

Test Report: BS EN 14476:2013 + A2:2019 Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of virucidal activity in the medical area- Test method and requirements (Phase 2/Step 1)

BluTest Laboratories Ltd Test Laboratory

5 Robroyston Oval, Nova Business Park, Glasgow, G33 1AP

Identification of sample

Name of the product Alcohol Free Sanitizer

Batch number Not supplied

Client Cleenol Group Limited

Client Address Neville House, Beaumount Road, Banbury, OX16 1RB

Project Code BT-CNL-03FT(2)-02 Date of Delivery 10 February 2020

Storage conditions **Ambient** Active substances Not supplied **Appearance** Liquid

Test Method and its validation

Method 1 part interfering substance + 1 part virus suspension + 8 parts

> biocide were mixed and incubated at the indicated contact temperature for the indicated contact times. Assays were validated by a cytotoxicity control, interference control, neutralisation control and a formaldehyde internal standard.

Neutralisation Dilution-neutralisation/gel filtration

Eagles Minimum Essential Medium + 5.0% v/v foetal bovine serum

at 4°C

Experimental Conditions

Period of analysis 21 February 2020 to 26 February 2020

Product diluents used Sterile distilled water 80.0% v/v; 50.0%; 10.0% v/v Product test concentrations Appearance product dilutions No changes noted-stable Turbidity observed at 10.0% v/v Appearance in test mixture

Contact times (minutes) 2 ± 10s Test temperature 20°C + 1°C

Interfering substances 0.3g/l bovine albumin 37°C + 1°C + 5% CO₂ Temperature of incubation

Identification and passage (P) of virus Vaccinia virus VR-1549 Elstree strain (P10)

Identification and passage (P) of cells Vero Cells (P 30) (Vaccinia Virus)



PROTOCOL SUMMARY

The basic virucidal efficacy test is set up with three concentrations of test product solution and a 2-minute contact time. Virus is exposed to disinfectant in 24-well plates, then neutralised, serially diluted and virus titred in 96-well tissue culture plates to determine the tissue culture infectious $dose_{50}$ (TCID₅₀) of surviving virus. *Vaccinia virus* VR-1549 Elstree strain / Vero cells are assayed in parallel in each test. TCID₅₀ is determined by the method of Karber¹.

Cytotoxicity control

The test product solution is measured for its effects on the host cells used to propagate the virus, to determine the sensitivity of the assay.

Interference control

The effect of the cells after treatment of the test product solution are verified to ensure the cells can show susceptibility for virus infection. This is compared against cells that have not been treated with test product.

Disinfectant suppression control VS1

Virus is added to the highest concentration of test product solution and then the mixture immediately removed and neutralised. The neutralised virus titre is then determined to assess the efficiency of the neutralisation procedure.

Disinfectant suppression control VS2

Internal control which adds virus to neutralised test product solution to assess the efficiency of the neutralisation procedure.

No column Control

Internal control on the highest contact time to assess any impact of the Microspin™ S 400 HR columns.

Virus recovery control

Virus titre is determined for virus in contact with sterile distilled water at t=0, t=2 and at t=15. The virus titre after 2 minutes is then compared to the recovery of disinfectant-treated virus to measure the log reduction in virus titre. The virus titre at 15 minutes is compared to the reference virus inactivation control.

Reference virus inactivation control

Virus is exposed to 0.7% W/V formaldehyde and the recovery of virus determined by TCID₅₀ after 5 and 15 minutes, in order to assess that the test virus has retained reproducible biocide resistance. In addition, the formaldehyde cytotoxicity of neutralised formaldehyde is determined, to measure assay sensitivity.

1Kärber, G.: Beitrag zur Kollektiven Behandlung Pharmakologischer Reihenversuche. Arch. Exp. Path. Pharmak. 162 (1931): 480-487.

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Vaccinia virus (VR-1549) Elstree strain Test Results

EN14476:2013 + A2:2019 Suspension test for the efficacy of Alcohol Free Sanitizer, BT-CNL-03-02 from Cleenol Group Limited against Vaccinia ATCC VR-1549 under Clean conditions

Test Results									
Concentration	10% (v/v)		50%	(v/v)	80% (v/v)				
Exposure Time	data	TCID ₅₀ /ml	data	TCID ₅₀ /ml	data	TCID ₅₀ /ml			
t = 2 min	0.00	3.16E+01	2.00	3.16E+03	2.00	3.16E+03			
Raw Data	000000	3.16E+01	660000	3.16E+03	660000	3.16E+03			
log		1.50		3.50		3.50			
log difference		4.50		2.50		2.50			

EN14476:2013 + A2:2019 Suspension test for the efficacy of Alcohol Free Sanitizer, BT-CNL-03-02 from Cleenol Group Limited

against Vaccinia ATCC VR-1549 under Clean conditions

Summary Table

Product: Interfering substance Concentration Level of cytotoxicity Cotoxicity Cytotoxicity Cytotoxicity Cotoxicity Cytotoxicity Cytotoxic

Summary Table									
Product:	Interfering substance	Concentration	Level of cytotoxicity		>4 lg reduction after 'X' Min				
				0 min	2 min	15 min	30 min	60 min	
Alcohol Free Sanitizer	0.3g/l BSA	80% (v/v)	3.50	3.50	3.50	n.a.	n.a	n.a.	>2 min
		50% (v/v)	3.50	n.a.	3.50	n.a.	n.a	n.a.	>2 min
		10% (v/v)	3.50	n.a.	1.50	n.a.	n.a	n.a.	<2 min
Virus Control	CLEAN			6.00	6.00	6.17	n.a	n.a.	n.a.
							5 min	15 min	
Formaldehyde	PBS	0.7% (w/v)	2.50				4.67	2.50	>60 mins



Vaccinia virus (VR-1549) Elstree strain Control Data

EN14476:20	13 + A2:2019	Suspension te	st for the ef	ficacy of Alco			03-02 from Cl	eenol Group	Limited again	st Vaccinia A	TCC VR-1549
						trols					
Virus Recovery 0 min		Virus Recovery 2 min		Virus Recovery 15 min		Cytotoxicity		Disinfectant Suppression VS		Disinfectant Suppression VS2	
raw data	TCID ₅₀ /ml	raw data	TCID ₅₀ /ml	raw data	TCID ₅₀ /ml	raw data	TCID ₅₀ /ml	raw data	TCID ₅₀ /ml	raw data	TCID ₅₀ /ml
4.50	1.00E+06	4.50	1.00E+06	4.67	1.48E+06	2.00	3.16E+03	2.00	3.16E+03	4.00	3.16E+05
666630	1.00E+06	666630	1.00E+06	666640	1.48E+06	660000	3.16E+03	660000	3.16E+03	666600	3.16E+05
	6.00		6.00		6.17		3.50		3.50		5.50
									2.50		0.50
		Formaldehyde	reference inact	ivation controls					No colum	n Control	
Cytot	oxicity	Exposure time			maldehyde				2 m	ins	
	•		5 min 15 min			min			raw data	TCID ₅₀ /ml	
raw data	TCID ₅₀ /ml		raw data	TCID ₅₀ /ml	raw data	TCID ₅₀ /ml			5.00	3.16E+06	
1.00	3.16E+02		3.17	4.68E+04	1.00	3.16E+02			666660	3.16E+06	
600000	3.16E+02		666100	4.68E+04	600000	3.16E+02				6.50	
	2.50	log		4.67		2.50					
		log difference		1.50		3.67					
									6. 1.6	(TOID)	
Interfere	Interference control		-4	-5	dilution -6	-7	-8		Stock Viru		
			1	1	0.83	0.33	0			E+ 08	
PBS (Control	3.16E+02	3.16E+02	3.16E+02	2.14E+02	6.76E+01	3.16E+01		66666		
	. 50 00		2.50	2.50	2.33	1.83	1.50				
Raw	Data	6	6	6	5	2	0				
	Product		1	1	0.83	0	0				
Pro			3.16E+02	3.16E+02	2.14E+02	3.16E+01	3.16E+01				
		2.50	2.50	2.50	2.33	1.50	1.50				
Raw	Data	6	6	6	5	0	0				
Log Difference		0.00	0.00	0.00	0.00	0.33	0.00				
Product Cyt Dilut	ion	-3	-3	-3	-3	-3	-3				
PBS Dilution		Neat	Neat	Neat	Neat	Neat	Neat				



CONCLUSION

Verification of the methodology

A test is only valid if the following criteria are fulfilled:

- a) The titre of the test suspension of at least 10⁸ TCID50 /ml is sufficiently high to at least enable a titre reduction of 4 lg to verify the method.
- b) Detectable titre reduction is at least 4 log₁₀.
- c) Difference of the logarithmic titre of the virus control minus the logarithmic titre of the test virus in the reference inactivation test is between:
 - Between 0.5 and 2.5 after 30 min and between 2.0 and 4.5 after 60 min for poliovirus
 - Between 3.0 and 5.0 after 30 min and between 3.5 and 5.5 after 60 min for adenovirus
 - Between 1.0 and 3.0 after 30 min and between 2.0 and 4.0 after 60 min for murine norovirus
 - Between 0.0 and 2.0 after 30 min and between 0.5 and 2.5 after 60 min for parvovirus
 - Between 0.75 and 3.5 after 5 min and between 2.0 and 4.0 after 15 min for Vaccinia virus
- d) Cytotoxicity of the product solution does not affect cell morphology and growth or susceptibility for the test virus in the dilutions of the test mixtures which are necessary to demonstrate a 4 log₁₀ reduction of the virus.
- e) The interference control result does not show a difference of $< 1.0 \log_{10}$ of virus titre for test product treated cells in comparison to the non-treated cells.
- e) Neutralisation validation. This is called the disinfectant suppression test in this protocol. The disinfectant was neutralised by column chromatography through an Illustra Microspin S-400 HR column to achieve the best possible neutralisation available for this test. The difference for virus is greater than 0.5 log₁₀ indicating rapid irreversible virucidal activity of the disinfectant by dilution at a concentration of 80.0% v/v for VS1. This neutralisation validation has been verified by VS2, which shows the product has been successfully neutralised.

According to EN 14476:2013 + A2:2019, **Alcohol Free Sanitizer POSSESSES VIRUCIDAL** activity at a concentration of **10.0% v/v** of the working concentration as tested after **2 MINUTES** at **20°C** under **CLEAN** conditions (0.3 g/l bovine albumin) against *Vaccinia virus* VR-1549 Elstree strain / Vero cells.

The cytotoxicity of the product has prevented at 4.0 log reduction being observed at 80.0% v/v and 50.0% v/v.

This product therefore is effective against all enveloped viruses as defined in EN 14476:2013 + A2:2019. This therefore includes all coronaviruses and SARS-CoV-2.

Signed

Dr Chris Woodall, Director BluTest Laboratories Ltd Glasgow, UK

Date: 27 February 2020.

DISCLAIMER

The results in this test report only pertain to the sample supplied.

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