



## CERTIFICATE OF ANALYSIS

PRODUCT NAME	CLEARKLENS CLEANSINALD SS VH09S
PRODUCT CODE	7516431
BATCH NUMBER	W026232
PRODUCTION DATE	06/2009
EXPIRY DATE	12/2010

TEST	METHOD	LIMITS	RESULTS
Appearance (20°C)	Visual	Clear Solution	clear solution
pH (neat)	GM 0130 01	9.0 – 10.5	10.5
Density (20°C)	GM 0110 01	0.990 – 1.010	1.001

QA Manager:  Date: 25/02/09

QA:  Date: 25-06-09

JD Approval:  Date: 25/06/09

*Date above relates to the date of signing of the document by each responsible person.*

*This document has been produced on behalf of JohnsonDiversey by authorised personnel.*

FLE001

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Flexible Medical Packaging Ltd  
Unit 8  
White Cross Industrial Estate  
Hightown  
Lancaster



**Isotron Limited**

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Facsimile: +44 (0)1274 686061  
Email: assistance@isotron.com  
Website: www.isotron.com

LA1 4XS

Cust. Ref: P014119  
Date Rec'd 28/05/09  
Date: 4/06/09

ITEM CODE ISOTRON BATCH	ITEM SPECIFICATION	QTY	ADDITIONAL DETAILS
B2FLE0010129 B209060046	C/SINALD SS 900ml BOTTLE 25-40kGy	11	B/No.W026227 - W026232  ACTUAL.....kGy DOSE      29.1 - 37.1  CUSTOMERS BATCH/CODE No.s AS ORDER No.P014119 DATED 27/5/09
B2FLE001VAL	VALIDATION		TRIPPLICATE VALIDATION
Total		11	Last Page 1 of 1

All in accordance with current  
technical agreement

This is to certify that the above items have been irradiated as  
specified above

Authorising Signature:

For and on behalf of ISOTRON LIMITED

## CERTIFICATE OF ANALYSIS

Mr J Jackson

 Flexible Medical Packaging Limited  
 Unit 8, Hightown  
 White Cross Industrial Estate  
 Lancaster LA1 4XS

**Date Received** : 26-Jun-09  
**Date Tested** : 30-Jun-09  
**Date Completed** : 14-Jul-09  
**Purchase Order No.** : PO14218

**Test Article** : Cleansinald SS Block 1

**Sample Code** : Works Order No. 26232

**Batch Reference** : 858

**Laboratory Reference** : LR-00867679

**Article Quantity** : 20 Bottles


**Test Required** : Sterility Method I: Membrane Filtration by Steritest. BP, Ph Eur and USP.

Test		Method	Result
Sterility Test by Membrane Filtration (Steritest) Method	*	MM107	Pass
Growth in Tryptone Soya Broth at 22°C after 14 days	*	MM107/01	0/1
Growth in Fluid Thioglycollate at 32°C after 14 days	*	MM107/02	0/1
Product Standard Data Sheet	*	MQ002/01	FM06

Day Book Ref: 574015

Volume Tested: 20 x 50mL

Test carried out according to current Pharmacopoeias.

  
 Mrs S L Wood, BSc., CBiol., MIBiol  
 Laboratory Manager - Pharmaceutical Microbiology

 K.A.Barker, C.Biol, M.I. Biol.,  
 Business Manager - Microbiology

 Date 16 July 2009

<b>GLP</b>
Compliant Laboratory

**Sample Booking Number:** CN\00137466  
 Flexible Medical Packaging Limited

Certificate of Analysis - OSMM

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