

Diversey Europe Operations BV Maarssenbroeksedijk 2 3542 DN Utrecht The Netherlands

Tel. +31 (0)30 247 6427 www.Diversey.com

CERTIFICATE OF ANALYSIS

Date: 09 June 2022

Product Name	CLEARKLENS BI-SPORE ACTIVATOR SOLUTION VH26 first off		
Product Code	7515828		
Batch Number	FMP22160 54305		
Production Date	09/06/2022		
Expiration Date	EXP 09/06/2024		

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	Limi Lower	ts - Upper	Results
Appearance	Visual	Clear Colou	ırless Liquid	Clear Colourless Liquid
pH (neat solution)	DM001	9.0	12.5	12.2
Specific Gravity (20°C)	DM004	1.004	1.020	1.005
Potential Chlorine Dioxide		100	180	121.5

On behalf of Diversey site	Name:	Justyna Staron Angelika Partynska
Quality Manger	Position	Quality Control Inspector

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COA Template Version: 02 Date of issuing: November 24th 2017



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CERTIFICATE OF ANALYSIS

Date: 09 June 2022

Product Name	CLEARKLENS BI-SPORE BASE SOLUTION VH26 first off		
Product Code	7515828		
Batch Number	FMP22160 54305		
Production Date	09/06/2022		
Expiration Date	EXP 09/06/2024		

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	Limi Lower	ts - Upper	Results
Appearance	Visual	Clear Colou	rless Liquid	Clear Colourless Liquid
pH (neat solution)	DM001	1.5	2.5	1.9
Specific Gravity (20°C)	DM004	1.010	1.030	1.018

On behalf of Diversey site	Name:	Justyna Staron Angelika Partynska
Quality Manger	Position	Quality Control Inspector

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STERIS: Gamma Certificate Of Processing

Prepared for: FLEXIBLE MEDICAL PACKAGING LTD (8245)

Gamma Process Run ID 2173-27514A

Product CodeLot NumberQuantityUOMBI-SP B3 Components DV5857543738Case

Validation Reference Number: 5857

Specified Dose Range (kGy):25.0 - 40.0Calculated Min Dose (kGy):30.3Reference Dose Range (kGy):26.1 - 35.8Calculated Max Dose (kGy):36.1

PO Number: 402355

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

Gamma Process Run Approval authorized by STERIS

Date/Time E-Signed: 16-Apr-2022 12:11 AM

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 WO54373 will be used in finished batch Fmp22160 54305.

Olga Kirchner

Document ID: 109039

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:

Bi-Spore Activator

Sample Description:

Sample Code: Fmp22160 54305

Lucideon Sample Number: UK222549-16275

Lucideon Report Number: UK222549-16275/MFEP

Issue Number:

Date Logged:

22-Jun-2022

Order Number:

402814

Date Reported:

13-Jul-2022

Date(s) of Test(s):

24-Jun-2022 to 08-Jul-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



Mrs Natalie Boot

Senior Business Support Administrator

PHARMACEUTICAL ANALYSIS REPORT



Flexible Medical Packaging

Unit 8 Hightown White Cross Industrial Estate Lancaster LA14XS

FAO:

Mrs.Olga Kirchner

Report of Tests on:

Bi-Spore Base

Sample Description:

Sample Code: Fmp22160 54305

Lucideon Sample Number: UK222549-16276

Lucideon Report Number: UK222549-16276/MFEP

Issue Number:

1

Date Logged:

22-Jun-2022

Order Number:

402814

Date Reported:

13-Jul-2022

Date(s) of Test(s):

24-Jun-2022 to 08-Jul-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



Mrs Natalie Boot

Senior Business Support Administrator