

Study Title:

Quantitative suspension test for evaluation of virucidal activity in the medical area (Phase 2 Step1)

Microbiological Solutions Limited (MSL) Gollinrod, Walmersley, Bury, BL9 5NB, UK

Angela Davies, CEO

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Megan Barrett Laboratory Manager Peter Thistlethwaite Technical Projects Manager

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The sample will be retained for 1 month unless otherwise requested in writing.

Bury, BL9 5NB

Tel: 0844 824 6003 Email: info@mls.io Web: www.msl.io Company number: 4218514



BS EN 14476:2013+A2:2019

Scope

The standard method BS EN 14476 describes a test method and the minimum requirements for virucidal activity of a chemical disinfectant and antiseptic products that form a homogenous physically stable preparation when diluted with hard water – or in the case of ready to use products that are not diluted when applied, - with water. Products can only be tested at a concentration of 80% (97% with a modified method for special cases) as some dilution is always produced by adding the test organisms and interfering substances. This European Standard applies to products that are used in the medical area in the fields of hygienic handrub, hygienic handwash, instrument disinfection by immersion, surface disinfection by wiping, spraying, flooding or other means and textile disinfection.

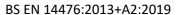
This European standard applies to areas and situations where disinfection is medically indicated. Such indication occurs in patient care, for example: In hospitals, in community medical facilities and in dental institutions or in clinics of schools, of kindergartens and of nursing homes, and may occur in the workplace and in the home. It may also include services such as laundries and kitchens supplying products directly for patients.

Outline of Test Method (Obligatory Test Conditions)

A sample of the test product is diluted in synthetic hard water in products diluted at point of use or water in the case of ready to use products is added to a test suspension of viruses in a solution of interfering substance. The mixture is maintained at one of the temperatures and contact times specified in the standard. At the end of this contact time, an aliquot is taken; the virucidal action in this portion is immediately suppressed by a validated method (dilutions of the sample in ice-cold cell maintenance medium). The dilutions are transferred into cell culture units either using monolayer or cell suspension. Infectivity tests are done either by plaque test or quantal tests. After incubation, the titres of infectivity are calculated according to Spearman and Käber or by plaque counting. Reduction of virus infectivity is calculated from differences of lg virus titres before (virus control) and after treatment with the product. The standard minimum spectrum of test organisms is Poliovirus, Adenovirus and Murine Norovirus.

Acceptance Criteria

The product when tested as above shall demonstrate at least a 4 \log_{10} reduction against the test virus. The test is deemed valid where all control requirements are met.





	Test information	Deviation
Name of Product	*** Protect Professional v2	
Batch Number & Expiry Date	N/S	
Date of Delivery	10/08/2020	
Period of Analysis	17/09/2020-24/09/2020	
Storage Conditions	Ambient	/
Appearance of the Product	Clear Liquid	/
Neutralisation Method	Dilution	
Product Diluent	Distilled water	
Test Concentrations	Neat (80%), Mid-range, Non active	
Experimental Conditions	Clean	
Interfering Substance	Clean 0.3g/l Bovine Albumin	
Test Temperature	20°C ± 1°C	
Temperature of Incubation	37°C ±1°C	
Identification of the Viral Strains:	Feline Coronavirus, Strain Munich	1
Contact Times	60 Minutes <u>+</u> 10s	
Stability and Appearance During Test	: No Change Observed (Homogenous)	

Deviations from Standard Method

1 – The product was tested against non standard organism Feline coronavirus, therefore reference inactivation controls were not performed due to no acceptance criteria available.

Test Result Summary

The test product received has achieved a 4-log reduction against Feline coronavirus when tested under the condition stipulated in this report.

See page 2 for acceptance criteria and raw data tables below for complete test results.



BS EN 14476:2013+A2:2019

Summary

Controls						
	MSL					
Conditions	SOLUTION PROVIDERS	Concentration	Contact time	log TCID50	log reduction	Control validation
Virus control (water)		N/A	60 minutes	6.50	N/A	Validated
Cytotoxicity (product)		Neat	N/A	< 2.25	N/A	Validated
Product supression control		Neat	Neat	6.79	-0.29	Validated

Interference controls	SOLUTION PROVIDERS					
Condition		Concentration	Contact time	log TCID50	Log difference	Control validation
Interference control (untr	reated)	N/A	N/A	7.38	N/A	N/A
Interference control (trea	ted)	Neat	N/A	7.29	0.08	Validated

Test Results	SOLUTION PROVIDERS						
Condition		Concentration	Contact time	log TCID50	log reduction	Pass/Fail	
Test product		Neat	60 minutes	2.50	4.00	Pass	
Test product		50%	60 minutes	2.75	3.75	Fail	
Test product		0.10%	60 minutes	6.38	0.13	Fail	



BS EN 14476:2013+A2:2019

Raw data

Virus cont	rol (water)			Contact ti	me	60 minutes		
Dilution	Counts						% CPE	p(1-p)
-2	4	4	4	4	4	4	1	0
-3	4	4	4	4	4	4	1	0
-4	4	4	4	4	4	4	1	0
-5	4	4	4	4	4	4	1	0
-6	4	4	4	3	3	3	0.875	0.109375
-7	1	1	1	0	0	0	0.125	0.109375
-8	0	0	0	0	0	0	0	0
-9	0	0	0	0	0	0	0	0

Organism	Feline Coro	navirus	
	Strain Mun	ich	
d	1		
sum px	2.00		
n	8		
SD50	-6.50		
SE	0.18		
хр	-5		

Cytotoxicity (product)			Product concentration			Neat		
Dilution	Counts						% CPE	p(1-p)
-2	3	3	3	3	3	3	0.75	0.1875
-3	0	0	0	0	0	0	0	0
-4	0	0	0	0	0	0	0	0
-5	0	0	0	0	0	0	0	0
-6	0	0	0	0	0	0	0	0
-7	0	0	0	0	0	0	0	0
-8	0	0	0	0	0	0	0	0
-9	0	0	0	0	0	0	0	0

Organism	Feline Coronavirus			
	Strain Munich			
d	1			
sum px	1.75			
n	8			
SD50	< -2.25			
SE	0.16			
хр	-1			

Product su	Product supression control			Product concentration			Neat	
Dilution	Counts						% CPE	p(1-p)
-2	4	4	4	4	4	4	1	0
-3	4	4	4	4	4	4	1	0
-4	4	4	4	4	4	4	1	0
-5	4	4	4	4	4	4	1	0
-6	4	4	4	3	3	3	0.875	0.109375
-7	1	1	1	2	2	2	0.375	0.234375
-8	0	1	0	0	0	0	0.04166667	0.039931
-9	0	0	0	0	0	0	0	0

Organism	Feline Coronavirus			
	Strain Munich			
d	1			
sum px	2.29			
n	8			
SD50	-6.79			
SE	0.23			
хр	-5			

Interference control (untreated)				Product concentration			Neat	
Dilution	Counts						% CPE	p(1-p)
-1	4	4	4	4	4	4	1	0
-2	4	4	4	4	4	4	1	0
-3	4	4	4	4	4	4	1	0
-4	4	4	4	4	4	4	1	0
-5	4	4	4	4	4	4	1	0
-6	4	4	4	4	4	4	1	0
-7	4	2	2	3	3	3	0.70833333	0.206597
-8	1	1	2	0	0	0	0.16666667	0.138889
-9	0	0	0	0	0	0	0	0
-10	0	0	0	0	0	0	0	0

Organism	Feline Coronavirus				
	Strain Mun	ich			
d	1				
sum px	1.875				
n	10				
SD50	-7.375				
SE	0.1959				
хр	-6				

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Gollinrod Walmersley Bury, BL9 5NB Tel: 0844 824 6003 Email: info@mls.io Web: www.msl.io Company number: 4218514





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Raw data

Interference control (treated)					oncentratio	Neat		
tion	Counts						% CPE	p(1-p)
-1	. 4	4	4	4	4	4	1	0
-2	. 4	4	4	4	4	4	1	0
-3	4	4	4	4	4	4	1	0
-4	4	4	4	4	4	4	1	0
-5	4	4	4	4	4	4	1	0
-6	4	4	4	4	4	4	1	0
-7	2	2	3	3	3	3	0.66666667	0.222222
-8	1	1	0	1	0	0	0.125	0.109375
-9	0	0	0	0	0	0	0	0
-10	0	0	0	0	0	0	0	0

Organism	Feline Coronavirus
	Strain Munich
d	1
sum px	1.7917
n	10
SD50	-7.292
SE	0.1919
хр	-6

Test product		Product concentration		Neat	Contact time		60 minute:	
Dilution	Counts						% CPE	p(1-p)
-2	4	4	4	4	4	4	1	0
-3	0	0	0	0	0	0	0	0
-4	0	0	0	0	0	0	0	0
-5	0	0	0	0	0	0	0	0
-6	0	0	0	0	0	0	0	0
-7	0	0	0	0	0	0	0	0
-8	0	0	0	0	0	0	0	0
-9	0	0	0	0	0	0	0	0

Organism	Feline Coronavirus
	Strain Munich
d	1
sum px	1.00
n	8
SD50	-2.50
SE	0.00
хр	-2

Test product		Product concentration			50% Contact time			60 minutes
Dilution	Counts						% CPE	p(1-p)
-2	4	4	4	4	4	4	1	0
-3	1	1	1	1	1	1	0.25	0.1875
-4	0	0	0	0	0	0	0	0
-5	0	0	0	0	0	0	0	0
-6	0	0	0	0	0	0	0	0
-7	0	0	0	0	0	0	0	0
-8	0	0	0	0	0	0	0	0
-9	0	0	0	0	0	0	0	0

Organism	Feline Coronavirus
	Strain Munich
d	1
sum px	1.25
n	8
SD50	-2.75
SE	0.16
vn	_2

Test product		Product concentration			0.10%	Contact time	60 minute	
Dilution	Counts						% CPE	p(1-p)
-2	4	4	4	4	4	4	1	0
-3	4	4	4	4	4	4	1	0
-4	4	4	4	4	4	4	1	0
-5	4	4	4	4	4	4	1	0
-6	3	3	3	4	4	2	0.79166667	0.164931
-7	1	0	0	0	0	1	0.08333333	0.076389
-8	0	0	0	0	0	0	0	0
-9	0	0	0	0	0	0	0	0

Organism	Feline Coronavirus
	Strain Munich
d	1
sum px	1.88
n	8
SD50	-6.38
SE	0.19
хр	-5

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KEY

CPE Cytopathic effect

Counts 0-4 indicating degree of cytopathic effect

0 = No effect, 1 = 25% CPE, 2 = 50% CPE, 3 = 75% CPE, 4 = 100% CPE

d Dilution factor (log)

Sum px Sum of % CPE from the highest dilution showing 100% CPE to the lowest dilution assessed.

n Number of dilutions

SD50 Dilution showing 50% of the end point according to Spearman-Kärber method

SE Standard error

xp Lowest dilution showing 100% CPE

TCID50 Titre causing 50% of the end point according to Spearman-Kärber

PASS = lg R greater than or equal to 4

FAIL = lg R less than 4

> greater than ≥ equal to or greater than < less than ≤ equal to or less than

Calculation notes

In cases where the highest dilution assessed has not shown 100% CPE, the value has been calculated assuming the dilution above this would give 100% CPE and the corresponding value has been assigned as <x.

The standard requires the product suppression control to show a <0.5 log reduction in viral titre. In cases where the product has failed to achieve the required 4 log reduction, but the product suppression control shows a >0.5 log reduction the result has been deemed as valid for fail as the consequence of inadequate suppression would be a partially extended contact time which would generate false positives, but not false negatives.

A similar approach has been taken in regards to the cytotoxicity controls. The standard requires a 4-log difference between the cytotoxicity level and the viral titre. In cases where this is not obtained, but the log reduction observed by the product is within the difference between the cytotoxicity levels and the viral titre the result is deemed acceptable for a fail as there will be no impact on the determination of efficacy.