

Toulouse, January 6th 2023

STUDY 20 - 2793

TEST REPORT N° 23-1990

Standard NF EN 17272 (Avril 2020) Determination of virucidal activity for aerial surface disinfection processes Human Coronavirus 229E

Medical area

Clean condition / Obligatory conditions

Promotor Company registration SIREN 448974253

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1. Test Laboratory

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2. Identification of the aerial disinfection system

Product: Formula N-5
Batch: A291020FD/1
Expiry date: 12/2022

Date of receipt: January/04/2021

Inetrnal code: 20-2793-1

Device: Diffuser PX-00

Serial number: 172X731

Concentration of disinfectant in the room: 5 mL/m³

One treatment - recovery of the discs after 60 minutes waiting at the end of the diffusion.

Promotor: Company registration SIREN 448974253

Storage conditions: Ambiant temperature

Period of testing: December 2021

Active Substance: Hydrogen peroxide (7.9%)

3- Experimental conditions

3-1 Virus/Receiving cells

Virus

Name Human Coronavirus 229E

Origin: ATCC
ATCC reference: VR-740
Batch number supplier: 58505270

Internal number Batch: SS-210920 (passage N°2)

Receiving cells

Name Vero cells
Origin: ATCC
ATCC reference: CCL-81
Batch number ATCC: 3372621

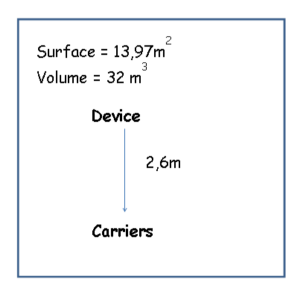
Internal number Batch: WCB-080205 (passage N°12)

3-2 Carriers

The selected tests surfaces are stainless steel discs, flats, corresponding to the requirements of paragraph 5.2.3.1 of the standard. The suppliers are MERCIER CLAUSSE.

3-3 Conditions of use of the device/product

- Room:



Relative humidity: start of test 64% - end of test 60% (requirement 50 - 70%). Temperature: start of test $18.5^{\circ}C$ - end of test $16.4^{\circ}C$ (requirement $20^{\circ}C_{\pm}2^{\circ}C$).

Test room volume: 32m³

- Carriers placement:

Distance between the device and the carriers: 2,6m (Annex B - table B.1).

The carriers were placed at a height of 1.31m, in a vertical position, towards the opposite side of the device

3-4 Interfering substance and culture media

-Interfering substance: BSA fraction V at 0,3g/l (Batch N°437)

-Culture media: MEM (Batch N°2980)

4- Validations Protocol

4-1 Control of sensitivity of cells to virus

- Add one volume of solution S or PBS + one volume of cellular suspension at 2.10^5 cells/ml for one hour in water bath at $36^{\circ}C\pm1^{\circ}C$
- The cells are centrifuged at 1600trs/min for 10 min and resuspended in culture media
- The virus is diluted from 1/4 to 1/4 on a 96-well microplate (15 dilutions)
- Add 100 μ l of cell suspension treated (Solution S) or not treated (PBS control) to each well of the microplate
- Incubate for 72 hours

The difference of titre reduction between cells treated by the solution S and cells treated by PBS shall be < 1 lg.

4-2 Control of efficiency for suppression of disinfectant activity

- Add 1 volume of BSA + 1 volume of virus suspension + 1 volume of solution S or distilled water
- Leave the mixture in the ice bath for 60 min at room temperature

5- Titration method

- Titrate the virus (method titration on cell in suspension) by following steps:
- Serial dilutions (1/4) are realized with culture medium in the glass tube
- Transfer 0,1 ml of each dilution into eight wells of a microplate plaque
- The last row of eight wells will receive 0,1 ml of culture medium (control untreated cells)
- Add 0,1 ml of cell suspension at 2.10⁵ cell/ml.
- Incubate for 72 hours at 36 ° $C \pm 1$ ° C under 5% $CO_2 \pm 2$ %.
- The viral cytopathic effect is read by using an inverted microscope

The estimated of infectious unite is determined by method KARBER-SPAERMAN calculating the negative logarithm of 50% endpoint (IgDIC50) by the following formula:

 $IgDICT50 = negative\ logarithm\ of\ the\ highest\ concentration\ of\ virus\ -\ [(Sum\ of\%\ affected\ to\ each\ dilution/100\ -\ 0.5)\ X\ (Ig\ dilution)]$

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6- Results

Virus suspension title assay: IgDICT50 = 8.13

No cytotoxicity was observed on the carrier without treatment which has been pretreated with the aerial disinfection system according to treatment.

Test 17/12/2021		
Start: 18.5°C / 64% RH	Degree of cytopathogenic	Logarithmic reduction
End: 18.4°C / 60% RH	effect (log)	
Sensitivity of cells to virus		
- With treatment (S1)		
Carrier 1	7.00	
Carrier 2	7.13	
Average		Difference <1 lg.
	7.07	
- Without traitement (S2)		
Carrier 1	6.88	
Efficiency for suppression of disinfectant activity		
- With treatment (D1)		
Carrier1	6.38	
Carrier 2	6.63	Difference <0,5 lg.
Average	6.50	
- Without traitement (D2)	-1	
Carrier 1	6.88	
Test control		
Carrier1	5.50	
Carrier 2	5.00	
Average	5.25	
Assay		
Support 1	0.88	
Support 2	1.00	4.33
Support 3	0.88	
Average	0.92	

7- Conclusion

According to the conditions of test for the standard NF EN 17272 (Avril 2020), the couple device/product: Diffuser PX-00 serial number 172X731 / Formula N-5 Batch A291020FD/1 for a use in medical area under clean condition, leds to a virucidal activity against Human coronavirus 229E (log reduction \geq 4), after treatment at 5 mL/m³ and 60 minutes waiting time.

These results are only valid for the product tested and apply to the sample as received.