

## **CERTIFICATE OF ANALYSIS**

**VIAL CONTENTS:** Endosafe® Control Standard Endotoxin is prepared from *E. coli* strain 055:B5. Each vial contains 10 ng of purified Lipopolysaccharide, freeze dried in a stabilized matrix.

RSE/CSE RATIO: The potency of this standard in Endotoxin units, (EU) has been determined to be						
EU/ng by the method formerly described in Appendix C ( <i>Gel-clot Technique</i> ) of the GUIDELINE ON VALIDATION OF THE LIMULUS AMEBOCYTE LYSATE TEST AS AN END PRODUCT ENDOTOXIN TEST FOR HUMAN AND ANIMAL PARENTERAL DRUGS, BIOLOGICAL PRODUCTS, AND MEDICAL DEVICES, published by the U.S. Food and Drug Administration.						
CSE Lot:	EX82043	CSE Exp. Date:	Aug 2021	-		
LAL Reagent Lot:	K2813L	LAL Exp. Date:	Apr 2022	-		
RSE Lot:	H0K354	RSE/CSE Ratio:	10	_EU/ng	Vial contents:10	0EU/vial
Geometric Mean Sensitivity with RSE:EU/mL						
IS/CSE RATIO: The Expert Committee on Biological Standardization of WHO has assigned a potency of the IS as 10,000 IU per vial of IS, so that 1 IU = 1 EU. The potency of this endotoxin standard in International (Endotoxin) Units, IU, has been designated as IU/ng.						
DIRECTIONS FOR USE: Reconstitute the lyophilized material with5.0mL of LAL reagent grade water to obtain20EU/mL or20IU/mL. Vortex mix vigorously for at least 5 minutes after rehydration, and for at least 1 minute immediately prior to each use.						
<b>STORAGE:</b> Store rehydrated material at 2-8°C for up to 4 weeks. Store lyophilized material at controlled room temperature or refrigerated as preferred. Diluted endotoxin should not be stored except under validated conditions.						
CAUTION: DO NOT FREEZE ENDOTOXIN SOLUTIONS						
Prepared By:	wed By: (1)	holsh	-		te: <u>30 Oct 2018</u> te: <i>0</i> 2 Nw 2018	
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