

CERTIFICATE OF INVESTIGATION STUDY

STUDY OF VALIDATION OF THE EFFICIENCY OF HAIER AIR CONDITIONER ON SARS-COV-2 BY NO GLP VIRAL CLEARANCE STUDY (FIO)

Study number: 1198/02
Study report for:

Sponsor: Qingdao Haier Air Conditioner General Co., LTD

#1, Haier Road
Haier High-tech Industrial Park, Laoshan District
Qingdao City, Shandong Province
China

Submitted by:

Test Facility: TEXCELL

Test facility management: Boisson Bruno

Signature:

Genavenir 5 1 rue Pierre Fontaine

91058 Evry cedex

France

In Haier conditions of use:

An inactivation and reduction titer of more 4.08 Log (99.991%) of Sars-Cov-2 in 24 hours was demonstrated on the device in 6.7m³ test lab.

Texcell

Immeuble Genavenir 5

1 rue Pierre Fontaine F-91058 EVRY Cedex

Téléphone : + 33 (0)1 60 91 33 10 Télécopie : + 33(0)1 64 93 33 24

www.texcell.com

Study number: 1198/02

March 29 2022 Page 2 of 30

CONTENT

1.	PE	RSONNEL INVOLVED IN THE PROJECT	3
2.	MA	TERIAL AND METHODS	3
	2.1.	Viruses (spiking test system)	3
	2.2.	Cells (titration test system)	4
	2.3.	Medium	5
3.	EX	PERIMENTS	5
	3.1.	Process Design	5
4.	Ini	FECTIVITY METHODS	6
	4.1.	Titration assay	6
	4.2.	Acceptance criteria of the titration assay	9
	4.3.	Determination of the viral titer	9
5.	RE	DUCTION FACTOR CALCULATION	12
6.	TA	BLES OF RESULTS	15
CC	ONCLUS	SIONS	24



1. PERSONNEL INVOLVED IN THE PROJECT

From Texcell

Bruno BOISSON

Test Facility Management

2 9 MARS 2022

Date of signature

Virology and Titration Group:

Franck Gaston Luck TY

2. MATERIAL AND METHODS

2.1. VIRUSES (SPIKING TEST SYSTEM)

• Severe Acute Respiratory Syndrome-Related Coronavirus 2 (SARS-CoV-2): (Istituto Nationale malattei infettive (INMI) "Lazzaro Spallanzani")

The human strain of SARS-COV-2 was obtained from a viral strain isolated in Italy at Istituto Nationale malattei infettive (INMI) from a sample collected on January 29, 2020. The virus identity has been confirmed by complete sequencing. The complete sequence was submitted to GenBank (SARS-Cov-2/INMI1-Isolate/2020/Italy: MT066156) and is available on the GISAID website (BetaVov/Italy/INMI1-isl/2020: EPI_ISL_410545) upon registration. 2019-nCoV/Italy-INMI1 strain was used as a virus source and propagated on vero cells as described in the journal articles listed below:

Capobianchi MR, Rueca M, Messina F, Giombini E, Carletti F, Colavita F, Castilletti C, Lalle E, Bordi L, Vairo F, Nicastri E, Ippolito G, Gruber CEM, Bartolini B. Molecular characterization of SARS-CoV-2 from the first case of COVID-19 in Italy. Clin Microbiol Infect. 2020 Jul;26(7):954-956. doi: 10.1016/j.cmi.2020.03.025.

Corman VM, Landt O, Kaiser M, Molenkamp R, Meijer A, Chu DKW, Bleicker T, Brünink S, Schneider J, Schmidt ML, Mulders DGJC, Haagmans BL, van der Veer B, van den Brink S, Wijsman L, Goderski G, Romette JL, Ellis J, Zambon M, Peiris M, Goossens H, Reusken C, Koopmans MPG, Drosten C. Detection of 2019 novel coronavirus (2019-nCoV) by real-time RT-PCR. Euro Surveill. 2020 Jan;25(3). doi: 10.2807/1560-7917.ES.2020.25.3.2000045.
Ogando NS, Dalebout TJ, Zevenhoven-Dobbe JC, Limpens RWAL, van der Meer Y, Caly L, Druce J, de Vries JJC, Kikkert M, Bárcena M, Sidorov I, Snijder EJ. SARS-coronavirus-2 replication in Vero E6 cells: replication kinetics, rapid adaptation and cytopathology. J Gen Virol. 2020 Jun 22. doi: 10.1099/jgv.0.001453.



Park WB, Kwon NJ, Choi SJ, Kang CK, Choe PG, Kim JY, Yun J, Lee GW, Seong MW, Kim NJ, Seo JS, Oh MD. Virus Isolation from the First Patient with SARS-CoV-2 in Korea. J Korean Med Sci. 2020 Feb 24;35(7):e84. doi: 10.3346/jkms.2020.35.e84.

Chu H, Chan JF, Yuen TT, Shuai H, Yuan S, Wang Y, Hu B, Yip CC, Tsang JO, Huang X, Chai Y, Yang D, Hou Y, Chik KK, Zhang X, Fung AY, Tsoi HW, Cai JP, Chan WM, Ip JD, Chu AW, Zhou J, Lung DC, Kok KH, To KK, Tsang OT, Chan KH, Yuen KY. Comparative tropism, replication kinetics, and cell damage profiling of SARS-CoV-2 and SARS-CoV with implications for clinical manifestations, transmissibility, and laboratory studies of COVID-19: an observational study. Lancet Microbe. 2020 May;1(1):e14-e23. doi: 10.1016/S2666-5247(20)30004-5.

2.1.1. Virus stock

Main characteristics of Sars-Cov-2 virus are described in the table below:

Virus	SARS-COV-2
Name	Severe Acute Respiratory Syndrome-Related Coronavirus 2
Family	Coronaviridae
Subfamily / Genus	Orthocoronavirus
Size in diameter	90-110 nm
Genome	One molecule of single-stranded RNA
Envelope	Enveloped
Strain	2019-nCoV/Italy-INMI1 strain
Production standard operating procedure (SOP)	TE1129
Titer of the virus stock used for spiking	≥ 6 Log ₁₀ TCID ₅₀ /mL
experiments	(50 % tissue culture infective dose per milliliter)
Titration assay and cells	Virus titer is determined by end-point dilution titration assay on Vero cells (operating procedure TE1088).

2.2. CELLS (TITRATION TEST SYSTEM)

(Freshney R.I., 1989, ATCC)

The cells are grown in accordance with Texcell's operating procedures TE1002, TE3001 and TE3011.

• **Vero cells**: (Institut Pasteur, Medical Virology Laboratory) (Simizu B. and Terasima T. 1988; Simizu B. et al., 1967).

Isolated from C. aethiops kidney on 27 Mar 1962. Vero cells are a lineage of <u>cells</u> used in <u>cell cultures</u>. The 'Vero' lineage was isolated from <u>kidney epithelial</u> cells extracted from an <u>African green monkey</u> (<u>Chlorocebus</u> sp.; formerly called Cercopithecus aethiops, this group of monkeys has been split into several different species). The lineage was developed on 27 March 1962, by Yasumura and Kawakita at the <u>Chiba University</u> in <u>Chiba</u>, <u>Japan</u>. The original cell line was named "<u>Vero</u>" after an <u>abbreviation</u> of <u>verda reno</u>, which means "green kidney" in <u>Esperanto</u>.

This is a cell line with the hypodiploid chromosome count. The modal chromosome number was 58 occurring in 66% of cells. In most cells, over 50% of the chromosomes in each cell complement belonged to structurally altered marker chromosomes. Normal A3, A4, B4, and B5 were absent; B2, B3 and B7 were occasionally paired; and B9, C1 and C5 were mostly paired. The rate of cells with higher ploidies was 1.7%. Other chromosomes were



mostly present in single copy. These cell lines from ATCC (CCL-81, CRL-1587, CRL1586), are used for the propagation, assay and isolation of numerous viruses like coronaviruses. This line is a clone from Vero 76. Vero E6 cells are known to show some contact inhibition, and are often suitable for propagating viruses that replicate slowly and was therefore used to isolate the strain of SARS-COV-2 virus used in this study. Virucidy assays are performed using either the Vero cell line from ATCC CCL-81 or from Institut Pasteur (IP)

2.3. MEDIUM

Vero cells:

Dilution medium: DMEM 2% glutamine, 1% gentamicine

Cell Culture medium: DMEM 4% SVF, 2% glutamine, 1% gentamicine

Titration medium: DMEM 2% glutamine, 1% gentamicine

3. EXPERIMENTS

3.1. PROCESS DESIGN

As requested by the sponsor the process below will be evaluated as one step:

As requested by the sponsor the process below will be evaluated as one step: Haier device

(A) Inactivation

- Virus gauze preparation with SARS-COV-2
- Place virus 2 gauze and Haier device into a workhouse (1 gauze on the device and 1 gauze at 1 meter in front of the DIU for horizontal position, adjust its height to 1.5 meters high)
- After the required incubation time (24h, 4h, 1h) the virus gauze is removed
- virus gauze in the test group and control group was eluted thoroughly with 10ml eluent, respectively
- Virus titration and calculation virus reduction factor R
 Process Duration (X) = 24h, 4h and 1h

Titrations:

Virus titration will be performed by end-point dilution assay (TCID50) for SARS-COV-2. Samples obtained during the spiking experiments will titrated immediately with the appropriate controls (depending of the Preliminary Assay result).



Study number: 1198/02 March 29 2022
Page 6 of 30

4. INFECTIVITY METHODS

4.1. TITRATION ASSAY

The samples are titrated according to operating procedure TE1088.

4.1.1. Principle of titration

The titration method is a quantitative assay in which the virus titer measurement is based on the detection of virus production in the infected cells, by observation of a specific cytopathic effect.

*Serial dilutions titration

Briefly:

Test sample is diluted with medium by serial 3-fold dilutions (eight replicates are performed for each dilution) across the 96 well plate (sample dilution plate).

Each well from the "sample dilution plate" is then inoculated on the corresponding well of a new plate (sample titration plate).

Cell suspension is added to each well of the "sample titration plate" and the plates are then incubated at appropriate temperature with or without CO₂ atmosphere (depending on viruses). After a period of incubation allowing viral replication and infection of adjacent cells, depending on viruses.

- wells with foci are counted after infection by observation under inverted light microscope.
- or a stain overlay (crystal violet) is added and wells are examined for cytopathic effect. The infected wells show up as clear areas whereas the non-infected wells are stained.

The infectious titer expressed as 50% tissue culture infective dose per milliliter ($TCID_{50}/mL$) is calculated using the Spearman-Kärber formula.

*Large Volume titration (LVT)

LVT assay could be performed (at the sponsor request) in order to improve the detection limit of the assay or the titer of the tested sample. Cells in 96 well plates or flasks (number according to desired sensitivity) are inoculated with a large volume of the lowest non-toxic and non-interfering dilution of the sample.

Cell suspension is then added to each well of the "sample titration plate" and the plates are incubated at appropriate temperature with or without CO₂ atmosphere (depending on viruses). After a period of incubation allowing viral replication and infection of adjacent cells, depending on viruses,

- wells with foci are counted after infection by observation under inverted light microscope.
- or a stain overlay (crystal violet) is added and wells are examined for cytopathic effect. The infected wells show up as clear areas whereas the non-infected wells are stained.

The infectious titer expressed as 50% tissue culture infective dose per milliliter ($TCID_{50}/mL$) is calculated using the appropriate formula.

4.1.2. Titration assay controls

*Negative control N1 (cell reference control)

During titration assay, in each 96-well plate, 8 wells are prepared as cell reference control. These cells are prepared in the same conditions as those used for the titration of the samples generated during the viral clearance experiments except that they are inoculated with unspiked medium.



*Positive reference control (virus reference control)

During each titration assay, a stock of each virus prepared at low final concentration of approximately $10^5 - 10^7$ TCID₅₀/mL (depending on virus) and used as virus reference control is titrated in the same conditions as those used for the titration of the samples generated during experiments.

4.1.3. Spiking experiments controls

*Cytotoxicity test control(s)

The evaluation of the cytotoxic effect is carried out by visual observation under inverted light microscope. The quality of the cell monolayer (confluence, refringence, aspect) of the tested samples is compared with that of the cell reference control N1.

Negative controls (N2 and N3) and each dilution of samples for which no total cytopathic effect is observed are evaluated for cytotoxicity by comparison with the cell reference control N1. Similar serial dilutions as those used for the titration assays are applied.

For a defined sample, the non-cytotoxic dilution is reached when no significant difference is observed in the 3-fold serial diluted sample compared with the cell reference control.

*Storage control(s)

The evaluation of the effect of the storage conditions on virus in the tested sample during the storage at \leq -70°C is carried out by comparison of the titers obtained in medium and in diluted sample after storage.

This evaluation is carried out with the starting material(s), and/or the final material(s) obtained from the Mock run when the generated samples are stored until titration.

The study is described as follows:

- The starting material and/or the final material is diluted at a defined dilution, and then spiked with virus (positive control or virus stock) at a low final concentration of approximately 10³-10⁴ TCID₅₀/mL.
- In parallel, medium is also spiked with the same virus (positive control or virus stock) to reach the same concentration as the starting material and/or final material samples.

Both storage controls samples are then stored at \leq -70°C until titration. These storage controls are prepared and titrated with the experiments generated samples

After titration, there is no storage effect on virus, when the difference between the two titers (titer in medium compared with titer in starting and/or final material(s)) is less than or equal to 1Log₁₀.

*Viral interference assay(s)

The evaluation of the potential interference of the lowest non-cytotoxic dilution of the sample on virus during the incubation period is carried out by comparison of the results obtained during the titration of the virus stock with medium, and the virus stock with diluted sample (final material/starting material and/or intermediate product).

The evaluation is carried out with the final material(s) (obtained from the mock run).

The results are used for the calculation of the reduction factor of the step.

The study of sample interference on virus titration is described as follows:



Study number: 1198/02 March 29 2022
Page 8 of 30

• First, the virus reference control (titration reference) at a low final concentration of approximately 10⁵-10⁷ TCID₅₀/mL, is titrated in the usual conditions i.e. serial 3-fold dilutions using medium as diluting medium

• In parallel, the same virus reference control is titrated by serial 3-fold dilutions using the lowest non-cytotoxic dilution of the sample to be tested as diluting medium.

The evaluation of the potential interference of the lowest non-cytotoxic dilution of the sample with the virus during the incubation period is carried out by comparison of the titers obtained in medium versus titers obtained in diluted sample.

For the defined non-cytotoxic dilution of the sample, there is no interference when the difference between the two titers is less than or equal to 1Log_{10} . This non-cytotoxic and non-interfering dilution is then used for the calculation of the infectious titer of the sample.



Study number: 1198/02 March 29 2022
Page 9 of 30

4.2. ACCEPTANCE CRITERIA OF THE TITRATION ASSAY

During each titration assay, a virus stock prepared at low final concentration of approximately $10^5 - 10^7$ TCID₅₀/mL (depending on virus), is used as reference control.

The titration assay is retained when:

- the cell reference control (N1) for each titration plate is conform to the expected result,
- the infectious titer of the virus reference control obtained is in the expected range.

4.3. DETERMINATION OF THE VIRAL TITER

(Schwartz D., 1993; Kaplan M. and Koprowski H., 1973)

Three situations may be predicted concerning the calculation of the viral titer.

Case	Subcase	Titer (T)
Infectious particles detected ≥ 12.5% positive wells/total tested wells		$T = T_{SK}$
Few infectious particles detected < 12.5% positive wells/total tested wells	$T_{\text{MaxL}} > dl$ $dl > T_{\text{MaxL}}$	$T = T_{MaxL}$ $T = dl$
No infectious particles detected 0% positive well/total tested wells		T < dl

with:

T = titer retained for the calculation of the reduction factor

 T_{SK} = infectious titer using the simplified Spearman-Kärber formula (Section 2.3.1)

 T_{MaxL} = infectious titer using the Maximum likelihood estimation (Section 2.3.2)

dl = detection limit using the Poisson formula with 95% precision (Section 2.3.3)

4.3.1. Calculation of the TCID₅₀ using Spearman-Kärber formula

The TCID₅₀ is evaluated by quantitative assay and defined as the virus dose capable of infecting 50% of the inoculated cultures.

The viral titer, T, expressed as the 50% tissue culture infective dose per milliliter (TCID $_{50}$ /mL), is defined by its mean value, m(T), and its confidence interval.

m(T) can be calculated according to the following formula:

$$m(T) = \frac{1}{Vo} 10m(a)$$

with v_0 = volume per replicate

"a" is also defined by its mean, m(a), and its standard deviation, S(a).

The viral titer can be calculated using Spearman-Kärber (SK) formula (Payment P. and Trudel M. 1993).

*This method is applicable firstly, in situations where cytopathic effect is observed, ranging from 0% to 100% of positive replicas per dilution in a same titration plate.

m(a) = T_{SK}, and S(a) is calculated using the following simplified Spearman-Kärber formula



$$m(a) = T_{SK} = -a_0 + \frac{k}{2} - k \sum_{i} p_i$$
 and $S(a)^2 = k^2 \sum_{i} \frac{p_i (1 - p_i)}{n_i - 1}$

with:

 $a = Log_{10}$ of the titer relative to the test volume

 $a_0 = Log_{10}$ of the reciprocal of the lowest dilution for which all wells are positive

 $k = Log_{10}$ of the dilution factor

 p_i = proportion of positive wells at the non-cytotoxic dilution d_i / r_i

r_i = number of positive wells at the non-cytotoxic dilution d_i

 n_i = number of replicates at the non-cytotoxic dilution d_i .

With a 95% precision, the confidence interval of "a" is the following:

$$a^{min} \le a \le a^{max}$$
 with: $a^{min} = m(a) - 2$ S(a)
$$a^{max} = m(a) + 2$$
 S(a)

With a 95% precision, the confidence interval of the titer T is the following:

$$T^{min} \leq T \leq T^{max} \text{ with:}$$

$$T^{min} = \frac{1}{Vo} 10^{amin}; T^{max} = \frac{1}{Vo} 10^{amax}$$

The dilutions of the samples retained for the calculation of the infectious titers are those for which no cytotoxicity is observed.

*Secondly, when less than 100% but \geq 12.5% of positive replicas per dilution is obtained for the lowest non-cytotoxic dilution tested, the virus titer is calculated assuming that the sample contains sufficient virus to infect 100% of tested wells at the previous serial dilution (worst-case).

*Total virus load

The viral load L is defined by its mean value, m(L), and its confidence interval.

$$m(L) = \frac{m(T)V_t}{c}$$

$$L\min \le L \le L\max: \qquad L^{min} = \frac{T^{min}V_t}{c} \text{ and } L^{max} = \frac{T^{max}V_t}{c}$$

with:

c = concentration factor of the ultracentrifugation (c = 1 when the samples are not ultracentrifuged).

 V_t = total volume of the sample during the scaled down process.

Study number: 1198/02

4.3.2. Large Volume Titration assay: Maximum Likelihood estimation

During LVT assay, when few positive wells are detected (< 12.5% of all tested wells), viral titers (TMaxL) in samples are calculated according to Maximum Likelihood estimation (Agut H., Calvez V., Barin F, 1997) as follows:

$$T_{MaxL} = (Ln (N/P) x d x (1000/v)) / Ln2$$

with:

 T_{MaxL} = titer relative to the test volume (TCID₅₀/mL)

N = number of all tested wells

P = number of negative wells

d = non-cytotoxic dilution factor of the sample

v = tested volume per well

 $Ln2 \approx 0.69 = corrective factor to convert PFU/mL into TCID₅₀/mL.$

When significant positive wells are detected (>12.5% of all tested wells) viral titers (T_{MaxL}) in samples are calculated using the Spearman-Kärber formula as described above.

*Total virus load

The viral load L is defined by its mean value, m(L_{MaxL}).

$$m(L_{MaxL}) = \frac{m(T_{MaxL})V_t}{c}$$

with:

c = concentration factor of the ultracentrifugation (<math>c = 1 when the samples are not ultracentrifuged).

 V_t = total volume of the sample during the scaled down process.

4.3.3. Detection limit of titration assay

When a sample contains a low concentration of infectious virus and only a fraction of the sample is tested for titration, there is a probability that the result of the tested fraction will be negative due to random (and unequal) distribution of the virus throughout the sample.

The detection limit, dl, for the titration assay corresponds to the lower theoretical titer which results in the detection of one infectious particle in one of the replicates performed. Since the infectious particle would be detected in a volume V_c (mL) of the dilution d_c , the detection limit, dl, is calculated with a 95% precision according to the Poisson formula (Löwer J., 1991):

$$p(95\%) = e^{-dl} [V_c d_c] = 0.05$$

i.e.
$$dl = \frac{-\ln(0.05)}{V_c d_c}$$

with:

$$V_c = [v_0 \ n_c]$$

v_o = volume per replicate

 n_c = number of replicates at non-cytotoxic dilution d_c

dl = detection limit



The non-cytotoxic dilution of the samples for which no positive wells are detected is then retained for the calculation of the infectious titer using the Poisson formula.

The infectious titers calculated with the Poisson formula are expressed as PFU/mL and are divided by ln2 to be converted into $TCID_{50}/mL$.

*Detection limit of the viral load of a whole fraction

The detection limit, DL, is the minimal viral load which could be theoretically detected in the total volume of a fraction belonging to the scaled down process.

DL is calculated according to the following formula:

$$DL = \frac{dl V_t}{c}$$

with:

c = concentration factor of the ultracentrifugation (c = 1 when the samples are not ultracentrifuged),

Vt = total volume of the sample during the scaled down process,

dl = detection limit for a titration assay.

The detection limit for the pre and post-treatment material are DLi and DLf respectively.

5. REDUCTION FACTOR CALCULATION

In accordance with the regulatory documents, the virus reduction factor (R) of an individual purification or inactivation step is defined as the Log10 of the ratio of the virus load (Li) in the pre-treatment material (starting material) and the virus load (Lf) in the post-treatment material (final material) which is ready for use in the next step of the process.

R is defined by its mean, m(R), and its confidence interval, [Rmin \le R \le Rmax]. If Rmin and/or Rmax is < 0 (negative value), then m(R) will present as \approx 0 without confidence interval.

Different cases have to be considered related to the effect of the pre-treatment material on the virus.

*Case 1

A studied manufacturing process step consists in the monitoring of a viral inactivation directly linked to a solution used during the process ("in process fraction", sanitization solution, etc.). In other terms, a putative virucidal effect associated with a solution is evaluated in a kinetics study.



*Case 2: Treatment reduction factor calculation m(Rt)

The pre-treatment material has no significant inactivating effect, i.e. the mean reduction factor, $m(R_0)$ is lower than or equal to 1. $m(R_0)$ is defined as the Log₁₀ of the ratio of the virus load in the medium (L_{cm}) and the virus load in the pre-treatment material (L_i).

In this case, the reduction factor of the step corresponds to the reduction factor of the treatment $m(R_t)$.

*Case 3: Clearance factor calculation m(R)

The pre-treatment material has a significant inactivating effect (excluding Case 1) i.e. the mean reduction factor, $m(R_0)$ is higher than 1.

- In a first approach, the clearance factor of the step, m(R), is calculated as the sum of the reduction factor associated with the initial inactivating effect, $m(R_o)$, and the reduction factor of the treatment, $m(R_t)$. The initial load is the virus load in medium (L_{cm}).
- In a second approach, the reduction factor of the treatment, $m(R_t)$ is calculated with the virus load in the pre-treatment material (L_i) .

In practice,

- if the volume of post-treatment material (V_f) = volume of pre-treatment material $(V_i) \pm 5\%$ (v/v), the reduction and clearance factors are calculated with viral titers and then X = T
- Otherwise, the reduction and clearance factors are calculated with viral loads and then X = L.



CASES	SUBCASES	EVALUATION OF THE REDUCTION FACTOR R _t AND CLEARANCE FACTOR (R)
Case 1		R is evaluated according to the three subcases of Case 2, with $X_i = X_{cm}$
Case 2	$m(x_f) < DL_f$	$R_t > Log_{10}\left(\frac{m(X_i)}{DL_f}\right)$ with $Rmin \le R_t \le Rmax$
Case 2 $m(R_O) \le 1$	$X_{i}^{min} \ge X_{f}^{max}$ $X_{i}^{min} < X_{f}^{max}$	$m(R_t) = Log_{10}\left(\frac{m(X_i)}{m(X_f)}\right)$ with $Rmin \le R_t \le Rmax$
	$X_i^{min} < X_f^{max}$	$m(R_t)$ is calculated according to subcase 2 $R_t \approx 0$ if Rmin, R_t or Rmax < 0 (negative value)
Case 3	$m(X_i) < DL_i$	R_{t} cannot be evaluated $R_{o} \ge Log\left(\frac{m(X_{cm})}{DL_{t}}\right) \text{ with } Rmin \le R_{o} \le Rmax$
$m(R_0) \ge 1$	$m(X_i) \ge DL_i$	$\begin{split} R_t \text{ can be evaluated at sponsor request, otherwise } R &= R_t + R_o \text{ is calculated} \\ R \text{ is evaluated according to the three subcases of Case 2 with } X_i &= X_{cm} \\ m(R_o) &= Log_{10} \bigg(\frac{m(X_{cm})}{m(X_i)} \bigg) \text{ with } Rmin \leq R_o \leq Rmax \end{split}$

X_i = initial sample titre (Ti) or load (Li)

 X_f = final sample titre (Tf) or load (Lf)

X_{cm} = culture medium titre (Tcm) or load (Lcm)

m(X) = mean titre or load

 $Rmin = R - 2 \sqrt{var Ti + var Tf}$

 $Rmax = R + 2 \sqrt{var Ti + var Tf}$

Var = Variance =
$$S(a)^2 = k^2 \sum_{i=1}^{\infty} \frac{p_i(1-p_i)}{n_i-1}$$

with:

- k = Log of the dilution factor

- p = proportion of positive wells at the non-cytotoxic dilution d / r

- r = number of positive wells at the non-cytotoxic dilution d

- n = number of replicates at the non-cytotoxic dilution d.

6. TABLES OF RESULTS

Table 1: STUDY OF EVALUATION OF THE EFFICIENCY OF HAIER AIR CONDITIONER (1H) ON THE SARS-COV2 BY NO GLP VIRAL CLEARANCE STUDY (FIO)

		TABLE 1.1: CONTROLS	: CON	FROLS		
	Sample	Infe	ctions 1	Infectious Titer (T)	Reduction fa	Reduction factor evaluation (R)
epoo	Definition	epoo	Mean	Mean titer [m(T)] and confidence interval	Mean reduction factor m(R) and confidence interval	Definition
		Negati	Negative controls	rols		
		Positiv	Positive controls	ols		
VI	Spiked (addition of virus stock) medium	TV1	5.24	5.00 \le T \le 5.48	NA	Virus stock control before process
V2	Spiked (addition of virus stock) medium	TV2	5.24	5.00 ≤ T ≤ 5.48	NA	Virus stock control after 1 hour (old control)
L1	Spiked (addition of virus stock) gauze and recovery	TL1	5.00	5.80 ≤ T ≤ 6.11	NA	Virus stock recovery control before process
1.2	Spiked (addition of virus stock) gauze and recovery	TL2	4.04	3.81 ≤ T ≤ 4.28	NA	Virus stock recovery control after 1 hour (old control)
			*	TA CALOR		

Infectious titers are expressed as Log10 50% tissue culture infectious dose per milliliter (Log10 TCID50/mL).



Table 1 (continued): STUDY OF EVALUATION OF THE EFFICIENCY OF HAIER AIR CONDITIONER (1H) ON THE SARS-COV2 BY NO GLP VIRAL CLEARANCE STUDY (FIO)

			TABLE 1.	TABLE 1.2: CONTROLS			
	Sample	I	Infectious Titer (T)	er (T)		Reduction	Reduction factor evaluation (R)
epoo	Definition	code	Mean ti confid	Mean titer [m(T)] and confidence interval	Mean re m(R) ar	Mean reduction factor m(R) and confidence interval	Definition
			Comparaisor	Comparaison of positive controls	slo		
VI	Spiked (addition of virus stock) medium	TVI	5.24	5.00 ≤ T ≤ 5.48	NA	NA	Virus stock control
							Experiment effect on virus in hold
	Spiked (addition of virus stock) medium		5.24	5.00 \le T \le 5.48		Not significant	control
V2		TV2			0~		(comparaison TV1 and TV2)
							Experiment effect on virus in recovery
	Spiked (addition of virus stock) gauze			5.80 ≤ T ≤ 6.11		Not significant	gauze control
	and recovery	TL1	5.00		0~		(comparaison TV1 and TL1)
							Experiment effect on virus in recovery
	Spiked (addition of virus stock) gauze			3.81 ≤ T ≤ 4.28		Not significant	gauze hold control
L2	and recovery	TL2	4.04		= 0.95		(comparaison TL1 and TL2)



Table 1 (continued): STUDY OF EVALUATION OF THE EFFICIENCY OF HAIER AIR CONDITIONER (1H) ON THE SARS-COV2 BY NO GLP VIRAL CLEARANCE STUDY (FIO)

	Reduction factor evaluation (R)	Mean reduction factor m(R) and confidence interval [m(R)] = Log10 (L1) - Log10 (LS)		Virus stock recovery gauze control			R = 3.22 (99.939%)	R = 0.48 (66.886%)
	ıd (L)	Mean load $[m(L)=m(T) \times Vt]$		5.80 ≤T ≤ 6.11		ı.	1.78 1.56 ≤ T ≤ 2.00	NA (no virus)
	Virus load (L)	[m(I		5.00		e facto	1.78	1.16
RIMENTS	Vi	epoo		LL1		d: Clearanc	LS1	LS2
TABLE 1.3: SPIKING EXPERIMENTS	Infectious Titer (T)	Mean titer [m(T)] and confidence interval	Positive control	5.80 < T < 6.11	STUDY	Samples collected, recovery and diluted: Clearance factor	1.56 ≤ T ≤ 2.00	4.22 ≤ T ≤ 4.82
E 1.3:	ctious 7	Mean		5.00		ted, re	1.78	4.52
TABI	Infe	epoo		TL1		imples collec	TS1	TS2
	Vol	(Vt) mL		1		Sa	-	-
	Sample	definition		Spiked (addition of virus stock)			Gauze on the evaporator, 10 centimeters away from the center of UVC module in horizontal position 1 hour	Gauze at 1 meter in front of the DIU for horizontal position, adjust its height to 1.5 meters high
		epoo		L1			S1	S2

Infectious titers are expressed as 50% tissue culture infectious dose per milliliter (Log10 TCID50/mL). Viral loads are expressed as 50% tissue culture infectious dose (Log10 TCID50).



Table 2: STUDY OF EVALUATION OF THE EFFICIENCY OF HAIER AIR CONDITIONER (4H) ON THE SARS-COV2 BY NO GLP VIRAL CLEARANCE STUDY (FIO)

Infectious titers are expressed as Log10 50% tissue culture infectious dose per milliliter (Log10 TCID50/mL).



Study number: 1198/02

Table 2 (continued): STUDY OF EVALUATION OF THE EFFICIENCY OF HAIER AIR CONDITIONER S (4H) ON THE SARS-COV2 BY NO GLP VIRAL CLEARANCE STUDY (FIO)

			TABLE 2	TABLE 2.5: CONTROLS			
	Sample	I	Infectious Titer (T)	er (T)		Reduction	Reduction factor evaluation (R)
epoo	Definition	code	Mean ti confid	Mean titer [m(T)] and confidence interval	Mean re m(R) ar ii	Mean reduction factor m(R) and confidence interval	Definition
		,	Comparaison	Comparaison of positive controls	sle		
V1	Spiked (addition of virus stock) medium	TV1	5.12	4.90 ≤ T ≤ 5.34	NA	NA	Virus stock control
							Experiment effect on virus in hold
	Spiked (addition of virus stock) medium		4.76	4.60 < T < 4.92		Not significant	control
V2		TV2			= 0.36		(comparaison TV1 and TV2)
							Experiment effect on virus in recovery
	Spiked (addition of virus stock) gauze			5.08 \le T \le 5.15		Not significant	gauze control
Γ 1	and recovery	TL1	5.24		0~		(comparaison TV1 and TL1)
							Experiment effect on virus in recovery
	Spiked (addition of virus stock) gauze			3.81 ≤ T ≤ 4.28		Not significant	gauze hold control
L2	and recovery	TL2	4.04		= 0.84		(comparaison TL1 and TL2)



Table 2 (continued): STUDY OF EVALUATION OF THE EFFICIENCY OF HAIER AIR CONDITIONER (4H) ON THE SARS-COV2 BY NO GLP VIRAL CLEARANCE STUDY (FIO)

	Reduction factor evaluation (R)	Mean reduction factor m(R) and confidence interval [m(R)] = Log ₁₀ (L1) - Log ₁₀ (LS)		Virus stock recovery gauze control			R = 3.70 (99.980%)	R = 0.83 (85.208%)
	Virus load (L)	Mean load $[m(L)=m(T) \times Vt]$		$5.08 \le T \le 5.15$		factor	1.54 1.54 \le T \le 1.54	4.40 4.12 ≤ T ≤ 4.69
SIMENTS	Vi	epoo		LL1		d: Clearance	LS1	LS2
TABLE 2.6: SPIKING EXPERIMENTS	Infectious Titer (T)	Mean titer [m(T)] and confidence interval	Positive control	5.08 ≤ T ≤ 5.15	STUDY	Samples collected, recovery and diluted: Clearance factor	1.54 ≤ T ≤ 1.54	4.12 ≤ T ≤ 4.69
E 2.6:	ctions	Mear		5.24		ted, re	1.54	4.40
TABI	Jule	əpoo		ILI		amples collec	TS1	TS2
	Vol	(Vt) mL		1		S	1	-
	Sample	definition		Spiked (addition of virus stock)			Gauze on the evaporator, 10 centimeters away from the center of UVC module in horizontal position 4 hours	Gauze at 1 meter in front of the DIU for horizontal position, adjust its height to 1.5 meters high
		əpoo		L1			S1	. S2

Infectious titers are expressed as 50% tissue culture infectious dose per milliliter (Log10 TCID50/mL). Viral loads are expressed as 50% tissue culture infectious dose (Log10 TCID50).



STUDY OF EVALUATION OF THE EFFICIENCY OF HAIER AIR CONDITIONER (24H) ON THE SARS-COV2 BY NO GLP VIRAL CLEARANCE STUDY (FIO)

		TABLE 3.7: CONTROLS	CONT:	ROLS		
	Sample	Infe	Infectious Titer (T)	iter (T)	Reduction fac	Reduction factor evaluation (R)
əpoo	Definition	apoo	Mean t	Mean titer [m(T)] and confidence interval	Mean reduction factor m(R) and confidence interval	Definition
		Negati	Negative controls	rols		
		Positiv	Positive controls	ols		
V1	Spiked (addition of virus stock) medium	TV1	5.48	5.32 < T < 5.63	NA	Virus stock control before process
V2	Spiked (addition of virus stock) medium	TV2	4.40	4.15 ≤ T ≤ 4.66	NA	Virus stock control after 24 hours (old control)
L1	Spiked (addition of virus stock) gauze and recovery	TL1	5.24	5.00 ≤ T ≤ 5.48	NA	Virus stock recovery control before process
L2	Spiked (addition of virus stock) gauze and recovery	TL2	3.45	3.16 ≤ T ≤ 3.73	NA	Virus stock recovery control after 24 hours (old control)
			**	1		

Infectious titers are expressed as Log10 50% tissue culture infectious dose per milliliter (Log10 TCID50/mL).



Table 3 (continued): STUDY OF EVALUATION OF THE EFFICIENCY OF HAIER AIR CONDITIONER (24H) ON THE SARS-COV2 BY NO GLP VIRAL CLEARANCE STUDY (FIO)

			TABLE 3.	TABLE 3.8: CONTROLS			
	Sample	Ir	Infectious Titer (T)	er (T)		Reduction	Reduction factor evaluation (R)
əpoo	Definition	opoo	Mean tit confid	Mean titer [m(T)] and confidence interval	Mean reo m(R) an ir	Mean reduction factor m(R) and confidence interval	Definition
			Comparaisor	Comparaison of positive controls	sle		
V1	Spiked (addition of virus stock) medium	TV1	5.48	5.32 ≤ T ≤ 5.63	NA	NA	Virus stock control
							Experiment effect on virus in hold
	Spiked (addition of virus stock) medium		4.40	4.15 \le T \le 4.66		Significant	control
V2	5	TV2			= 1.07		(comparaison TV1 and TV2)
22							Experiment effect on virus in recovery
	Spiked (addition of virus stock) gauze			5.00 \le T \le 5.48		Not significant	gauze control
I	and recovery	TL1	5.24		= 0.24		(comparaison TV1 and TL1)
							Experiment effect on virus in recovery
	Spiked (addition of virus stock) gauze			3.16 ≤ T ≤ 3.73		Significant	gauze hold control
L2	and recovery	TL2	3.45		= 1.79		(comparaison TL1 and TL2)



Table 3 (continued): STUDY OF EVALUATION OF THE EFFICIENCY OF HAIER AIR CONDITIONER (24H) ON THE SARS-COV2 BY NO GLP VIRAL CLEARANCE STUDY (FIO)

TABLE 3.9: SPIKING EXPERIMENTS	Reduction factor evaluation (R)	Mean reduction factor m(R) and confidence interval $[m(R)] = Log_{10} (L1) - Log_{10} (LS)$	Positive control	Virus stock recovery gauze control	STUDY	Samples collected, recovery and diluted: Clearance factor	R > 4.08 (>99.991%)	R = 1.67 (97.862%)
	Virus load (L)	Mean load [m(L)=m(T) x Vt]		5.08 \le T \le 5.15			1.16 NA (No virus)	3.57 3.27 ≤ T ≤ 3.87
		əpoo		LL1			LS1	LS2
	Infectious Titer (T)	Mean titer [m(T)] and confidence interval		5.00 ≤ T ≤ 5.48			NA	3.27 ≤ T ≤ 3.87
		Mean		5.24			1.16	3.57
		эроэ		ITI			TS1	TS2
	Vol	(Vt) mL		1			1	1
	Sample	definition		Spiked (addition of virus stock)			Gauze on the evaporator, 10 centimeters away from the center of UVC module in horizontal position 24 hours	Gauze at 1 meter in front of the DIU for horizontal position, adjust its height to 1.5 meters high
		epoo		L1			S1	S2

Infectious titers are expressed as 50% tissue culture infectious dose per milliliter (Log10 TCID30/mL). Viral loads are expressed as 50% tissue culture infectious dose (Log10 TCID30).



CONCLUSIONS

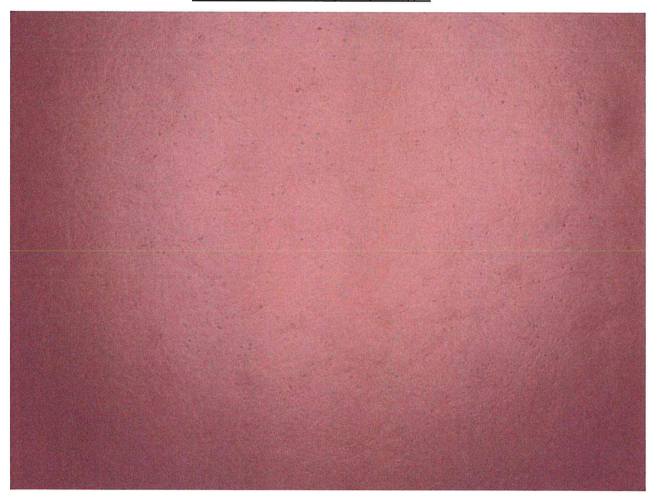
The resulted virus reduction factor (R) of the "Haier's air conditioner" for the time tested in this study is more 99.991% (>4.08 log) with 1.67 log due to the experiments duration.



Study number: 1198/02 March 29, 2022
Page 25 of 30

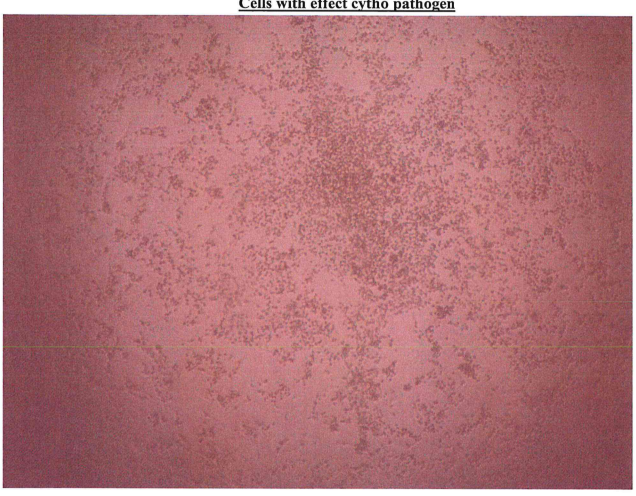
APPENDED

Cells without effect cytho pathogen



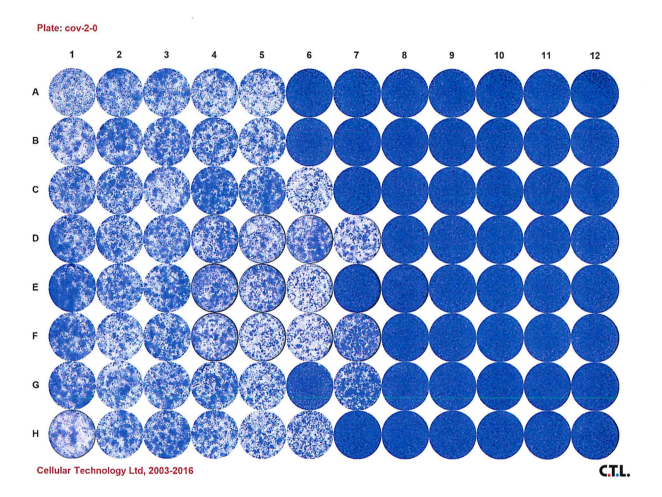








Plates of titration by TCID50/mL with Sars-Cov-2









Study number: 1198/02 March 29, 2022 Page 29 of 30







