

SARS-CoV-2 Inactivation Testing: Interim Report

Report identifier	HCM/CoV2/059/v1		
Report date	10 November 2020		
Undertaken by High Containment Microbiology, NIS Laboratories, National Infection			
Service, Public Health England			
N.B. This is an interim report and may be updated as further results are obtained			

Product/treatment details		
Product/treatment	virusPHIX-P9	
Manufacturer	Rapid Labs/RNAssist	
Product code	RD-VP9-50	
Manufacturer's recommended ratio of sample to product	Swab samples: swab to be added directly to tube containing 1ml product. Saliva samples: 1 volume saliva to be added to 3 volumes product	

Sample details		
Sample type tested	Tissue culture fluid containing 5% (v/v) foetal calf	
1 31	serum	
Virus strain tested	SARS-CoV-2 England 2	
Ratio of spiked virus stock to	Not applicable; tissue culture fluid used undiluted	
sample matrix	,	

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Experimental conditions		
Ratio of sample to product tested	1 volume sample to 3 volumes product	
Contact time/s	10 minutes; 30 minutes	
Temperature of incubation	Room temperature	
Brief description of tests performed	Triplicate samples were treated with test buffer for indicated contact time/s or mock-treated in triplicate with an equivalent volume of PBS. All samples were then diluted two-fold to reduce product viscosity and subjected to a purification step to remove cytotoxic buffer components. PBS-treated samples were subjected to the same dilution and purification procedure in parallel. Purified samples were titrated on Vero E6 cells to establish virus titre. This test is quantitative and reports the titre of virus in each treatment condition in TCID50 per ml. Reduction in virus titre following treatment is given as the difference between the mean log ₁₀ TCID50/ml for treated conditions and the PBS control.	

Table of results		
(0)	Mean virus titre in	Titre reduction in
	log ₁₀ TCID50/ml	log ₁₀ TCID50/ml
	[95% confidence interval]	[95% confidence interval]
PBS-treated	5.8 [5.5-6.1]	-
Test buffer-treated (10 minutes)	≤1.4 [†] *	≥4.4 [4.1-4.6]
Test buffer-treated (30 minutes)	≤1.0 ^{†*}	≥4.8 [4.5-5.1]

[†]Limit of detection for test was 1.0 log₁₀ TCID50/ml. Mean titres are reported as ≤ when at least one replicate was below the limit of detection.

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^{*95%} confidence interval cannot be calculated

Interpretation

Treatment with virusPHIX-P9 reduced SARS-CoV-2 titre by ≥4.4 and ≥4.8 log₁₀ TCID50/ml after 10 and 30 minutes, respectively. 30 minute treatment reduced levels of virus to below the limit of detection of the test.

Demonstrating complete inactivation is dependent on the starting titre of virus used for testing. Complete inactivation is likely if samples contained lower levels of infectious virus than those tested here, but sample treatments that inactivate virus effectively in our testing may fail to inactivate samples containing higher levels of virus than those evaluated in this study.

This test has been performed using tissue culture fluid. The effectiveness of this treatment against SARS-CoV-2 may vary when used to inactivate clinical samples or other types of sample matrix. Any results of inactivation testing using other sample matrices will be released as they become available.

Inactivation reagents should not be assumed to be 100% effective against SARS-CoV-2.

Suitability of products and treatments for inactivation of other pathogens has not been evaluated in this study.

All COVID-19 laboratory testing workflows must be subjected to suitable and sufficient risk assessment, with consideration given to any inactivation step. Risk assessments should be reviewed regularly as new information on the inactivation of SARS-CoV-2 becomes available.

The impact of chosen inactivation method on the sensitivity of subsequent SARS-CoV-2 detection should also be assessed locally.

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Summary of revisions

Version 1: New document

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Queries regarding this report or HCM inactivation testing should be directed to HCMgroup@phe.gov.uk

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