



Monkeypox Virus Inactivation Testing Report

Report identifier	HCM/MPx/011/v1
Report date	5 October 2022
Testing laboratory	High Containment Microbiology, UK Health Security Agency (UKHSA)

Product details	
Product name	NeuMoDx Viral Lysis Buffer
Product code	401600
Lot number	118384
Manufacturer	NeuMoDx
Storage conditions	Ambient temperature
Active substances and concentrations (if known)	Guanidine hydrochloride (33.5%)
Instructions for use	1 volume sample treated with 1 volume of product

Report identifier and version number: HCM/MPx/011/v1

Report date: 5 October 2022

Page 1 of 4

UNCONTROLLED WHEN PRINTED

Experimental conditions	
Period of analysis	23 September – 05 October 2022
Product test concentrations	1 volume product to 1 volume test sample
Test temperature	Ambient temperature
Treatment times tested	10 minutes
Sample type tested and virus details	Monkeypox virus stock: Monkeypox virus isolate UK2 (GenBank entry MT903344), in tissue culture fluid containing 5% foetal bovine serum
Description of test	<p>Triplicate samples of monkeypox virus tissue culture fluid were treated with product at the indicated test concentration for indicated contact times. Mock-treatments were carried out in triplicate using an equivalent volume of minimum essential medium (MEM) instead of product. After treatment, all samples were subjected to a filtration step to reduce cytotoxic buffer components, using Pierce Detergent Removal Spin Columns in accordance with the manufacturer's instructions. MEM-treated samples were subjected to the same filtration procedure in parallel. All samples were immediately titrated on Vero E6 cells and plates immunostained using an anti-vaccinia virus antibody to establish virus titre. Product only controls (purified and unpurified) were additionally titrated to determine product cytotoxicity before and after filtration.</p> <p>This test is quantitative and reports the virus titre for each treatment condition in focus forming units (FFU)/mL. Reduction in virus titre following treatment is given as the difference between the mean log₁₀ FFU/mL for treated conditions and the MEM control.</p>

Table of results				
Treatment condition	Mean virus titre in FFU/mL	Mean virus titre in log₁₀ FFU/mL [95% CI]	Titre reduction in log₁₀ FFU/mL [95% CI]	% reduction in virus titre
MEM-treated	6.8x10 ⁶	6.8 [6.5-7.2]	-	-
10-minute treatment	5.2x10 ⁵	5.7 [5.5-6.0]	1.1 [0.8-1.5]	92.338%

Results interpretation and limitations

Treatment with NeuMoDx Viral Lysis Buffer (1 volume product to 1 volume sample) for 10 minutes resulted in a 1.1 log₁₀ reduction in monkeypox virus titre in these tests. This is a modest reduction compared to the inactivation effectiveness of other molecular lysis buffers, and a considerable level of infectious virus remaining following treatment with this product. This product should not be relied upon to completely inactivate monkeypox virus.

This test has been performed using tissue culture fluid. The effectiveness of this treatment against Monkeypox Virus may vary when used to inactivate clinical samples or other types of sample matrix.

Nucleic acid stability in this product has not been examined, nor has the suitability of this product for inactivation of other pathogens been evaluated in this study.

Disclaimer

UKHSA does not in any way recommend any particular product for virus inactivation; and UKHSA shall not be responsible for the choice of product or treatment for virus inactivation, and it is the responsibility of users of the product to ensure that any such product or treatment implemented has undergone the necessary verification and validation; and UKHSA shall not be liable, to the greatest extent possible under any applicable law, for any claim, loss or damage arising out of or connected with use of this and related reports and choice of virus inactivation products or treatments.

UKHSA is an executive agency of the Department of Health and Social Care.
Unauthorised use of the UKHSA name and/or logo is prohibited.

Summary of revisions

Version 1: New document