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CERTIFICATE OF ANALYSIS

Date: 06/03/2019

Product Name	CLEARKLENS CLEANSINALD SC VH9
Product Code	7516429
Batch Number	FMP19065, 46864
Production Date	06.03. 2019
Expiration Date	EXP 06/03/2021

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	Test Method	Limits		Results
		Lower	Upper	
Appearance	Visual	Clear	Slightly Yellow Liquid	clear slightly yellow liquid
pH (neat solution)	DM001	12.0	13.0	12.7
Specific Gravity (20°C)	DM004	1.040	1.060	1.050

On behalf of Diversey site Quality Manager	Name :	J. Staan
	Position	Quality Control Inspector

This document being issued electronically does not bear a signature

COA Template

Version : 02

Date of Issuing : November 24th 2017



Wickham Laboratories

Contract Analytical Services

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Date Received: 12 Mar 2019
Date Tested: 01 Apr 2019
Date Test Completed: 15 Apr 2019
Purchase Order: 32737

CERTIFICATE OF ANALYSIS

Laboratory Reference Number: 0047671/3
Test Required: Sterility by Membrane Filtration Steritest
Date Received: 12/03/2019
Test Article: Cleansinald SC
Sample Code: FMP19065 46864
Batch Ref: 628503
Qty Received: 20 x 25mL Bottles

Test	Method Item	Result
Sterility Test by Membrane Filtration (Steritest) Method	MM107/00	Pass
Growth in Tryptone Soya Broth at 20-25°C after 14 days	MM107/01	No growth in one broth
Growth in Fluid Thioglycollate Medium at 30-35°C after 14 days	MM107/02	No growth in one broth
Volume Tested	MM107/05	20 x 25 mL
Tested in Accordance with Ph Eur, USP & JP	MQ005/07	EP 9.0 2.6.1, USP 41 <71> & JP XVII 4.06
Product Standard Data Sheet	FG047/psd	FM06-01

Approval is provided by Electronic Signature. Their name and position is shown below.

 BSc, CBiol MRSB

Date: 23 Apr 2019 10:02:31

Mrs P. Pham-Lengoc
Documentation Manager – Microbiology

STERIS: Gamma Certificate Of Processing

Prepared for: FLEXIBLE MEDICAL PACKAGING LTD (8245)

Gamma Process Run ID: 2173-1263A

<u>Product Code</u>	<u>Lot Number</u>	<u>Quantity</u>	<u>UOM</u>
C/SINALD BOTTLES/CAPS/BAGS DV4894	WO46875	11	Case

Validation Reference Number: 4894

PO Number: PO32425

Processing Run Start Date: 17-Jan-19 10:56 PM

Processing Run End Date: 18-Jan-19 04:40 AM

Specified Dose Range (kGy):	25.0 - 40.0	Calculated Min Dose (kGy):	29.6
Reference Dose Range (kGy):	27.4- 36.5	Calculated Max Dose (kGy):	35.9

Signature Manifest

Reviewed and E-Signed By: Mark Morse (QA Officer)

Date/Time E-Signed: 2019-01-18 11:42 AM

Document Content Revision: 1

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
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