



STUDY REPORT

Study Title

ASTM E1052

Standard Test Method to Assess the Activity of Microbicides against Viruses in Suspension

Product Identity

Quadsil Hand Gel

Test Microorganism

Human Coronavirus, Strain 229E, ATCC VR-740

Study Identification Number

NG15327

Author

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Study Completion Date

13JUL2020

Testing Facility

Microchem Laboratory
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Study Sponsor

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STUDY REPORT SUMMARY

General Study Information

Study Title: ASTM E1052
Standard Test Method to Assess the Activity of
Microbicides against Viruses in Suspension

Study Identification Number: NG15327

Test System

Test Microorganism: Human Coronavirus, Strain 229E, ATCC VR-740

Host Cell: MRC-5 (CCL-171)

Test Substance: Quadsil Hand Gel

Test Substance Receipt Date: 05MAY2020

Test Parameters

Test Substance Dilution: Ready to use

Total Organic Soil Load: No additional soil load incorporated into the
inoculum

Number of Replicates Per Lot: Single

Contact Time(s): 60 seconds and 5 minutes

Exposure Temperature: Room temperature

Neutralization Method(s): 2% FBS EMEM

Study Dates

Experimental Start Date/Time: 23JUN2020 / 1214

Experimental Termination Date/Time: 30JUN2020 / 1423

Study Completion Date: 13JUL2020



TEST PROCEDURE

Summary

- Stock virus was thawed and was not supplemented with an organic soil load.
- Test and virus control substances were dispensed in 9-part equivalent volumes into sterile vessels.
- Test and virus control substances were each inoculated with 1-part equivalent volumes of the test virus.
- The test suspensions were held for the contact time(s) specified by the Study Sponsor, and then neutralized by ten-fold serial dilutions into the appropriate solution.
- The virus control suspension was neutralized in the same manner as the test suspensions.
- Following neutralization, the viral suspensions were quantified to determine the levels of infectious virus using standard cell culture (e.g. TCID₅₀) assay techniques.
- The cell culture plates were incubated for the period most suitable for the virus-host cell system (e.g. 7 days).
- After the incubation period, the assay was scored for the presence/absence of test virus and cytotoxic effects. The appropriate calculations were performed (e.g. Spearman-Kärber) to determine viral titers and levels of test substance cytotoxicity, where applicable.
- Log₁₀ and percent reductions were computed for test suspensions relative to the control suspensions, and reported to the Study Sponsor.
- Unless otherwise noted, no modifications to the method were made for this study.

SUCCESS CRITERIA

The following measures are met to ensure the acceptability of virucidal efficacy data:

- The virus titer control demonstrate obvious and or typical cytopathic effects on the monolayers unless a detection method other than cytopathic effect is used.
- Neutralization of the test substance with a low titer (e.g. 1000-5000 infective units) of the test virus is demonstrated.
- Quantification of the test and control parameters are conducted at a minimum of four determinations per dilution.

The product performance criteria follows:

- The log and percent reduction of the test virus following exposure to the test substance are calculated however, there is no minimum reduction level to qualify as "passing" or an "efficacious" product.



CALCULATIONS AND STATISTICAL ANALYSIS

The TCID₅₀ (Tissue Culture Infectivity Dose) represents the endpoint dilution where 50% of the cell cultures exhibit cytopathic effects due to infection by the test virus. The endpoint dilution at which 50% of the host cell monolayers exhibit cytotoxicity is termed the Tissue Culture Dose (TCD₅₀). The TCID₅₀, and TCD₅₀ was determined using the Spearman-Kärber method and calculated as follows:

Negative logarithm of endpoint titer =

$[-\text{Log of first dilution inoculated}] - [(\text{sum of \% mortality at each dilution}/100) - 0.5] \times \text{Logarithm of dilution}$

The result of this calculation is expressed as TCID₅₀/0.1 ml (or volume of dilution inoculated) for the test, virus control, and neutralization control and TCD₅₀/0.1 ml (or volume of dilution inoculated) for the cytotoxicity control.

Calculation of the Log Reduction

The log reduction in viral titer was calculated as follows:

Plate Recovery Control Log₁₀ TCID₅₀ – Virus-Test Substance Log₁₀ TCID₅₀

Calculation of the Percent Reduction

The percent reduction in viral titer was calculated as follows:

Percent Reduction = $1 - (C/B) \times 100$, where:

B = Average TCID₅₀ of virus in control suspensions.

C = Average TCID₅₀ of virus in virus-test suspensions.

The presence of any test substance cytotoxicity were taken into account when calculating the log and percent reductions in viral titer.

If multiple virus control and test replicates were performed, the average TCID₅₀ of each parameter was calculated and the average result used to calculate the log reductions in viral titer.



RESULTS

Table 1: Virus Titer and Virus Control

		Virus Titer	Virus Control
Cell Control		0 0 0 0	0 0 0 0
Dilution	10 ⁻¹	+ + + +	N/A
	10 ⁻²	+ + + +	+ + + +
	10 ⁻³	+ + + +	+ + + +
	10 ⁻⁴	+ + + +	+ + + +
	10 ⁻⁵	+ + + 0	+ + + +
	10 ⁻⁶	+ + 0 0	+ 0 0 0
	TCID ₅₀ per 0.1 ml		5.75 Log ₁₀

Key: + = Virus recovered; 0 = Virus not recovered and/or no cytotoxicity observed;
T = Cytotoxicity observed

Table 2: Test Results — Quadsil Hand Gel

		Contact Time	
		60 seconds	5 minutes
Cell Control		0 0 0 0	0 0 0 0
Dilution	10 ⁻¹	N/A	N/A
	10 ⁻²	T T T T	T T T T
	10 ⁻³	0 0 0 0	0 0 0 0
	10 ⁻⁴	0 0 0 0	0 0 0 0
	10 ⁻⁵	0 0 0 0	0 0 0 0
	10 ⁻⁶	0 0 0 0	0 0 0 0
	TCID ₅₀ per 0.1 ml		≤2.50 Log ₁₀
Log ₁₀ Reduction		3.25 Log ₁₀	3.25 Log ₁₀
Percent Reduction		99.68%	99.68%

Key: + = Virus recovered; 0 = Virus not recovered and/or no cytotoxicity observed;
T = Cytotoxicity observed



RESULTS (cont.)

Table 3: Cytotoxicity Control Results

		Cytotoxicity Control
Cell Control		0 0 0 0
Dilution	10 ⁻¹	N/A
	10 ⁻²	T T T T
	10 ⁻³	0 0 0 0
	10 ⁻⁴	0 0 0 0
TCID ₅₀ per 0.1 ml		≤2.50 Log ₁₀

Key: + = Virus recovered; 0 = Virus not recovered and/or no cytotoxicity observed;
 T = Cytotoxicity observed

Table 4: Test Substance Neutralization Control Results

		Neutralization Control
Cell Control		0 0 0 0
Dilution	10 ⁻¹	N/A
	10 ⁻²	T T T T
	10 ⁻³	+ + + +
	10 ⁻⁴	+ + + +
Neutralized at TCID ₅₀ per 0.1 ml		≤2.50 Log ₁₀

Key: + = Virus recovered; 0 = Virus not recovered and/or no cytotoxicity observed;
 T = Cytotoxicity observed



STUDY CONCLUSION

The purpose of the study was to determine the virucidal efficacy of Quadsil Hand Gel against Human Coronavirus, Strain 229E, with no additional soil load incorporated into the inoculum, at contact times of 60 seconds and 5 minutes and at an exposure temperature of room temperature.

The Virus Control demonstrated an average viral titer of 5.75 Log₁₀ TCID₅₀ per 0.1 ml.

The Test Substance Neutralization Control demonstrated that the test substance was neutralized at ≤2.50 Log₁₀ for the single lot assayed.

Test substance cytotoxic effects to the host monolayer were observed at ≤2.50 Log₁₀ TCD₅₀ per 0.100 ml.

Taking the cytotoxicity and neutralization control results into consideration, the evaluated test substance, Quadsil Hand Gel demonstrated a 3.25 Log₁₀ reduction in viral titer (99.68%) at contact times of 60 seconds and 5 minutes.

The test substance will be disposed of 30 days after the completion of this study, unless otherwise requested by the Study Sponsor.

The results of this study apply to the tested substances(s) only. Extrapolation of findings to related materials is the responsibility of the Sponsor.

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