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CERTIFICATE OF ANALYSIS

Date:

24 January 2022

Product Name	CLEARKLENS CLEANSINALD SS VH9S		
Product Code	100848254		
Batch Number	FMP22024	53587	
Production Date	24/01/2022		
Expiration Date	EXP 24/07/2023		

This is to certify that the above batch of product has been tested and conforms to the manufacturing Quality Control specification for the product.

	Test Method	Limits		Results	
		Lower	- Upper		
Appearance	Visual	Clear Slightly	Yellow Liquid	Clear Slightly Yellow Liquid	
pH (neat solution)	DM001	9.0	11.4	10.7	
Specific Gravity (20°C)	DM004	0.990	1.010	0.999	

On behalf of Diversey site Quality Manger	Name:	Urszula Haraburda Angelika Partynska
	Position	Quality Control Inspector

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COA Template	Version : 02	Date of issuing: November 2	24th 2017
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STERIS: Gamma Certificate Of Processing

Prepared for: FLEXIBLE MEDICAL PACKAGING LTD (8245)

Gamma Process Run ID 2173-25458A

C/SINALD SS 900ml Bottle DV4673 53624 13 Case

Validation Reference Number: 4673

Processing Run Start Date: 14-Jan-2022 12:56 PM

Processing Run End Date: 14-Jan-2022 6:37 PM

Specified Dose Range (kGy): 25.0 - 40.0 Calculated Min Dose (kGy): 28.5

Reference Dose Range (kGy): 27.3 - 36.6 Calculated Max Dose (kGy): 35.0

PO Number: 401686

Product meets Customer specifications; zero nonconformities occurred during this irradiation run.

Gamma Process Run Approval authorized by STERIS

Date/Time E-Signed: 14-Jan-2022 8:34 PM

Operating facilities are in compliance with applicable regulations providing services under a certified quality system which meets the requirements of FDA QSR, EN/ISO 13485 current certified version, and is in alignment with EN ISO 11137 current certified version

Processing Location

Synergy Health Sterilisation UK Limited, a STERIS Company Brunel Close Drayton Fields Industrial Estate Daventry Northants NN11 8RB

Phone: + 44(0) 1327 706 111

Items Irradiated under WO53624 will be used in finished batch FMP22024 53587.

Olga Kirchner

Document ID: 100044

N/A Last Revised in Rel 2.0.0.0 Rel Date: 13-Aug-2018 Page 1 of 1

PHARMACEUTICAL ANALYSIS REPORT



Flexible Medical Packaging

Unit 8 Hightown White Cross Industrial Estate Lancaster LA14XS

FAO:

Mrs.Olga Kirchner

Report of Tests on:

Cleansinald SS

Sample Description:

Sampe Code: FMP22024 53587

Lucideon Sample Number: UK22415-2738

Lucideon Report Number: UK22415-2738/MFEP

Issue Number:

Date Logged:

28-Jan-2022

Order Number:

PO401809

Date Reported:

17-Feb-2022

Date(s) of Test(s):

01-Feb-2022 to 15-Feb-2022

Sterility Testing Membrane Filtration EP

Test Results:

Result: Pass

The test results meet the EP/USP criteria: Yes

Tests carried out in accordance with cGMP

End of Test Report

JC8001 17-660-22

Mrs Natalie Boot

Senior Business Support Administrator