial	Salmonella species	Absent	Absent	
Content	TAMC	$\leq 100 \text{ CFU/g}$	\leq 10 CFU/g	
	TYMC	$\leq 100 \text{ CFU/g}$	$\leq 10 \text{ CFU/g}$	
Nitrogen Content		$\leq 0.005\%$	≤ 0.005 %	
рН @ 25°С		4.5 - 6.5	5.7	
Residual Ethanol		$\leq 5000 \text{ ppm}$	$\leq 5000 \text{ ppm}$	
Residual Isopropyl Alcohol		$\leq 5000 \text{ ppm}$	$\leq 5000 \text{ ppm}$	
Residual Methanol		\leq 3000 ppm	\leq 3000 ppm	
Residue on Ignition		$\leq 0.1\%$	≤ 0.1 %	
Soluble Starch		Passes Test	Passes Test	
Specific Rotation @ 20°C		+197° to +201°	+198 °	
(NF)		$\leq 0.0200\%$	\leq 0.0200 %	
Sulfate	(EP)	$\leq 0.0200\%$	\leq 0.0200 %	
	(JP)	$\leq 0.024\%$	\leq 0.024 %	
Water (Karl Fischer)		9.0% to 11.0%	9.3 %	

COUNTRY OF ORIGIN: U.S.A.

TEST METHOD REFERENCE: DCN: 18-002375

<u>INTENDED USE:</u> Material represented by this Certificate of Analysis is suitable to be used only as the following: ICH Q7 Compliant cGMP Manufactured Excipient for use in further Manufacturing. The material represented by this Certificate of Analysis is not suitable to be used as an Active Pharmaceutical Ingredient, Drug Product or Household Item.

Prepared by: <u>Nyle Myler / QA Specialist</u> Date: <u>3/23/20</u> Reviewed by: <u>Canada A Supervisor</u> Date: <u>3/23/20</u>