

EVS-EN 17111:2018 INTERFLO OÜ LABORATORY Maagi 6a, 74114, Maardu Estonia Tel.+372 58098890

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Quantitative carrier test for the evaluation of virucidal activity in the medical area (phase 2, step 2)

Test report no 786

1. General information

Client: Medisept Sp. z o.o., ul. Ludwika Spiessa 4, 20 270 Lublin, Poland

Date of order: 2023/04/03

2. Identification of sample

Name of the product: VIRUTON PULVER

Batch number: 220902_4

Aplication Solid product for disinfection of medical instruments





Manufacturer: Medisept Sp. z o.o

Date of delivery: 2022/11/21

Storage conditions: room temperature and darkness

Apperance of the product: Dry blend of white and blue granules and powder

Active substance and concentration 44 % Sodium percarbonate, 26 % TAED

3. Test conditions

Test period: 2023/05/19 - 2023/05/30

End of testing 2023/05/30

Product test concentrations: 0,05%, 0,5%, 1,0%, 2,0%

Exposure time: 15 min.

Test temperature: $19.5^{\circ}\text{C} \pm 0.5^{\circ}\text{C}$

Interfering substance: for dirty conditions (bovine albumine 3,0 g/l and sheep erythrocytes 3 ml/l)

No flocculation, no precipitation

Neutralization of active substance: Immediate dilution, 30 min, +4°C

Diluent PBS, EMEM + 2% FBS

Stability of product in the mix with virus and interfering substance

virus and interfering substance

Test organisms: Adenovirus type 5 ATCC VR-5
Murine norovirus strain S99 Berlin

4. Materials

1.1 Culture medium and reagents

- Eagle's Minimum Essential Medium (EMEM, Corning, REF 10-009-CVR)
- Dulbecco's modification of Eagle's Medium (DMEM, Corning, REF 10-014-CVR)
- Fetal Bovine Serum (FBS, Corning, REF 35-016-CV)
- 50% % Glutaraldehyde solution (Thermo Scientific, A10500)
- Aqua bi-distillate (Interflo OU, Estonia)



- PBS (Phosphate Buffered Saline) (Interflo OU, Estonia)
- BSA (Bovine Serum Albumin fr.V) (Roche Diagnostic GMBH)
- ShE (Defibrinated sheep erythrocytes) (BioTRADING Benelux B.V.)

1.2 Virus and cells

Strain: Adenovirus type 5 ATCC VR-5, LOT 70024114 (LGC Standards GmbH, Wesel, Germany)

HeLa cells ATCC-CCL-2, LOT 70037076. Morphology – epithelial-like (LGC Standards GmbH, Wesel, Germany)

Murine Norovirus S99, RVB-0651 (Friedrich-Loeffler-Institute, Greifswald, Germany)

Cells: RAW 264.7 CCLV-RIE 0996 cells, (ATCC- TIB-71), Home mouse (*Mus musculus*). Morphology – monocyte/macrophage (Friedrich-Loeffler-Institute, Greifswald, Germany)

- 1.3 Apparatus, glassware and small items of equipment
- CO2 incubator Memmert GmbH & Co. KG, ICO50 med, no D218.1233, Germany
- Agitator Vortex V-3, no 7EE0011 Biosan SIA, Latvia
- Digital pH-meter, Instrument: HI2002, SN:CO646088, Hanna, USA
- Centrifuge Centric 322A, Code: 464.000, Domel, d.o.o. BU Laboratory system, Slovenia
- Inverted microscope Motic AE2000 Series, Motic China Group, Ltd, China
- Water bath WB/OB 7-45 WBU 45 Memmert GmbH & Co. KG, Germany
- Adjustable and fixed-volume pipettes, Sartorius Lab Instruments GmbH & Co. KG Germany
- Tissue Culture Plate, 96-well, Falcon, REF 353072 Corning Incorporated, USA
- Tissue Culture flask Falcon pour culture cellulaire, REF 353108 Corning Incorporated, USA
- Tissue Culture flask Falcon pour culture cellulaire, REF 353136 Corning Incorporated, USA
- 50 mL Polypropylene Conical Tube Falcon a Corning Brand, REF 352070, Corning Science Mexico S.A. de C.V., Mexico
- Frosted Glass Carriers 15mm x60mm x1mm, one surface sandblasted, Interflo OU, Estonia



5. Methods

5.1 Preparation of test virus suspension

For the preparation of the test virus suspension according to EN 5.4.1 cells which were cultivated with Eagle's Minimum Essential Medium (EMEM) supplemented with L-glutamine, sodium pyruvate and 10 % fetal calf serum (FCS), were infected with a multiplicity of infection of 0,1 at 37°C. After cells showed a cytopathic effect, they were subjected to a threefold freeze/thaw procedure followed by a low speed centrifugation in order to sediment cell debris. After aliquotation of the supernatant, test virus suspension was stored at –80°C.

5.2 Preparation of disinfectant (dilutions)

The test product was evaluated as 100 % -+solution. Due to the addition of the fresh prepared hard water (5.2.2.7) heated to $35 \degree C \pm 5.0 \degree C$ and mix during the 20 min. (5 min. for dilution and 15 min for product activation) were prepared solutions for test. The test product was evaluated as 2.0 %, 1.0%, 0.5% solution and 0.05% solutions (demonstrations of non –active range).

5.3 Inoculation of the carriers

Glass carriers were placed in Petri dishes in horizontal position. Pipette 0,05ml of the mixture of test suspension and interfering substance on the inoculation square of the carrier. Inoculum were dried under an air laminar flow at 20°C in 15 min.

5.4 Infectivity assay

Infectivity was determined as endpoint titration according to EN 5.5 transferring 0.1 ml of each dilution into eight wells of a microtiter plate to 0.1 ml of freshly trypsinised cells (10 3-10 4 cells per well), beginning with the highest dilution. Microtitre plates were incubated at 37° C in a 5 % CO₂ – atmosphere. The cytopathic effect was read by using an inverted microscope after seven days. Calculation of the infective dose TCID₅₀ /ml was calculated with the method of Spearman (2) and Kärber 3) with the following formula:

$$-\log$$
10 TCID 50 = Xo $-0.5 + \sum r/n$





meaning

Xo = log 10 of the lowest dilution with 100 % positive reaction

r = number of pos. determinations of lowest dilution step with 100 % positive and all higher positive dilution steps

n = number of determinations for each dilution step.

5.5 Calculation and verification of virucidal activity

The virucidal activity of the test disinfectant was evaluated by calculating the decrease in titer in comparison with the control titration without disinfectant. The difference is given as reduction factor (RF).

According to the EN 17111, a disinfectant or a disinfectant solution at a particular concentration is having virus-inactivating efficacy if the titre is reduced at least by $4 \log 10$ steps within the recommended exposure period. This corresponds to an inactivation of $\geq 99,99$ %.

5.6 Inactivation assay

Determination of virucidal activity has been carried out in accordance to EN 5.5. The test product was examined as 2,0 % 1,0%, 0,5% solutions and as 0,05% solution (demonstrations of non –active range) in water at 20°C according to EN 17111, 15 min. were chosen as contact times.

Immediately at the end of the chosen contact time, activity of the disinfectant was stopped by dilution to 10 - 8 at +4°C.

Titrations of the virus control were performed after the longest exposure time (EN 5.5.6).

Furthermore, a cell control (only addition of medium) was incorporated.

Inactivation tests were carried out in sealed test tubes in a water bath at $20^{\circ}\text{C} \pm 1.0^{\circ}\text{C}$. Aliquots were retained after appropriate exposure times, and residual infectivity was determined.

5.7 Determination of cytotoxicity

Determination of cytotoxicity was performed according to EN 5.5.3.1.



5.8 Cell sensitivity to virus

For the control of cell sensitivity to virus 0.3 parts by volume of water were mixed with 9.7 parts by volume of the lowest apparently non-cytotoxic dilution of the product. This mixture or PBS as control was added to a volume of double concentrated cell suspension. After 1 h at 37°C the cells were centrifuged and re-suspended in cell culture medium (EN 5.5.3.2b).

Finally, a comparative titration of the test virus suspension was performed on the pre-treated (disinfectant) and non-pre-treated (PBS) cells as described above.

5.9 Control of efficacy for suppression of disinfectant's activity

Furthermore, a control of efficiency for suppression of disinfectant's activity was performed according (EN 5.5.4).

5.10 Reference virus inactivation test

As reference for the test validation 50 % glutaraldehyde solution according to EN 5.5.5 was included. 5 min was chosen as contact time. In addition, cytotoxicity of glutaraldehyde test solution was determined following EN 5.5.3.1.

6. Verification of the methodology

The following criteria as mentioned in EN 5.7 were fulfilled:

- a) The titre of the test virus suspension allowed the determination of a $\geq 4 \log_{10}$ reduction (maximal virus reduction $\geq 6,00\pm0,00$).
- b) The difference of the logarithmic titre of the virus water control minus the logarithmic titre of the test virus in the reference inactivation test (see 5.5.5) was ≥4,00 log10 after 5 min for both viruses.
- c) The test product (2,0 %) showed cytotoxicity in the 1:100 dilutions thus allowing the detection of a 4 log 10 reduction of virus titre
- d) The comparative titration on pre-treated (disinfectant) and non-pre-treated (PBS) BGM cells showed no significant difference <1 log 10 of virus titre: 8,75 (PBS) versus 8,50 (1:1000 dilutions of 2.0% of disinfectant) log 10 TCID50/ml for Adenovirus and 8,75 (PBS) versus 8,25 (1:1000 dilutions of 2.0% of disinfectant) log 10 TCID50/ml for Norovirus.





e) The control of efficacy for suppression of disinfectant's activity (1,0%) showed no decrease in virus titer (9,00 versus 8,50 log10 TCDI50/ml for Adenovirus and 9,00 versus 8,63 log10 TCDI50/ml for Norovirus).

Since all criteria according EN 5.7 fulfilled, examination according to EN 17111 is valid.

7. Results

Results for examination are shown in tables 1 to 7. Tables 1 to 6 demonstrate the raw data, whereas table 7 gives a summary of results.

The diluted test product (2,0%, 1,0% assay) was able to inactivate Adenovirus and Norovirus after 15 min. in this quantitative carrier test (Table 1,2). The reduction factor was $\geq 6,00\pm0,00$ at this time point. This corresponded to in inactivation of ≥ 99.99 %. The test product in 0,5 % assay was also able to inactivate Adenovirus and Norovirus after 15 min. in this quantitative suspension test (Table 3). The reduction factor was $4,88\pm0,74$ for Adenovirus and $5,50\pm1,00$ for Norovirus at this time point.

8. Conclusion

The disinfectant for medical instruments VIRUTON PULVER tested with concentrations 2,0%, 1,0%, 0,5 % demonstrated effectiveness against Adenovirus and Norovirus after an exposure time of 15 min. under dirty conditions.

Therefore, the disinfectant VIRUTON PULVER (LOT 220902_4) can be declared as active against viruses for instrumental disinfection with treatment temperature \leq 40°C.

0,5% 15 MIN

Tallinn, 2023/05/30

Dr. Ljudmila Šljapnikova



9. Reference

1. EN 17111:2018: Chemical disinfectants and antiseptics – Quantitative carrier test for the evaluation of virucidal activity for instruments used in the medical area – Test method and requirements (phase 2, step 2)

Annex:

Legend to the tables

- Table 1: Raw data for VIRUTON PULVER (2,0%) tested against Adenovirus
- Table 2: Raw data for VIRUTON PULVER (1,0%) tested against Adenovirus
- Table 3: Raw data for VIRUTON PULVER (0,5%) tested against Adenovirus
- Table 4: Raw data for VIRUTON PULVER (0,05%) tested against Adenovirus
- Table 5: Raw data for VIRUTON PULVER (2,0%) tested against Norovirus
- Table 6: Raw data for VIRUTON PULVER (1,0%) tested against Norovirus
- Table 7: Raw data for VIRUTON PULVER (0,5%) tested against Norovirus
- Table 8: Raw data for VIRUTON PULVER (0,05%) tested against Norovirus
- Table 9: Raw data for GLUTARLDEHYDE solution (50 %) tested against Adenovirus, Norovirus
- Table 10: Raw data for control of efficacy for suppression of disinfectant's activity (1,0 %)
- Table 11: Raw data for Adenovirus cell sensitivity (2,0 %)







Table 12: Raw data for Norovirus cell sensitivity (2,0 %)

Table 13: Summary of results with VIRUTON PULVER and Adenovirus

Table 14: Summary of results with VIRUTON PULVER and Norovirus

Legend to the Figures

Figure 1: Adenovirus-inactivating properties of VIRUTON PULVER + ref. Glutaraldehyde (50 %)

Figure 2: Norovirus-inactivating properties of VIRUTON PULVER + ref. Glutaraldehyde (50 %)





Table 1: Raw data for VIRUTON PULVER (2,0 %) tested against Adenovirus at 20 °C (quantal test; 8 wells)

Product	Concentration	Interfering substance	Contact time				Dilution	range (log	g 10)		
				2	3	4	5	6	7	8	9
Test product	2,0%	3,0 g/L BSA +	15 min	tttt	0000	0000	0000	0000	0000	0000	0000
_		3,0 ml/L ShE		tttt	0000	0000	0000	0000	0000	0000	0000
Test product	2,0 %	3,0 g/L BSA +	n. a.	tttt	0000	0000	0000	n.d.	n.d.	n.d.	n.d.
cytotoxicity		3,0 ml/L ShE		tttt	0000	0000	0000				
Virus	n.a.	3,0 g/L BSA +	0 min	4444	4444	4444	4444	4444	4444	4444	3343
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3443
Virus	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4444	4444	3333
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3443
Water	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4433	2222	0000
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4443	2222	0000

n.d. – not done

0 – no virus present

T-cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)



Table 2: Raw data for VIRUTON PULVER (1,0 %) tested against Adenovirus at 20 °C (quantal test; 8 wells)

Product	Concentration	Interfering substance	Contact time				Dilution	range (log	g 10)		
				2	3	4	5	6	7	8	9
Test product	1,0%	3,0 g/L BSA +	15 min	tttt	0000	0000	0000	0000	0000	0000	0000
_		3,0 ml/L ShE		tttt	0000	0000	0000	0000	0000	0000	0000
Test product	1,0 %	3,0 g/L BSA +	n. a.	tttt	0000	0000	0000	n.d.	n.d.	n.d.	n.d.
cytotoxicity		3,0 ml/L ShE		tttt	0000	0000	0000				
Virus	n. a.	3,0 g/L BSA +	0 min	4444	4444	4444	4444	4444	4444	4444	3343
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3443
Virus	n. a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4444	4444	3333
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3443
Water	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4433	2222	0000
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4443	2222	0000

n.d. - not done

0 – no virus present

t – cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)



Table 3: Raw data for VIRUTON PULVER (0,5 %) tested against Adenovirus at 20 °C (quantal test; 8 wells)

Product	Concentration	Interfering substance	Contact time				Dilution	range (log	g 10)		
				2	3	4	5	6	7	8	9
Test product	0,5%	3,0 g/L BSA +	15 min	tttt	2220	0000	0000	0000	0000	0000	0000
_		3,0 ml/L ShE		tttt	2222	2200	0000	0000	0000	0000	0000
Test product	0,5 %	3,0 g/L BSA +	n. a.	tttt	0000	0000	0000	n.d.	n.d.	n.d.	n.d.
cytotoxicity		3,0 ml/L ShE		tttt	0000	0000	0000				
Virus	n.a.	3,0 g/L BSA +	0 min	4444	4444	4444	4444	4444	4444	4444	3343
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3443
Virus	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4444	4444	3333
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3443
Water	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4433	2222	0000
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4443	2222	0000

n.d. – not done

0 – no virus present

T – cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)





Table 4: Raw data for VIRUTON PULVER (0,05 %) tested against Adenovirus at 20 °C (quantal test; 8 wells)

Product	Concentration	Interfering substance	Contact time				Dilution	range (log	g 10)		
				2	3	4	5	6	7	8	9
Test product	0,05%	3,0 g/L BSA +	15 min	tttt	4444	4334	2232	2000	0000	0000	0000
_		3,0 ml/L ShE		tttt	4444	4433	2332	0000	0000	0000	0000
Test product	0,05 %	3,0 g/L BSA +	n. a.	tttt	0000	0000	0000	n.d.	n.d.	n.d.	n.d.
cytotoxicity		3,0 ml/L ShE		tttt	0000	0000	0000				
Virus	n.a.	3,0 g/L BSA +	0 min	4444	4444	4444	4444	4444	4444	4444	3343
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3443
Virus	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4444	4444	3333
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3443
Water	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4433	2222	0000
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4443	2222	0000

n.d. – not done

0 – no virus present

T – cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)





Table 5: Raw data for VIRUTON PULVER (2,0 %) tested against Norovirus at 20 °C (quantal test; 8 wells)

Product	Concentration	Interfering	Contact				Dilution	range (log	g 10)		
		substance	time		ı	1	1		r	r	
				2	3	4	5	6	7	8	9
Test product	2,0%	3,0 g/L BSA +	15 min	tttt	0000	0000	0000	0000	0000	0000	0000
		3,0 ml/L ShE		tttt	0000	0000	0000	0000	0000	0000	0000
Test product	2,0 %	3,0 g/L BSA +	n. a.	tttt	0000	0000	0000	n.d.	n.d.	n.d.	n.d.
cytotoxicity		3,0 ml/L ShE		tttt	0000	0000	0000				
Virus	n.a.	3,0 g/L BSA +	0 min	4444	4444	4444	4444	4444	4444	4443	3322
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3233
Virus	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4444	4444	3333
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4433	2223
Water	n. a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	3333	2222	0000
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4433	2222	0000

n.d. – not done

0 – no virus present

T-cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)



Table 6: Raw data for VIRUTON PULVER (1,0 %) tested against Norovirus at 20 °C (quantal test; 8 wells)

Product	Concentration	Interfering substance	Contact time				Dilution	range (log	g 10)		
				2	3	4	5	6	7	8	9
Test product	1,0%	3,0 g/L BSA +	15 min	tttt	0000	0000	0000	0000	0000	0000	0000
		3,0 ml/L ShE		tttt	0000	0000	0000	0000	0000	0000	0000
Test product	1,0 %	3,0 g/L BSA +	n. a.	tttt	0000	0000	0000	n.d.	n.d.	n.d.	n.d.
cytotoxicity		3,0 ml/L ShE		tttt	0000	0000	0000				
Virus	n.a.	3,0 g/L BSA +	0 min	4444	4444	4444	4444	4444	4444	4443	3322
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3233
Virus	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4444	4444	3333
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4433	2223
Water	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	3333	2222	0000
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4433	2222	0000

n.d. - not done

0 – no virus present

t – cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)





Table 7: Raw data for VIRUTON PULVER (0,5 %) tested against Norovirus at 20 °C (quantal test; 8 wells)

Product	Concentration	Interfering substance	Contact time				Dilution	range (log	g 10)		
				2	3	4	5	6	7	8	9
Test product	0,5%	3,0 g/L BSA +	15 min	tttt	2200	0000	0000	0000	0000	0000	0000
_		3,0 ml/L ShE		tttt	0220	0000	0000	0000	0000	0000	0000
Test product	0,5 %	3,0 g/L BSA +	n. a.	tttt	0000	0000	0000	n.d.	n.d.	n.d.	n.d.
cytotoxicity		3,0 ml/L ShE		tttt	0000	0000	0000				
Virus	n.a.	3,0 g/L BSA +	0 min	4444	4444	4444	4444	4444	4444	4443	3322
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3233
Virus	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4444	4444	3333
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4433	2223
Water	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	3333	2222	0000
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4433	2222	0000

n.d. – not done

0 – no virus present

T – cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)





Table 8: Raw data for VIRUTON PULVER (0,05 %) tested against Norovirus at 20 °C (quantal test; 8 wells)

Product	Concentration	Interfering substance	Contact time				Dilution	range (log	g 10)		
				2	3	4	5	6	7	8	9
Test product	0,05%	3,0 g/L BSA +	15 min	tttt	4444	3434	2222	0000	0000	0000	0000
_		3,0 ml/L ShE		tttt	4444	4433	2222	0000	0000	0000	0000
Test product	0,05 %	3,0 g/L BSA +	n. a.	tttt	0000	0000	0000	n.d.	n.d.	n.d.	n.d.
cytotoxicity		3,0 ml/L ShE		tttt	0000	0000	0000				
Virus	n.a.	3,0 g/L BSA +	0 min	4444	4444	4444	4444	4444	4444	4443	3322
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4444	3233
Virus	n. a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	4444	4444	3333
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4444	4433	2223
Water	n.a.	3,0 g/L BSA +	15 min	4444	4444	4444	4444	4444	3333	2222	0000
control		3,0 ml/L ShE		4444	4444	4444	4444	4444	4433	2222	0000

n.d. – not done

0 – no virus present

T – cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)



Table 9: Raw data for Glutaraldehyde solution (50,0 %) tested at 20 °C (quantal test; 8 wells)

Product	Concentration	Interfering substance	Contact time				Dilution	range (lo	g 10)		
				2	3	4	5	6	7	8	9
Glutaraldehyde	50,0 %	PBS	5 min	tttt	tttt	tttt	0000	0000	0000	0000	0000
at Adenovirus	(m/V)			tttt	tttt	tttt	0000	0000	0000	0000	0000
Glutaraldehyde	50,0 %	PBS	n. a.	tttt	tttt	tttt	0000	n.d.	n.d.	n.d.	n.d.
CCL-2	(m/V)			tttt	tttt	tttt	0000				
cytotoxicity											
Adenovirus	n.a.	PBS	5 min	4444	4444	4444	4444	4444	4444	2323	0000
water control				4444	4444	4444	4444	4444	4444	2222	0000
Glutaraldehyde	50,0 %	PBS	5 min	tttt	tttt	tttt	0000	0000	0000	0000	0000
at Norovirus	(m/V)			tttt	tttt	tttt	0000	0000	0000	0000	0000
Glutaraldehyde	50,0 %	PBS	n.a.	tttt	tttt	tttt	0000	n.d.	n.d.	n.d.	n.d.
TIB-71	(m/V)			tttt	tttt	tttt	0000				
cytotoxicity											
Norovirus	n. a.	PBS	5 min	4444	4444	4444	4444	4444	4444	3223	0000
water control				4444	4444	4444	4444	4444	4444	2233	0000

n.d. - not done

0 – no virus present

T-cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)



Table 10: Raw data for control efficacy for suppression of VIRUTON PULVER disinfectant activity 1,0 %,

	Concentration	Interfering substance				Dilution	range (log	g 10)		
			2	3	4	5	6	7	8	9
Adenovirus	1,0 %	3,0 g/L BSA +	tttt	4444	4444	4444	4444	3443	2222	0000
		3,0 ml/L ShE	tttt	4444	4444	4444	4444	3333	2222	0000
Adenovirus	n.a.	3,0 g/L BSA +	4444	4444	4444	4444	4444	4444	4433	2000
control		3,0 ml/L ShE	4444	4444	4444	4444	4444	4444	4334	2022
Norovirus	1,0 %	3,0 g/L BSA +	tttt	4444	4444	4444	4444	4443	3322	0000
		3,0 ml/L ShE	tttt	4444	4444	4444	4444	3344	2333	0200
Norovirus	n.a.	3,0 g/L BSA +	4444	4444	4444	4444	4444	4443	3233	2220
control		3,0 ml/L ShE	4444	4444	4444	4444	4444	4444	3332	0022

n.d. – not done

0 – no virus present

t – cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)



Table 11: Raw data (Adenovirus) for cell sensitivity (2,0%)

Product	Dilution	Interfering substance				Dilution	range (lo	g 10)				
			2 3 4 5 6 7 8 9									
PBS	-	-	4444	4444	4444	4444	4444	4444	2333	0002		
			4444	4444	4444	4444	4444	4444	2333	0200		
VIRUTON	1:1000	3,0 g/L BSA +	4444	4444	4444	4444	4444	4444	2222	0000		
PULVER		3,0 ml/L ShE	4444	4444	4444	4444	4444	4333	2222	0000		

n.d. – not done

0 – no virus present

t-cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)



Table 12: Raw data (Norovirus) for cell sensitivity (2,0%)

Product	Dilution	Interfering				D:1-4:-	(1-	- 10)				
		substance		T	T	Dilution	range (lo	g 10)	1	,		
			2	3	4	5	6	7	8	9		
PBS	-	-	4444	4444	4444	4444	4444	4444	2333	0000		
			4444	4444	4444	4444	4444	4444	2333	0022		
VIRUTON	1:1000	3,0 g/L BSA +	4444	4444	4444	4444	4444	3333	0222	0000		
PULVER		3,0 ml/L ShE	E 4444 4444 4444 4444 4444 4434 2202 0000									

n.d. – not done

0 – no virus present

t – cytotoxic

1 to 4 – virus present (degree of CPE in 8 cells units) (wells of microtiter plates)



Table 13: Summary of results of activity of the product VIRUTON PULVER with Adenovirus type 5 ATCC VR-5

					log ₁₀ TCID ₅₀ / ml				Reduction with
Assay	Interfering	Concentration	Cytotoxicity		after			the confidence	
	substance		level	0 min	5 min	15 min	30 min	60 min	interval of 95 %
			\log_{10}						
	3,0 g/L BSA +	2,0 %	2,5			≤2,50			≥6,00±0,00
Test product	3,0 ml/L ShE	1,0 %	2,5			≤2,50			≥6,00±0,00
		0,5 %	2,5			$3,63\pm0,37$			4,88±0,74
		0,05 %	2,5			5,63±0,13			2,88±0,25
Water control	3,0 g/L BSA +	0%	n.a.			8,50±0,00			
	3,0 ml/L ShE								
Virus control	3,0 g/L BSA +	n.a.	n.a.	9,50±0,00		9,50	9,00		n.a.
	3,0 ml/L ShE								
Glutaraldehyde	PBS	50 % (w:v)	4,5		≤4,50				≥4,00
Water control for	PBS	n.a.	n.a.		8,50				n.a.
Glutaraldehyde									
Suppression of	3,0 g/L BSA +	1,0 %	2,5				8,50		n.a.
disinfectant activity	3,0 ml/L ShE								
Cell sensitivity	PBS	n.a.	n.a.					8,75	n.a.
control									
PBS									
Cell sensitivity	3,0 g/L BSA +	1:1000	n.a.					8,50	n.a.
control test product	3,0 ml/L ShE								





Table 14: Summary of results of activity of the product VIRUTON PULVER with Murine norovirus strain S99 Berlin

					log ₁₀ TCID ₅₀ / ml				Reduction with
Assay	Interfering	Concentration	Cytotoxicity		after			the confidence	
	substance		level	0 min	5 min	15 min	30 min	60 min	interval of 95 %
			\log_{10}						
	3,0 g/L BSA +	2,0 %	2,5			≤2,50			≥6,00±0,00
Test product	3,0 ml/L ShE	1,0 %	2,5			≤2,50			≥6,00±0,00
		0,5 %	2,5			3,00±0,50			5,50±1,00
		0,05 %	2,5			5,50±0,00			3,00±0,00
Water control	3,0 g/L BSA +	0%	n.a.			8,50±0,00			
	3,0 ml/L ShE								
Virus control	3,0 g/L BSA +	n.a.	n.a.	9,50±0,00		9,50	9,00		n.a.
	3,0 ml/L ShE								
Glutaraldehyde	PBS	50 % (w:v)	4,5		≤4,50				≥4,00
Virus control for	PBS	n.a.	n.a.		8,50				n.a.
Glutaraldehyde									
Suppression of	3,0 g/L BSA +	0,5 %	1,5				8,63		n.a.
disinfectant activity	3,0 ml/L ShE								
Cell sensitivity	PBS	n.a.	n.a.					8,75	n.a.
control									
PBS									
Cell sensitivity	3,0 g/L BSA +	1:1000	n.a.					8,25	n.a.
control test product	3,0 ml/L ShE								



Figure 1

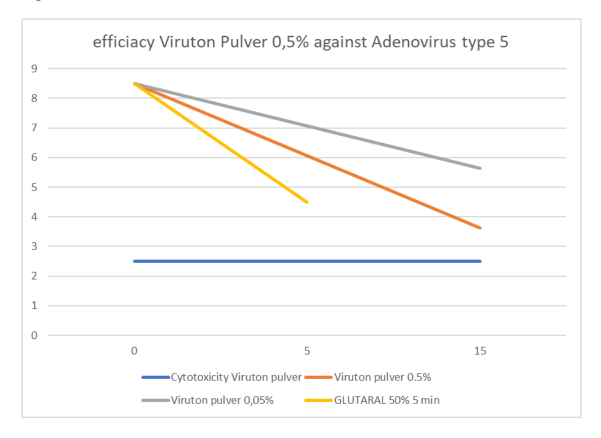




Figure 2

