

A) IDENTYFICATION OF THE SAMPLE	
Name of the product/Details about the product	DEZINFECTANT UNIVERSAL "BIO-DEZ"
1	Expiration date: 01.10.2024.
	Manufacturer (supplier): Ecochim-Grup SRL.
	Storing conditions: Dry, without sun, 5-25 Celsius degree.
	Conditions of use: Hygienic handrub, surface disinfection,
	medical instruments disinfection, surgical handrub
Active compound/s and its concentration/s	Ethyl alcohol 72-76%, CAS 64-17-5 and CE 200-578-6.
There compounds and its concentration's	Benzalkonium chloride 0.024- 0.029%, CAS 68424-85-1 and
	CE 270-325-2, Methylthionibium chloride 0.00024%, CAS 61-
	73-4 and 200-515-2
Concentrations requested for the assay	Pure (80%).
B) TEST METHOD	Ture (80%).
,	UNE-EN 13727: 2012 + A2: 2015. Chemical disinfectants and
Performed in accredited contracted partner laboratory,	
Scope of Accreditation Nr. 648/LE1286	antiseptics. Quantitative suspension test for the evaluation of
Report Registration No. D/21/B0645	bactericidal activity of chemical disinfectants for instruments
Quantitative evaluation assay of the bactericidal activity	used in Medicine. Test method and requirements (phase 2, step
under dirty conditions, in the medical area (phase 2, step	1). AENOR.
1) with product Desinfectant Universal "Bio-Dez", (UNE-	
EN 13727: 2012 + A2: 2015 Standard).	
Testing method	DESIN-1031-b //EN 13727: 2012 + A2: 2015
C) INFORMATION ABOUT SAMPLE RECEPTI	ON
Date of reception of order with test conditions	21.10.2021
Date of reception of the sample	25.10.2021
Aspect of the received product	Blue liquid in plastic package
D) METHOD OF ASSAY AND ITS VALIDATION	N (UNE-EN 13727: 2012+A2: 2015 Standard.)
Method used	Dilution-neutralization
Neutralizer	Tryptone 5 g/L, yeast extract 2.5 g/L, dextrose 10 g/L, sodium
	thioglycolate 1 g/L, sodium thiosulfate 1 g/L, sodium bisulphite
	2.5 g/L, soya lecithin 7 g/L, polysorbate-80 5 g/L, glycine 1 g/L,
	1-histidine 1 g/L and saponin 30 g/L.
E) EXPERIMENTAL CONDITIONS	1 instraine 1 g/L and supoint 30 g/L.
Assay period	2021/11/10 to 2021/11/14.
Solvent of the product used in the assay	Sterile distilled water.
Product concentrations for the assay	Pure (80%), 50%, 0.1%
Aspect of the dilutions of the product	Pure (80%) and 50% blue liquid;
Aspect of the dilutions of the product	0.1% transparent.
Contact time	60 seconds
	+20°C ± 1°C
Assay temperature	
Interfering substance	Bovine serum albumin 3 g/L and erythrocytes 3 mL/L.
Stability of the mixture (interfering substance and	Stable
product diluted in sterile distilled water)	200 400
Temperature of incubation	+36°C± 1°C
Identification of the strain used	- Pseudomonas aeruginosa (CECT 116 = ATCC 15442).
Identification of the strain used	- Pseudomonas aeruginosa (CECT 116 = ATCC 15442). - Staphylococcus aureus (CECT 239 = ATCC 6538).
Identification of the strain used	

Laboratory: Bucharest 041914, 8 Berceni Street.

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Enclosore no. 1 subcontracted tests

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Date: 08.12.2021

Authorized by: Mariana Ilinca, Manager of Microbiological laboratory



ENCLOSORE NO. 1 SUBCONTRACTED TESTS TO REPORT OF ANALYSIS NO 80249/21/ROBCH Results of the assay

Evaluation of bactericidal activity....... See tables 3, 6, 9 and 12.

Number of replicates per assay organism .. 1

Special remarks

- All controls and validation were between the basic limits.
- At least one concentration of the sample showed a log reduction lower than 5 log.
- At least one concentration of the sample showed a log reduction higher than 5 log.
- The highest concentration that can be assayed in the test (80%) is due to the mixtures to perform the assay.
- No precipitate formed during the test procedure (the test mixtures were homogeneous).

Conclusion

The product **Desinfectant Universal**"Bio-Dez", batch not indicated, when is pure (80%), shows bactericidal activity after 60 seconds at 20°C ± 1°C, under dirty conditions (bovine serum albumin 3 g/L and erythrocytes 3 mL/L), for the reference strains *Pseudomonas aeruginosa* (CECT 116 = ATCC 15442), *Staphylococcus aureus* (CECT 239 = ATCC 6538), *Enterococcus hirae* (CECT 4081 = ATCC 10541) and *Escherichia coli* K12 (CECT 433 = NCTC 10538), when tested according to UNE-EN 13727: 2012 + A2: 2015 Standard.

Note: The results obtained correspond to the sample received in the laboratory.

Quality Assurance Review:

The assay development and the results obtained have been supervised by the Director of the study.

The Quality Assurance Director has inspected the development of the assay, proving that has been realized following the proper procedure and using the adequate media, materials, and reagents, following the Good Laboratory Practices (GLPs) as well and the final report contains the primary data obtained.

Reference

 UNE-EN 13727: 2012 + A2: 2015. Chemical disinfectants and antiseptics. Quantitative suspension test for the evaluation of bactericidal activity of chemical disinfectants for instruments used in Medicine. Test method and requirements (phase 2, step 1). AENOR.

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Results of the assay (Bactericidal suspension) with Pseudomonas aeruginosa (CECT 116 = ATCC 15442).

Seeding: Pour plate; No. of plates: 1/mL.

Table 1.-Validation and controls

1	spension dation (i		l l		Contro	Control of neutralizer (B)			Validation of the method (<i>C</i>) Sample concentration: Pure (80%)		
Vc1	61	X=	Vc_1	53	X= 54	Vc_1	46	X= 48	Vc_1	42	<i>X</i> =
V_{C2}	56	58.5	Vc2	<i>C</i> 2 55			50		Vc2	37	39.5
30 ≤ x ($30 \le x \text{ of } Nv_0 \le 160$? $x \text{ of } A \text{ es } \ge 0.5 \text{ X de}$		$x \text{ of } B \text{ es } \ge 0.5 X \text{ de}$			x of C	c es ≥ 0.:	5X of			
				Nv_0 ?		Nv_0 , or 0.0005 Nv_B ?			Nvo?		
	Yes			Yes		Yes			Yes		
	spension						X = 60				
vali	dation (/	Vv_B)	Vc1: 57 Vc2: 63		$30 \le x \text{ de } Nv_B/1000 \le$						
						160?					
							Yes				

Table 2.-Suspension of the assay

	N	Vc ₁	Vc_2	
Suspension of assay (N and N_{θ})	10 ⁻⁶	217	234	$Xwm = 2.25 \times 10^8$, $\lg N = 8.35$ $N_0 = N/10$; $\lg N_0 = 7.35$
	10-7	21	22	$7.17 \le \lg N_0 \le 7.70$? Yes

Table 3.-Results of the activity assays with the sample

Concentrations of the sample (%)	Dilutions steps	Vc ₁	Vc2	$Lg Na = lg$ $(X \times 10 \text{ o}$ $Xwm \times 10)$	Lg R (Lg N ₀ =7.35)	Time of contact (seconds)	
Pure (80%)	Na ⁰	<14	<14	<2.15	>5.20	60	
Fute (0070)	Na -1	<14	<14	~2.13	-5.20	00	
50%	Na ⁰	<14	<14	<2.15	>5.20	60	
30%	Na -1	<14	<14	~2.13	~5.20	00	
0.1%	Na ⁰	>330	>330	>4.52	<2 92	60	
0.170	Na -1	>330	>330	~4.3 2	<2.83	60	

Explanations:

Vc = number per mL (one or two plates); Xwm = weighted mean of X.

 $X = \text{mean of } Vc_1 \text{ and } Vc_2 \text{ (duplicate of } 1 + 2);$ Logarithmic reduction R: $(\lg R = \lg N_0 - \lg Na)$.

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Results of the assay (Bactericidal suspension) with Staphylococcus aureus (CECT 239 = ATCC 6538).

Seeding: Pour plate; No. of plates: 1/mL.

Table 4.-Validation and controls

1	spension dation (i		Control of experimental conditions (A)		Contro	Control of neutralizer (B)			Validation of the method (C) Sample concentration: Pure (80%)		
Vc ₁	77	X= 74	Vc_1	68	<i>X</i> =	Vc_1	75	<i>X</i> =	Vc1	70	X= 66
V_{C2}	71		Vc2	69	68.5	Vc_2	82	78.5	Vc2	62	
30 ≤ x	$0 \le x \text{ of } Nv_0 \le 160$? $x \text{ of } A \text{ es } \ge 0.5 \text{ X de}$		$x \text{ of } B \text{ es } \ge 0.5 X \text{ de}$			x of C	c es ≥ 0.:	5X of			
				Nv_0 ?		Nv_0 , or 0.0005 Nv_B ?			Nv_0 ?		
	Yes			Yes		Yes			Yes		
Su	spension	ı of	Vc1: 79 Vc2: 86				X = 82.5				
vali	dation (7	Vv_B)			$30 \le x \text{ de } Nv_B/1000 \le$						
						160?					
							Yes				

Table 5.-Suspension of the assay

	N	Vc ₁	Vc_2	
Suspension of assay (N and N_{θ})	10 ⁻⁶	>330	>330	$Xwm = 3.40 \times 10^8$, $1g N = 8.53$ $N_0 = N/10$; $1g N_0 = 7.53$
	10-7	33	35	$7.17 \le \lg N_0 \le 7.70$? Yes

Table 6.-Results of the activity assays with the sample

Concentrations of the sample (%)	Dilutions steps	Vc1	Vc2	$Lg Na = lg$ $(X \times 10 \text{ o}$ $Xwm \times 10)$	Lg R (Lg No=7.53)	Time of contact (seconds)	
Pure (80%)	Na ⁰	<14	<14	<2.15	>5.38	60	
1 1120 (0070)	Na -1	<14	<14	2.22	2.20		
50%	Na ⁰	<14	<14	<2.15	>5.38	60	
3076	Na -1	<14	<14	~2.13	-5.56	00	
0.1%	Na ⁰	>330	>330	>4.52	<3.01	60	
0.1%	Na -1	>330	>330	~4.32	₹5.01	60	

Explanations:

Vc = number per mL (one or two plates); Xwm = weighted mean of X.

 $X = \text{mean of } Vc_1 \text{ and } Vc_2 \text{ (duplicate of } 1 + 2);$

Logarithmic reduction R: $(\lg R = \lg N_0 - \lg Na)$.

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Results of the assay (Bactericidal suspension) with *Enterococcus hirae* (CECT 4081 = ATCC 10541).

Seeding: Pour plate; No. of plates: 1/mL.

Table 7.-Validation and controls

1	spension dation (<i>I</i>		Control of experimental conditions (A)		Contro	Control of neutralizer (B)			Validation of the method (<i>C</i>) Sample concentration: Pure (80%)		
Vc1	41	X= 43	Vc1	44	<i>X</i> =	Vc_1	37	<i>X</i> =	Vc1	35	X= 36
V_{C2}	45		Vc_2	43	41.5	Vc_2	40	38.5	Vc2	37	
30 ≤ x	$30 \le x \text{ of } Nv_0 \le 160$? $x \text{ of } A \text{ es } \ge 0.5 \text{ X de}$		5 X de	$x \text{ of } B \text{ es } \ge 0,5 X \text{ de}$			x of C	C es ≥ 0.:	5X of		
				Nv_0 ?		Nv_0 , or 0.0005 Nv_B ?			Nv_0 ?		
	Yes			Yes		Yes			Yes		
Su	spension	of					X = 40.5				
vali	dation (7	Vv_B)	Vc1:	Vc1: 39 Vc2: 42		$30 \le x \text{ de } Nv_B/1000 \le$					
						160?					
							Yes				

Table 8.-Suspension of the assay

	N	Vc ₁	Vc_2	
Suspension of assay (N and N_{θ})	10-6	167	154	$Xwm = 1.61 \times 10^8$, $1g N = 8.20$ $N_0 = N/10$; $1g N_0 = 7.20$
	10-7	17	16	$7.17 \le \lg N_0 \le 7.70$? Yes

Table 9.-Results of the activity assays with the sample

Concentrations of the sample (%)	Dilutions steps	Vc1	Vcı	Lg Na = 1g (X x 10 o Xwm x 10)	Lg R (Lg No=7.20)	Time of contact (seconds)	
Pure (80%)	Na ⁰	<14	<14	<2.15	>5.05	60	
Fuie (00/0)	Na -1	<14	<14	~2.15	~5.05	00	
50%	Na ⁰	15	14	2.16	5.04	60	
30%	Na -1	<14	<14	2.10	3.04	60	
0.1%	Na ⁰	>330	>330	>4.52	<2.68	60	
0.1%	Na -1	>330	>330	~4.32	~2.08	60	

Explanations:

Vc = number per mL (one or two plates); Xwm = weighted mean of X.

 $X = \text{mean of } Vc_1 \text{ and } Vc_2 \text{ (duplicate of } 1 + 2);$

Logarithmic reduction R: $(\lg R = \lg N_0 - \lg Na)$.

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ENCLOSORE NO. 1 SUBCONTRACTED TESTS TO REPORT OF ANALYSIS NO 80249/21/ROBCH Results of the assay (Bactericidal suspension) with *Escherichia coli K12* (CECT 433 = NCTC 10538).

Seeding: Pour plate; No. of plates: 1/mL.

Table 10.-Validation and controls

1	spension dation (I		I I		Control of neutralizer (B)			Validation of the method (C) Sample concentration: Pure (80%)				
Vc_1	58	X= 60	Vc_1	44	X= 45	Vc_1	51	X= 50	Vc_1	43	X= 41	
V_{C2}	62		Vc_2	46		Vc2	49		Vc2	39		
30 ≤ x ($30 \le x \text{ of } Nv_0 \le 160$? $x \text{ of } A \text{ es } \ge 0$,		5 X de	$x \text{ of } B \text{ es } \ge 0.5 X \text{ de}$			x of C	c es ≥ 0.:	5X of			
				Nv_0 ?		Nv_0 , or 0.0005 Nv_B ?			Nv ₀ ?			
	Yes			Yes			Yes			Yes		
Su	spension	ı of					X = 55					
vali	dation (7	Vv_B)	Vc1: 54 Vc2: 56		$30 \le x \text{ de } Nv_B/1000 \le$							
							160?					
							Yes					

Table 11.-Suspension of the assay

	N	Vc ₁	Vc_2	
Suspension of assay (N and N_{θ})	10-6	241	259	$Xwm = 2.49 \times 10^8$, $\lg N = 8.40$ $N_0 = N/10$; $\lg N_0 = 7.40$
	10 ⁻⁷	22	25	$7.17 \le \lg N_0 \le 7.70$? Yes

Table 12.-Results of the activity assays with the sample

Concentrations of the sample (%)	Dilutions steps	Vc1	Vc2	Lg Na = 1g (X x 10 o Xwm x 10)	Lg R (Lg N ₀ =7.40)	Time of contact (seconds)	
Pure (80%)	Na ⁰	<14	<14	<2.15	>5.25	60	
Fure (8078)	Na -1	<14	<14	~2.13	-5.25	00	
50%	Na ⁰	<14	<14	<2.15	>5.25	60	
30%	Na -1	<14	<14	~2.13	~3.23	60	
0.1%	Na ⁰	>330	>330	>4.52	<2.88	60	
0.1%	Na -1	>330	>330	~4.32	~2.00	60	

Explanations:

Vc = number per mL (one or two plates); Xwm = weighted mean of X.

 $X = \text{mean of } Vc_1 \text{ and } Vc_2 \text{ (duplicate of } 1 + 2);$

Logarithmic reduction R: $(\lg R = \lg N_0 - \lg Na)$.

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A) IDENTYFICATION OF THE SA	MPLE:
Name of the product	DEZINFECTANT UNIVERSAL "BIO-DEZ"
	Sample quantity: 2 pcs x 1 L
	Production date: 26.01.2021
	Expiration date: 26.01.2024
	Sampling date: 22.02.2021
	Sample temperature: 15°C
	Reception hour: 15:00
	Responsible for sampling: Crestinov Alexandr
The active substance	Ethyl alcohol 72-76% CAS 64-17-5 CE 200-578-6
	Benzalkonium chloride 0,024-0,029% CAS 68424-85-1 CE 270-325-2
	Methylthioninium chloride 0,00024% CAS 61-73-4 and 200-515-2
B) TEST METHOD:	
Method	EN 1500:2013 Chemical disinfectants and antiseptics - Hygienic handrub - Test method and requirements (phase 2, step 2)
Neutralizer	Polysorbate 80 30 g/l, saponine 30g/l, histidine 1g/l, cysteine 1g/l
C) EXPERIMENTAL CONDITIONS	S:
Product test concentrations (%V/V)	100%
Test temperature	20°C
Contact time	3ml of the preparation for 60s
Incubation temperature	36±1 °C
Test-organism	E. coli K12 NCTC 10538

Date: 27.07.2021

Authorized by: Daria Depa, Senior Specialist Analyst, Cosmetics Microbiology Laboratory Approved by: Hanna Wachowska, Laboratory Director (Approved with qualified electronic signature)

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Table 1. PROCEDURE FOR REFERENCE HYGIENIC HANDRUB

PRODUCT: Standard 2-propanol 60% (V/V) TEST ORGANISM: E. coli K12 NCTC 10538

NUMBER IN CONTAMINATION FLUID: 2,4 x 10⁸ cfu/ml

VO	lunteer		r	number of cfu p	er plate from o	dilution 10x			
	Hand		prevalues			postva	lues		Reduction
Nr	left/right	x10 ⁻⁴	x10 ⁻⁵	log x	$x10^{0}$	x10 ⁻¹	x 10 ⁻²	log y	log z
	1	288	29		61	7	0		
1	r	247	22	6,42	33	3	0	1,65	4,77
	I	167	17		51	5	0		
2	r	291	28	5,81	36	4	0	1,63	4,18
	I	175	11		42	5	0		
3	r	275	25	6,33	29	2	0	1,54	4,79
	I	220	21		30	3	0		
4	r	192	19	6,31	68	6	0	1,65	4,66
	I	164	15		37	3	0		
5	r	301	33	6,35	52	5	0	1,64	4,71
	I	200	20		23	2	0		
6	r	198	18	6,30	37	4	0	1,46	4,83
	I	287	22		60	6	0		
7	r	288	29	6,45	42	5	0	1,70	4,75
_	I	298	28		31	4	0		
8	r	213	21	6,40	58	5	0	1,63	4,77
_	I	283	23		34	3	0		
9	r	311	33	5,96	51	5	0	1,62	4,34
4.0	I	313	32	0.45	53	6	0	4.0=	4.00
10	r	251	25	6,45	36	4	0	1,65	4,80
44		175	18	0.05	54 47	5	0	4.00	4.00
11	r	295	22	6,35	72	3	0	1,69	4,66
40	-	183	19	F 74		7	0	4 74	4.00
12	r	171 206	17 22	5,74	36 29	4 2	0	1,71	4,03
13	-	317	33	6,41	49 49	5	0	1 57	4 9 4
13	1	295	28	0,41	55	6	0	1,57	4,84
14	r	279	25	6,45	64	7	0	1,78	4,68
14	1	248	22	0,45	72	7	0	1,70	4,00
15	r	256	26	6,40	66	6	0	1,84	4,56
10	i I	301	31	0,40	46	5	0	1,04	7,50
16	r r	261	26	6,45	27	3	0	1,55	4,90
- 10	i	259	24	0,10	41	4	0	1,00	1,00
17	r	271	28	6,42	22	1	0	1,47	4,96
	i	259	22	5,	61	6	0	.,	.,00
18	lr	288	23	6,43	33	3	0	1,65	4,78
	ı	223	21	.,	35	4	0	,	, -
19	r	205	20	6,33	45	5	0	1,60	4,72
	ı	297	28	· ·	54	6	0	,	,
20	r	257	24	5,90	28	3	0	1,59	4,31
X _{śr}		•	•	6,28				1,63	4,65
S	╡			0,23				0,09	

log x-logarithm of the average value of the initial left and right hand

log y-logarithm of the average value of the final left and right hand

log z-logarithm reduction

x śr- overall average of log x, log y, log z

Date: 27.07.2021

Authorized by: Daria Depa, Senior Specialist Analyst, Cosmetics Microbiology Laboratory Approved by: Hanna Wachowska, Laboratory Director (Approved with qualified electronic signature)

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Table 2. HYGIENIC HANDRUB PROCEDURE WITH THE PRODUCT

PRODUCT P 136987/21/JSHR

TEST ORGANISM: E. coli K12 NCTC 10538

NUMBER IN CONTAMINATION FLUID: 2,4 x 108 cfu/ml

VC	olunteer		n	umber of cfu pe	er plate from o	dilution 10x			
	Hand		prevalues			postva			Reduction
Nr	left/right	x10 ⁻⁴	x10 ⁻⁵	log x	$x10^{0}$	x10 ⁻¹	x 10 ⁻²	log y	log z
	I	132	14		103	11	1		
1	r	224	21	6,24	92	9	0	1,98	4,26
	I	>330	125		89	7	0		
2	r	304	31	6,27	78	4	0	1,68	4,59
	l I	144	15		97	9	0		
3	r	132	11	6,14	78	5	0	1,93	4,21
	I	328	34		87	8	0		
4	r	>330	85	6,20	99	9	0	1,89	4,32
	I	164	11		116	11	2		
5	r	132	12	6,16	99	8	0	2,03	4,13
	l l	>330	121		61	3	0		
6	r	320	32	6,27	83	9	0	1,67	4,60
	l l	328	33		61	4	0		
7	r	288	29	6,49	71	7	0	1,81	4,68
	Į.	>330	58		91	9	0		
8	r	>330	22	5,51	72	6	0	1,82	3,69
•	l I	336	36		79	8	0	4.00	
9	r	>330	21	5,90	106	12	2	1,96	3,94
40		296	28	0.00	74	7	0	4.00	4.40
10	ır	>330	41	6,02	85 93	9	0	1,90	4,12
4.4		228 104	21	6.40	93 80	8 5		4.00	4.00
11	l I	>330	11 48	6,19	107		0	1,93	4,26
12		200	20	5,97	94	11 9	1 0	1,98	4,00
12	1	248	25	5,97	112	14	2	1,90	4,00
13	'_	212	22	6,36	113	11	1	2,06	4,31
13	1	>330	48	0,30	89	8	0	2,00	4,31
14	l'r	255	22	6,02	91	9	0	1,95	4,07
17	i i	278	28	0,02	99	7	0	1,55	4,07
15	r r	169	17	6,34	67	6	0	1,77	4,57
	i	178	11	5,5 .	104	11	1	.,	.,
16	r	255	25	6,32	69	7	o O	1,93	4,39
	i	274	28	5,52	79	8	0	.,,,,	.,
17	r	231	24	6,40	107	12	2	1,97	4,44
	I	225	22	-, -	92	9	0	,	,
18	r	183	19	6,31	66	7	0	1,89	4,42
	I	199	17		53	5	0	•	
19	r	252	23	6,35	89	8	0	1,83	4,51
	I	266	22		97	9	0		
20	r	231	21	6,39	68	7	0	1,91	4,48
X _{śr}				6,19				1,89	4,30
S				0,22				0,10	0,25

log x-logarithm of the average value of the initial left and right hand log y-logarithm of the average value of the final left and right hand log z-logarithm reduction

x śr- overall average of log x, log y, log z

Date: 27.07.2021

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Table 3. LIST OF COMPUTED IG VALUES AND IG REDUCTIONS

volu	volunteer R 2-propanol 60% (V/V)				Р			
Nr		log x	log y	log z	log x	log y	log z	
1	R-P	6,42	1,65	4,77	6,24	1,99	4,25	
2	R-P	5,81	1,63	4,18	6,27	1,91	4,36	
3	R-P	6,33	1,54	4,79	6,14	1,93	4,21	
4	R-P	6,31	1,65	4,66	6,20	1,96	4,24	
5	R-P	6,35	1,64	4,71	6,16	2,03	4,13	
6	P-R	6,30	1,46	4,83	6,27	1,84	4,43	
7	P-R	6,45	1,70	4,75	6,49	1,81	4,68	
8	P-R	6,40	1,63	4,77	5,51	1,90	3,61	
9	P-R	5,96	1,62	4,34	5,90	1,96	3,94	
10	P-R	6,45	1,65	4,80	6,02	1,90	4,12	
11	R-P	6,35	1,69	4,66	6,19	1,93	4,26	
12	R-P	5,74	1,71	4,03	5,97	2,00	3,97	
13	R-P	6,41	1,57	4,84	6,36	2,06	4,31	
14	R-P	6,45	1,78	4,68	6,02	1,95	4,07	
15	R-P	6,40	1,84	4,56	6,34	1,90	4,43	
16	P-R	6,45	1,55	4,90	6,32	1,93	4,39	
17	P-R	6,42	1,47	4,96	6,40	1,97	4,44	
18	P-R	6,43	1,65	4,78	6,31	1,89	4,42	
19	P-R	6,33	1,60	4,72	6,35	1,83	4,51	
20	P-R	5,90	1,59	4,31	6,39	1,91	4,48	
X ₂₀		6,28	1,63	4,65	6,19	1,93	4,26	
X10(R-P)		6,26	1,67	4,59	6,19	1,97	4,22	
X10 (P-R)		6,31	1,59	4,72	6,20	1,90	4,30	

Criteria:

Rs(R-P) = 4,59-4,22=0,37Rs (P-R)= 4,72-4,30=0,42

Abs= 0,37-0,42=-0,05<2

logx(R) = 6,28 > 5logx(P) = 6,19 > 5

logz(P), logz(R) > 3

Validation conditions of neutralizer and methods have been satisfied

Date: 27.07.2021

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Table 4. COMPUTATION OF INDIVIDUAL DIFFERENCES OF Ig R-P

volunteer	log	RF	difference	difference	
	R	Р	R-P	high to low	Range +/-
1	4,77	4,25	0,52	1,16	1
2	4,18	4,36	-0,18	0,68	2
3	4,79	4,21	0,59	0,61	3
4	4,66	4,24	0,42	0,59	4
5	4,71	4,13	0,58	0,58	5
6	4,83	4,43	0,40	0,53	6
7	4,75	4,68	0,07	0,52	7
8	4,77	3,61	1,16	0,52	8
9	4,34	3,94	0,40	0,51	9
10	4,80	4,12	0,68	0,42	10
11	4,66	4,26	0,40	0,40	11
12	4,03	3,97	0,06	0,40	12
13	4,84	4,31	0,53	0,40	13
14	4,68	4,07	0,61	0,36	14
15	4,56	4,43	0,13	0,21	15
16	4,90	4,39	0,51	0,13	16
17	4,96	4,44	0,52	0,07	17
18	4,78	4,42	0,36	0,06	18
19	4,72	4,51	0,21	-0,17	-19
20	4,31	4,48	-0,17	-0,18	-20
	sum of ranks	(+): 171			
	sum of ranks	(-): 39		<u> </u>	

SORTING OF INDIVIDUAL DIFFERENCES AND COMPUTATION FOR HODGES-LEHMANN 97,5% UPPER CONFIDENCE LIMITS FOR THE DIFFERENCE IN IG BETWEEN R-P

_		1,16	0,68	0,61	0,59	0,58	0,53	0,52	0,52	0,51
1	1,16	1,16								
2	0,68	0,92	0,68							
3	0,61	0,89	0,65	0,61						
4	0,59	0,87	0,63	0,60	0,59					
5	0,58	0,87	0,63	0,59	0,58	0,58				
6	0,53	0,85	0,61	0,57	0,56	0,55	0,53			
7	0,52	0,84	0,60	0,57	0,56	0,55	0,53	-0,52		
8	0,52	0,84	0,60	0,56	0,55	0,55	0,53	-0,52	-0,52	
9	0,51	0,83	0,59	0,56	0,55	0,54	0,52	-0,52	-0,51	-0,51
10	0,42	0,79	0,55	0,52	0,50	0,50	0,48	-0,47	-0,47	-0,21
11	0,40	0,78	0,54	0,51	0,49	0,49	0,47	-0,46	-0,46	-0,20
12	0,40	0,78	0,54	0,51	0,49	0,49	0,47	-0,46	-0,46	
13	0,40	0,78	0,54	0,50	0,49	0,49	0,47	-0,46		
14	0,36	0,76	0,52	0,49	0,47	0,47	0,45			
15	0,21	0,69	0,45	0,41	0,40	0,39				
16	0,13	0,65	0,41	0,37	0,36					
17	0,07	0,62	0,38	0,34						
18	0,06	0,61	0,37							
19	-0,17	0,50								
20	-0,18									

Date: 27.07.2021

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Table 6. WILCOXON'S TMATCHED PAIRS SIGNED-RANKS TEST: CRITICAL VALUES LESS WITH RANG SUM (+) OR (-) AT DIFFERENT LEVELS OF SIGNIFICANCE

n	one-sided level of significance								
	0,05	0,025	0,01						
18	47	40	32						
19	53	46	27						
20	60	52	43						
21	68	59	49						
22	75	66	56						

For the designated level of significance 0,025 for n=20 the value read from the table 6 is 52.

Hence c = 52+1 = 53.

For the distribution of 53 Table 5 assigns a value of 0,55 which is less than the agreed inferiority margin of 0,6.

Therefore, the hypothesis of inferiority of PP compared to the reference RP is rejected.

The test preparation (PP) is non-inferior to RP.

Date: 27.07.2021

Authorized by: Daria Depa, Senior Specialist Analyst, Cosmetics Microbiology Laboratory

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A) IDENTYFICATION OF THE SAMPLE	
Name of the product/Details about the product	DEZINFECTANT UNIVERSAL "BIO-DEZ"
	Expiration date: 01.10.2024.
	Manufacturer (supplier): Ecochim-Grup SRL.
	Storing conditions: Dry, without sun, 5-25 Celsius degree.
	Conditions of use: Hygienic handrub, surface disinfection,
	medical instruments disinfection, surgical handrub
Active compound/s and its concentration/s	Ethyl alcohol 72-76%, CAS 64-17-5 and CE 200-578-6. Benzalkonium chloride 0.024-0.029%, CAS 68424-85-1 and CE
	270-325-2, Methylthionibium chloride 0.00024%, CAS 61-73-4 and 200-515-2
Concentrations requested for the assay	Pure (80%).
B) TEST METHOD	
Performed in accredited contracted partner	UNE-EN 13624:2014 Chemical disinfectants and antiseptics.
laboratory, Scope of Accreditation Nr. 648/LE1286	Quantitative suspension test for the evaluation of the fungicidal
Report Registration No. D/21/B0644	or yeasticidal activity in the medical area. Test method and
Quantitative evaluation assay of yeasticidal activity	requirements (phase 2, step 1). AENOR.
under dirty conditions, in the medical area (phase 2,	
step 1), with product Desinfectant Universal"Bio-	
Dez", (UNE-EN 13624: 2014 Standard).	
Testing method	Procedure DESIN-1058-b // EN 13624:2014
C) INFORMATION ABOUT SAMPLE REC	EPTION
Date of reception of order with test conditions	21.10.2021
Date of reception of the sample	25.10.2021
Aspect of the received product	Blue liquid in plastic package
D) METHOD OF ASSAY AND ITS VALIDA	TION (UNE-EN 13624: 2014 Standard)
Method used	Dilution-neutralization
Neutralizer	Tryptone 5 g/L, yeast extract 2.5 g/L, dextrose 10 g/L, sodium
	thioglycolate 1 g/L, sodium thiosulfate 1 g/L, sodium bisulphite
	2.5 g/L, soy lecithin 7 g/L, polysorbate-80 5 g/L, glycine 1 g/L,
	1-histidine 1 g/L and Saponin 30 g/L.
E) EXPERIMENTAL CONDITIONS	
Assay period	2021/11/08 to 2021/11/14.
Solvent of the product used in the assay	Sterile distilled water.
Product concentrations for the assay	Pure (80%), 50%, 0.1%
Aspect of the dilutions of the product	Pure (80%) and 50% blue liquid;
	0.1% transparent.
Contact time	60 seconds
Assay temperature	$+20^{\circ}\text{C} \pm 1^{\circ}\text{C}$
Interfering substance	Bovine serum albumin 3 g/L and erythrocytes 3 mL/L.
Stability of the mixture (interfering substance and product diluted in sterile distilled water)	Stable
Temperature of incubation	+30°C± 1°C
Identification of the strain used	Candida albicans CECT-1394 (ATCC 10231)

Laboratory: Bucharest 041914, 8 Berceni Street.

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Enclosore no. 1 subcontracted tests

PGL 09 F 04 Ed. 1 Rev. 0

Page 1 of 3

Date: 08.12.2021

Authorized by: Mariana Ilinca, Manager of Microbiological laboratory

^{*}Test method accredited # Test performed by subcontractor. Ø Non accredited methods.



ENCLOSORE NO. 1 SUBCONTRACTED TESTS TO REPORT OF ANALYSIS NO 80248/21/ROBCH Results of the assay

•	Assay of validation	See tables 1 and 2
•	Evaluation of yeasticidal activity	See table 3.
•	Number of replicates per assay organism	

Special remarks

- All controls and validation were between the basic limits.
- At least one concentration of the sample showed a log reduction lower than 4 log.
- At least one concentration of the sample showed a log reduction higher than 4 log.
- The highest concentration that can be assayed in the test (80%) is due to the mixtures to perform the assay.

1

No precipitate formed during the test procedure (the test mixtures were homogeneous).

Conclusion

The product **Desinfectant Universal"Bio-Dez**", batch not indicated, when is pure (80%), shows yeasticidal activity after 60 seconds at $20^{\circ}\text{C} \pm 1^{\circ}\text{C}$, under dirty conditions (bovine serum albumin 3 g/L and erythrocytes 3 mL/L), for the reference strain *Candida albicans* (CECT 1394 = ATCC 10231), when tested as required by the **UNE-EN 13624: 2014** Standard.

Note: The results obtained correspond to the sample received in the laboratory.

Quality Assurance Review:

The assay development and the results obtained have been supervised by the Director of the study.

The Quality Assurance Director has inspected the development of the assay, proving that has been realized following the proper procedure and using the adequate media, materials, and reagents, following the Good Laboratory Practices (GLPs) as well and the final report contains the primary data obtained.

Reference

UNE-EN 13624: 2014. Chemical disinfectants and antiseptics. Quantitative suspension test
for the evaluation of the fungicidal or yeasticidal activity in the medical area. Test method
and requirements (phase 2, step 1). AENOR.

Laboratory: Bucharest 041914, 8 Berceni Street.

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ENCLOSORE NO. 1 SUBCONTRACTED TESTS TO REPORT OF ANALYSIS NO 80248/21/ROBCH Results of the assay with *Candida albicans* (CECT 1394 = ATCC 10231).

Seeding: Pour plates. No. of plates: 1 /mL.

Table 1.-Validation and controls.

Suspension of validation (Nv ₀)		Control of experimental conditions (A)		Contro	Control of neutralizer (B)			Validation of the method (<i>C</i>) Sample concentration:			
									P	ure (80%	6)
Vc_1	86	X= 90	Vc_1	72	X= 74	Vc_1	75	X= 73	Vc_1	66	<i>X</i> =
V_{C2}	94]	Vc2	76]	Vc2	71]	Vc2	61	63.5
30 ≤ x ($30 \le x \text{ of } Nv_0 \le 160$?		$x \text{ of } A \text{ es } \ge 0.5 X \text{ de}$		x of E	es ≥ 0,5	5 X de	x of C	$x \text{ of } C \text{ es } \ge 0.5 X \text{ of }$		
			Nv_0 ?		Nv_0 , or 0.0005 Nv_B ?		Nv_0 ?				
	Yes			Yes		Yes			Yes		
Su	Suspension of		X=78								
vali	validation (NvB) Vc1: 79 Vc2: 77		$30 \le x \text{ de } Nv_B/1000 \le$								
						160?					
							Yes				

Table 2. -Suspension of the assay.

	N	Vc_1	Vc_2	$Xwm = 3.35 \times 10^7$
Suspension of assay	10-5	>330	>330	$\lg N = 7.53$
(N and N ₀)	10	- 550		$N_0 = N/10$
	10 ⁻⁶	32		$lg N_0 = 6.53$ 6.17 $\leq lg N_0 \leq 6.70$? Yes

Table 3.-Results of the activity assays with the sample.

Concentrations of the sample (%)	Dilutions steps	Vc ₁	Vc2	Lg Na = lg (X x 10 or Xmw x 10)	$LgR $ (lg N_{θ} = 6.53)	Time of contact (seconds)	
Pure (80%)	Na^0	<14	<14	<2.15	>4.38	60	
Pute (80%)	Na ⁻¹	<14	<14 <14 <2.1		Z4.30	00	
50%	Na^0	<14	<14	<2.15	>4.38	60	
30%	Na ⁻¹	<14	<14	~2.13	∠4.30	00	
0.19/	Na^0	>330	>330	>4.52	<2.01	60	
0.1%	Na ⁻¹	>330	>330	~4.3 2	~2.01	60	

Explanations:

Vc = number per mL (one or two plates); Xwm = ponderated mean of X.

 $X = \text{mean of Vc}_1$ and Vc_2 (duplicate of 1 + 2); R (reduction): ($\lg R = \lg N_0 - \lg N_a$).

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Enclosore no. 1 subcontracted tests

^{*}Test method accredited # Test performed by subcontractor. Ø Non accredited methods.



A) IDENTYFICATION OF THE SAMPLE	
Name of the product/Details about the product	DEZINFECTANT UNIVERSAL "BIO-DEZ"
	Sample quantity: 1 pcs x 1 L
	Production date: 26.01.2021
	Expiration date: 26.01.2024
	Manufacturer(supplier): ECOCHIM-GRUP
	Dry, without sun, 5-25°C
Active(s) Substance(s) and its concentration(s)	Ethyl alcohol 72-76%, CAS 64-17-5 and CE 200-578-6 Benzalkonium chloride 0,024-0,029%, CAS 68424-85-1 and CE 270-325-2; Methylthioninium chloride 0,00024%, CAS 61-73-4 and 200.
Concentration ordered for the assay	3%
B) TEST METHOD	
Performed in accredited subcontracted partner laboratory: Scope	NF EN 14476:2013+A2:2019 Standard. Virucidal
of Accreditation Nr. 648/LE1286	quantitative suspension test for chemical disinfectants and
Report D/21/V0204- Modified report	antiseptics used in human medicine. Test method and
Virucidal test with the sample DEZINFECTANT	requirements (Phase 2/Step1).AFNOR
UNIVERSAL "BIO-DEZ" against Poliovirus type 1,	
Adenovirus type 5 and Murine Norovirus (NF EN	
14476:2013+A2:2019 Standard)	
Testing method	Procedure DESIN-1078 (NF EN 14476:2013+ A2:2019
	Standard)
C) INFORMATION ABOUT SAMPLE RECEPTION	
Date of reception of the sample	22.03.2021
Date of reception of order with test conditions	22.03.2021
Aspect of the received product	Blue transparent liquid in a plastic container
D) EXPERIMENTAL CONDITIONS	
Assay period	From 22.03.2021 to 01.04.2021
Assay temperature	37°C ± 1°C
Titration method	TCID50 (Tissue Culture Infective Dose 50%)
Product concentrations for the assay	80%, 3% and 0.03%
Contact time	60 seconds
Contact temperature	$20^{\circ}\text{C} \pm 1^{\circ}\text{C}$
Procedure to stop product cytotoxicity	Molecular sieving (< 4 columns)
Procedure to stop product activity	Cooling with ice
Solvent of the product used in the assay	Hard water
Aspect of the dilutions of the product	Transparent
Stability of the mixture (interfering substance and product	
diluted in sterile hard water/distilled water)	Stable
Interfering substance	Clean conditions in the presence of bovine serum albumin
<u> </u>	0.3 g/L.
Identification of the origin of viral stains and number of passes	Poliovirus type 1 aliquot: 2021/01/07 passage 2;
	Adenovirus type 5 aliquot: 2021/01/14 passage 2;
	Murine Norovirus aliquot: 2021/02/11 passage 2;
Cell lines (name, origin, number of passes)	Vero ref: FTVE, working aliquot 12 passages 13 and 17;
	Raw 264.7, Public Health England, working aliquot 12,
	passages 12, 16 and 18;

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Enclosore no. 1 subcontracted tests

PGL 09 F 04 Ed. 1 Rev. 0

Page 1 of 13

Date: 13.09.2021

Authorized by: Mariana Ilinca, Manager of Microbiological laboratory



Note: Modified report due to the client requests to add the 80% concentration results in the conclusions section.

Validation of assay results

Poliovirus type 1 (ATCC VR-192)

Titre of the viral suspension for the virus control (at the requested test time): • Clean conditions
Maximum level of virus inactivation detectable (difference between the titre of the viral suspension and the cytotoxicity level): • Clean conditions
Adenovirus type 5 (ATCC VR-5)
Titre of the viral suspension for the virus control (at the requested test time): • Clean conditions
Maximum level of virus inactivation detectable (difference between the titre of the viral suspension and the cytotoxicity level): • Clean conditionslog 10 ^{-5.83}
Murine Norovirus (strain S99 Berlin)
Titre of the viral suspension for the virus control (at the requested test time): • Clean conditions
Maximum level of virus inactivation detectable (difference between the titre of the viral suspension and the cytotoxicity level): • Clean conditions
Reference test (formaldehyde 1.4%)
Cytotoxicity level of formaldehyde 0.7% log10 ^{-0.50}
Viral quantification in the reference test (formaldehyde) after 60 minutes and with Poliovirus type 1

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Enclosore no. 1 subcontracted tests



ENCLOSORE NO. 1 SUBCONTRACTED TESTS TO REPORT OF ANALYSIS NO 17899/21/ROBCH/Z1 Confidence interval

Titre of virus with 95% confidence interval with Poliovirus type 1 (at the requested test time): O Clean conditions
Titre of virus with 95% confidence interval with Adenovirus type 5 (at the requested test time): \circ Clean conditions
Titre of virus with 95% confidence interval with Murine Norovirus (at the requested test time): $ \qquad $
Reduction with the confidence interval of 95 %
Sensitivity of cells to virus
 Viral quantification of Poliovirus type 1 with cells not treated by the test solution with the test sample
reduction of the title of the virus $< 1 \log_{10}$.
Control of the effectivity of the disinfectant detection activity
 Viral quantification of Poliovirus type 1 after 30 minutes on bath ice without exposing the virus to the test sample
• Viral quantification of Adenovirus type 5 after 30 minutes on bath ice without

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- Viral quantification of Murine Norovirus after 30 minutes on bath ice without exposing the virus to the test samplelog10^{-6,32}
- Viral quantification of Murine Norovirus exposing the virus to "the test sample and incubated 30 minutes on ice bathlog10^{-5,99}

Note: The difference between decimal logarithm of titre without exposing the virus to the sample and of the test suspension should be ≤ 0.5

Special remarks

- The sample is tested at 80%; 3% and 0.03%. The highest concentration that can be tested in the test is 80%, because of the mixtures made during the test.
- All controls and validation were between the basic limits.
- One concentration at least showed a log reduction less than 4 log.
- One concentration at least showed a log reduction equal or higher than 4 log.

Assay results

Description of the results under the requested test conditions:

Virus of assay	Test concentrations, reduction obtained with the confidence interval of 95 % and virucidal activity								
	80%	3%	0.03%						
Poliovirus	5.33 ± 0.49 TCID ₅₀	0.83 ± 0.54 TCID ₅₀	0.17 ± 0.55 TCID ₅₀						
type 1	Shows	Does not show	Does not show						
Adenovirus	≥ 5.83 ± 0.40 TCID ₅₀	1.51 ± 0.52 TCID ₅₀	0.17 ± 0.56 TCID ₅₀						
type 5	Shows	Does not show	Does not show						
Murine	≥ 5.66 ± 0.34 TCID ₅₀	2.17 ± 0.48 TCID ₅₀	0.09 ± 0.53 TCID ₅₀						
Norovirus	Shows	Does not show	Does not show						

Virucidal activity exists when the titre of virus shows a reduction ≥ 4 log₄ TCID₅₀: Tissue Culture Infectious Dose 50%.

Tables of results and graphics

See tables 1 to 6 and figure 1 to 3.

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The disinfectant sample "DEZINFECTANT UNIVERSAL, BIO-DEZ", batch not indicated, under clean conditions (bovine serum albumin 0.3 g/L), at 3% concentration, requested by the client, and during 60 seconds of contact time and 20°C of temperature, does not show virucidal activity against the three mandatory viruses (Poliovirus type 1, Adenovirus type 5 and Murine Norovirus) when the activity is assayed according with the NF EN 14476: 2013 + A2: 2019 Standard. However, the disinfectant sample "DEZINFECTANT UNIVERSAL, BIO-DEZ", batch not indicated, under clean conditions (bovine serum albumin 0.3 g/L), at 80% concentration and during 60 seconds of contact time and 20°C of temperature, shows virucidal activity against the three mandatory viruses (Poliovirus type 1, Adenovirus type 5 and Murine Norovirus) when the activity is assayed according with the NF EN 14476: 2013 + A2: 2019 Standard.

Therefore, the disinfectant tested, does not show general virucidal activity, diluted at 3% and it shows general virucidal activity, diluted at 80% when the activity is evaluated according to the NF EN 14476: 2013 + A2: 2019 Standard.

Note 1: The results obtained correspond to the sample received in this laboratory.

Note 2: The information that depend on the information received from the client and are not facilitated by the same one, shown as "not indicated".

Quality Assurance Review:

The assay development and the results obtained have been supervised by the Director of the study.

The Quality Assurance Director has inspected the development of the assay, proving that has been realized following the proper procedure and using the adequate media, materials and reagents, following as well the Good Laboratory Practices (GLPs) and the final report contains the primary data obtained.

Reference:

 NF EN 14476: 2013 + A2: 2019 Standard. Virucidal quantitative suspension test for chemical disinfectants and antiseptics used in human medicine. Test method and requirements (Phase 2/Step 1). AFNOR.

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Table 1. Results of activity of the sample test sample with Poliovirus type 1 (ATCC VR-192) under test conditions requested by the client.

Assay	Concen- tration*	Interfering substance	Cytoto- xicity level	0 min		TCID ₅₀ er 30 min	60 min	Reduction with the confidence interval of 95 %
	80%		0.5	-	1.91	-	-	5.33 ± 0.49
Test sample	3%	0.3 g/L BSA	0.5	-	6.41	-	-	0.83 ± 0.54
	0.03%		0.5	-	7.07	-	-	0.17 ± 0.55
Virus control	NA	0.3 g/L BSA	NA	7.32	7.24	-	-	NA
Formaldehyde	0.7% (w:v)	NA	0.5	NR	NR	5.16	3.32	NA
Virus control Formaldehyde	0.7% (w:v)	NA	0.5	7.07	NR	NR	6.99	NA

Control of sensitivity of cells to virus (difference between decimal logarithm of titre using

Control of the effectiveness of the disinfectant detection activity (difference between decimal logarithm of titre without exposing the virus to the sample and of the test suspension).....log10^{-0.41}

NA: not applicable; NR: not realized

Times recommended by Standard for surfaces: maximum 5 or 60 minutes.

Times recommended by Standard for instruments: maximum 60 minutes.

Times recommended by Standard for Hygienic treatment of hands by friction and hygienic handwashing: between 30 or 120 seconds.

PBS: phosphate buffered saline; BSA: bovine serum albumin.

Virucidal activity exists when the titre of virus shows a reduction ≥4 log.

*: see Special remarks to understand the values of these concentrations.

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Table 2. Results of the activity of the test sample, with Poliovirus type 1 (ATCC VR-192)

(Assay of titration with 12 wells), under test conditions requested by the client.

Assay	Concen-	Interfering	Time of contact			Г	ilutions	(log10)	1,0		
Assay	tration *	substance	(sec/min)	1	2	3	4	5	6	7	8
	tration	Substance	(occinin)	4444	2000	0000	0000	0000	0000	0000	_
	80%		60 sec	4444	3000	0001	0000	0000	0000	0000	N
				4444	0320	0000	0000	0000	0000	0000	
				4444	4444	4444	4444	4444	0230	1000	
Test sample 39	3%	0.3 g/L BSA	60 sec	4444	4444	4444	4444	4444	3424	0201	N
				4444	4444	4444	4444	4444	0403	0000	
				4444	4444	4444	4444	4444	3042	0201	00
	0.03%		60 sec	4444	4444	4444	4444	4444	3343	1302	00
				4444	4444	4444	4444	4444	2444	0020	11
0.000	908/	0.2 - 7 DCA		0000	0000	0000	0000	0000	0000	0000	3.1
Cytotoxicity	80%	0.3 g/L BSA	NA	0000	0000	0000	0000	0000	0000	0000	N
				4444	4444	4444	4444	4444	4444	3023	00
			0	4444	4444	4444	4444	4444	4444	4040	01
				4444	4444	4444	4444	4444	4444	3320	00
Virus control	NA	0.3 g/L BSA		4444	4444	4444	4444	4444	4444	3000	00
	l NA		60 sec	4444	4444	4444	4444	4444	4444	3402	00
		00 300	4444	4444	4444	4444	4444	4444	3023	00	
Formaldehyde 0.7% (w:v)			4444	4444	4444	4444	2300	0002	0000		
			30 min	4444	4444	4444	4444	0302	0010	0000	N
	0.7%		50 11111	4444	4444	4444	4444	2000	0100	0000	
		NA		4444	4444	0320	0001	0000	0000	0000	
	(**.*)		60 min	4444	4444	2301	0010	0000	0000	0000	N
				4444	4444	0332	0000	0000	0000	0000	
Control of				0000	0000	0000	0000	0000	0000	0000	
formaldehyde	0.7%	0.3 g/L BSA	NA	0000	0000	0000	0000	0000	0000	0000	N
cytotoxicity	(w:v)	0.5 g/L 155A	1421	0000	0000	0000	0000	0000	0000	0000	٠.
cytotoxicity	(w.v)			4444	4444	4444	4444	4444	4444	0300	00
			0	4444	4444	4444	4444	4444	4444	0204	00
Virus control				4444	4444	4444	4444	4444	4444	2030	00
formaldehyde	0.7%	NA		4444	4444	4444	4444	4444	3240	2301	00
Tormandenyde	(w:v)	142	60 min	4444	4444	4444	4444	4444	2444	0100	10
				4444	4444	4444	4444	4444	3403	3202	00
			C-11	CCCC	CCCC	CCCC	CCCC	CCCC	CCCC	0C0C	CO
			Cells not	CCCC	CCCC	CCCC	cccc	cccc	0CCC	CC0C	00
Sensitivity control of cells to virus			treated	CCCC	CCCC	CCCC	CCCC	CCCC	CCCC	0CCC	00
	NA.	NA	0.11	CCCC	CCCC	CCCC	cccc	CCCC	C0CC	0C00	00
			Cells	CCCC	CCCC	CCCC	CCCC	CCCC	CCC0	00C0	00
			treated	CCCC	CCCC	CCCC	cccc	cccc	CCC0	000C	00
			*****	CCCC	CCCC	CCCC	CCCC	CCCC	CCCC	C00C	00
Effectiveness			Without	CCCC	CCCC	CCCC	CCCC	CCCC	CCCC	CC0C	00
control of the			sample	CCCC	CCCC	CCCC	CCCC	CCCC	CCCC	0000	CO
	NA (0.3 g/L BSA			2446		2444				
	NA.	0.3 g/L BSA		CCCC	cccc	CCCC	CCCC	cccc	COCC	0000	0.0
disinfectant detection activity	NA	0.5 g/L BSA	With	CCCC	CCCC	CCCC	CCCC	CCCC	C0CC	0C00 0C0C	00

a): 1 to 4, virus present and grade of cytophatic effect in 12 units of cellular culture, or grade of cellular lesions in the cytotoxicity assay.

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C = cytopathic effect with presence of virus (in this case and according to Standard/coe's not take into account the degree of cytopathic effect only, the presence or absence of the same).

^{0 =} no virus present or absence of cellular lesions in the cytotoxicity assay; NA: not applicable; NR; not realized; BSA: Bovine serum albumin; PBS: phosphate buffered saline.

sec: seconds; min: minutes.*: sec Special remarks to understand the values of these concentrations.

^{*}Test method accredited # Test performed by subcontractor. Ø Non accredited methods.



Table 3. Results of activity of the test sample with Adenovirus type 5 (ATCC VR-5), under test conditions requested by the client.

Assay	Concentration*	Interfering substance	Cytoto- xicity level	0 min	_	TCID ₅₀ er 30 min	60 min	Reduction with the confidence interval of 95 %
	80%		0.5	-	0.50	-	-	≥ 5.83 ± 0.40
Test sample	3%	0.3 g/L BSA	0.5	-	4.82	-	-	1.51 ± 0.52
	0.03%		0.5	-	6.16	-	-	0.17 ± 0.56
Virus control	NA	0.3 g/L BSA	NA	6.41	6.33	-	-	NA
Formaldehyde	0.7% (w:v)	NA	0.5	NR	NR	1.99	1.41	NA
Virus control Formaldehyde	0.7% (w:v)	NA	0.5	6.08	NR	NR	5.98	NA

Control of sensitivity of cells to virus (difference between decimal logarithm of titre using treated and untreated cells)log10^{-0.41}

NA: not applicable; NR: not realized

Times recommended by Standard for surfaces: maximum 5 or 60 minutes

Times recommended by Standard for instruments: maximum 60 minutes

Times recommended by Standard for Hygienic treatment of hands by friction and hygienic handwashing: between 30 or 120 seconds.

PBS: phosphate buffered saline; BSA: bovine serum albumin.

Virucidal activity exists when the titre of virus shows a reduction ≥4 log.

*: see Special remarks to understand the values of these concentrations.

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Table 4. Results of the activity of the test sample, with Adenovirus type 5 (ATCC VR-5)

(Assay of titration with 12 wells), under test conditions requested by the client.

			Time of				Dilutions	(log10)			
Assay	Concen- tration *	Interfering substance	contact (sec/min)	1	2	3	4	5	6	7	8
	80%		60 sec	0000 0000 0000	0000 0000	0000 0000 0000	0000 0000	0000 0000	0000 0000	0000 0000 0000	NE
Test sample	3%	0.3 g/L BSA	60 sec	4444 4444 4444	4444 4444 4444	4444 4444 4444	3342 0324 4344	0200 1202 0002	0000 0000	0000 0000 0000	NI
	0.03%		60 sec	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	0200 0330 2020	0001 0002 0100	NI
Cytotoxicity	80%	0.3 g/L BSA	NA	0000 0000 0000	0000 0000	0000 0000	0000 0000 0000	0000 0000	0000 0000	0000 0000 0000	N
			0	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	3044 3023 4220	0001 0000 2000	N
Virus control	NA	0.3 g/L BSA	60 sec	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	0200 3304 0332	1000 0020 1000	N
	0.7%	0.7% (w:v) NA	30 min	4344 2324 4233	0200 1020 0002	0000 0000 1100	0000 0000 0000	0000 0000 0000	0000 0000	0000 0000	NI
			60 min	3002 3304 3422	0000 1000 0200	0000 0000	0000 0000 0000	0000 0000	0000 0000 0000	0000 0000	N
Control of formaldehyde cytotoxicity	0.7% (w:v)	0.3 g/L BSA	NA	0000 0000 0000	0000 0000	0000 0000	0000 0000	0000 0000 0000	0000 0000	0000 0000 0000	N
Virus control			0	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	3404 2430 3043	0201 0311 0220	0010 2000 1000	N
formaldehyde	0.7% (w:v)	NA	60 min	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	3403 4442 3434	0201 0202 0010	0000 0000 1100	N
Sensitivity control	N7.4	N7.	Cells not treated	CCCC CCCC	CCCC CCCC	CCCC	CCCC CCCC	CCCC 0CCC	OCOC CCOC COOO	0000 0C00 0C00	N
of cells to virus	NA	NA NA	Cells treated	CCCC CCCC	CCCC	CCCC	CCCC	COCC CCCO	0C00 C0CC 00C0	0000 0000 0000	N
Effectiveness control of the			Without sample	CCCC	CCCC	CCCC	CCCC	cccc	C00C C0C0 0CCC	0000 000C	N
disinfectant detection activity	NA	0.3 g/L BSA	With sample	CCCC	CCCC	CCCC	CCCC	CCCC CCCC	0C00 C0C0 0C80	0000 0000 00C0	N

a): 1 to 4, virus present and grade of cytophatic effect in 12 units of cellular culture of grade of cellular lesions in the cytotoxicity assay.

C = cytopathic effect with presence of virus (in this case and according to Standard does not take into account the degree of cytopathic effect only, the presence or absence of the same).

0 = no virus present or absence of cellular lesions in the cytotoxicity assay; NA: not approache. NR: not realized; BSA: Bovine serum albumin; PBS: phosphate buffered saline.
900; seconds; min: minutes.

*: see Special remarks to understand the values of these concentrations.

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Table 5. Results of activity of the test sample, with Murine Norovirus, strain S99 Berlin, under test conditions requested by the client.

Assay	Concentration*	Interfering substance	Cytoto- xicity level	0 min	-	TCID ₅₀ er 30 min	60 min	Reduction with the confidence interval of 95 %
	80%		0.5	-	0.50	-	-	≥ 5.66 ± 0.34
Test sample	3%	0.3 g/L BSA	0.5	-	3.99	-	-	2.17 ± 0.48
	0.03%		0.5	-	6.07	-	-	0.09 ± 0.53
Virus control	NA	0.3 g/L BSA	NA	6.25	6.16	-	-	NA
Formaldehyde	0.7% (w:v)	NA	0.5	NR	NR	4.24	2.66	NA
Virus control Formaldehyde	0.7% (w:v)	NA	0.5	6.41	NR	NR	6.25	NA

NA: not applicable; NR: not realized

Times recommended by Standard for surfaces: maximum 5 or 60 minutes

Times recommended by Standard for instruments: maximum 60 minutes

Times recommended by Standard for Hygienic treatment of hands by friction and hygienic handwashing; between 30 or 120 seconds.

PBS: phosphate buffered saline; BSA: bovine serum albumin.

Virucidal activity exists when the titre of virus shows a reduction ≥4 log.

*: see Special remarks to understand the values of these concentrations.

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Table 6. Results of the activity of the test sample, with Murine Norovirus strain S99 Berlin (Assay of titration with 12 wells), under test conditions requested by the client.

			Time of					(log10)			
Assay	Concen- tration *	Interfering substance	contact (sec/min)	1	2	3	4	5	6	7	8
	80%		60 sec	0000 0000	NR						
Test sample	3%	0.3 g/L BSA	60 sec	4444 4444 4444	4444 4444 4444	4444 4444 4444	3000 2303 0040	0001 0000 0000	0000 0000 0000	0000 0000	NR
	0.03%		60 sec	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	3443 4203 4434	0200 2012 0302	0011 0000 0000	NR
Cytotoxicity	80%	0.3 g/L BSA	NA	0000 0000 0000	0000 0000 0000	0000 0000 0000	0000 0000 0000	0000 0000 0000	0000 0000 0000	0000 0000	NR
Virus control NA		0.2 -/T DCA	0	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	3444 2334 4434	2011 2201 3201	0000 0000 0000	NR
	NA	0.3 g/L BSA	60 sec	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	3000 4403 0234	0001 0000 0000	NR
Formaldehyde 0.7% (w:v)	0.7%	NA	30 min	4444 4444 4444	4444 4444 4444	4444 4444 4444	3430 4203 0220	0001 0000 0000	0000 0000 0000	0000 0000 0000	NF
			60 min	4444 4444 4444	3320 3223 0442	0100 2000 0110	0010 0100 0000	0000 0000 0000	0000 0000 0000	0000 0000	NR
Control of formaldehyde cytotoxicity	0.7% (w:v)	0.3 g/L BSA	NA	0000 0000 0000	0000 0000 0000	0000 0000 0000	0000 0000 0000	0000 0000 0000	0000 0000	0000 0000 0000	NR
Virus control	0.7%		0	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	3004 0320 3330	0010 0002 0021	000 000
formaldehyde	(w:v)	NA	60 min	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	4444 4444 4444	0303 0404 3200	0010 0020 1000	000 000
ensitivity control			Cells not treated	CCCC CCCC	CCCC CCCC	CCCC	CCCC	CCCC	CC0C 0CC0	000C 00C0 0000	NF
of cells to virus	NA	NA NA	Cells treated	CCCC	CCCC	CCCC	CCCC	CCCC CCCC	00C0 00CC 0000	0000 0000 0000	NF
Effectiveness control of the			Without sample	CCCC	CCCC	CCCC	CCCC	CCCC	C00C CCC0 CCC0	0000 0C0C 0000	NF
disinfectant detection activity	NA	0.3 g/L BSA	With sample	CCCC	CCCC	CCCC	CCCC	COCC COCC	0C00 CC0C 0CCC	00C0 00C0 0000	NR

a): 1 to 4, virus present and grade of cytophatic effect in 12 units of cellular culture, or grade of cellular lesions in the cytotoxicity assay.

NR: not realized; BSA: Bovine serum albumin; PBS: phosphate buffered saline

sec: seconds; min: minutes

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C = cytopathic effect with presence of virus (in this case and according to Standard does not take into account the degree of cytopathic effect only, the presence or absence of the same)

^{0 =} no virus present or absence of cellular lesions in the cytotoxicity assay; NA: not applicable;

^{*:} see Special remarks to ungerstand the values of these concentrations.

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Figure 1. Results of the activity of the test sample under test conditions requested by the client with Poliovirus type 1 (ATCC VR-192).

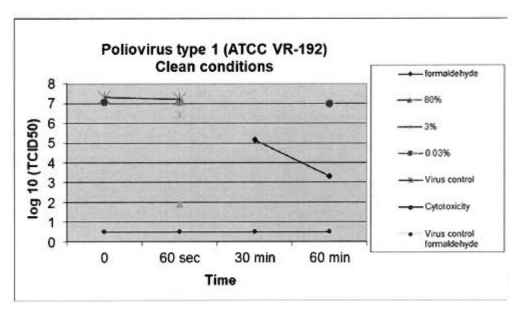
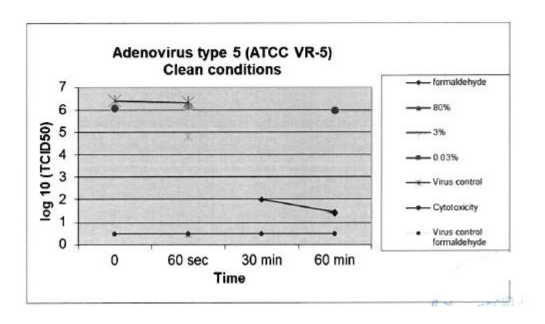


Figure 2. Results of the activity of the test sample under test conditions requested by the client with Adenovirus type 5 (ATCC VR-5).



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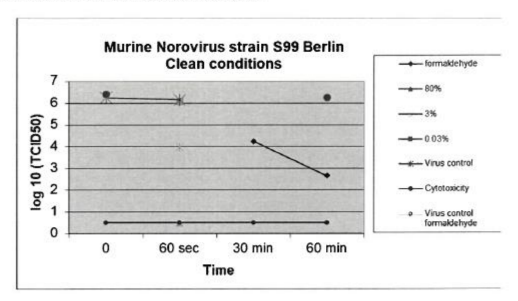
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Figure 3. Results of the activity of the test sample under test conditions requested by the client with Murine Norovirus strain S99 Berlin.



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Date: 13.09.2021

Authorized by: Mariana Ilinca, Manager of Microbiological laboratory



A) IDENTYFICATION OF THE SAMPLE								
Name of the product/Details about the product	DEZINFECTANT UNIVERSAL "BIO-DEZ"							
	Expiration date: 26.01.2024							
	Manufacturer (supplier): ECOCHIM-GRUP							
	Storing conditions: Dry, without sun, 5-25 Celsius							
	degree.							
	Conditions of use: Handrub							
Active(S) substance (S) and its concentration(s)	Ethyl alcohol 72-76%, CAS 64-17-5 and CE 200-578-							
Tetro(b) substance (b) and its concentration(s)	6; Benzalkonium chloride 0.024-0.029%, CAS 68424-							
	85-1 and CE 270-325-2; Methylthioninium chloride							
	0.00024%, CAS 61-73-4 and 200							
C								
Concentrations requested for the assay	3%/ on May 5 the client requested to perform the test at							
D) THE CELL TERM OF	80% concentration (Pure).							
B) TEST METHOD	TIME EN 14240 2005 CI : 1 11: C							
Performed in accredited subcontracted partner laboratory: Scope of Accreditation Nr. 648/LE1286	UNE-EN 14348: 2005. Chemical disinfectants and							
	antiseptics. Quantitative suspension test for the evaluation of							
Report D/21/B0152 Mycobactericidal and tuberculocidal activity of chemical disinfectants in the medical area including	mycobactericidal activity of chemical disinfectants in the medical area including instrument disinfectants. Test							
instrument disinfectants under clean conditions with the product	methods and requirements (phase 2, step 1). AFNOR							
DEZINFECTANT UNIVERSAL "BIO-DEZ" with	methods and requirements (phase 2, step 1). At Work							
deviations from the standard (UNE-EN 14348: 2005 Standard)								
C) METHOD OF ASSAY AND ITS VALIDATION (UNE-EN 14348: 2005 Standard)								
Testing method	DESIN-1052-b // EN 14348: 2005							
Method used	Dilution-neutralization							
Neutralizer	Tryptone 5 g/L, yeast extract 2.5 g/L, dextrose 10 g/L,							
1,000,000	sodium thioglycolate 1 g/L, sodium thiosulfate 1 g/L, sodium							
	bisulphite 2.5 g/L, soya lecithin 7 g/L, polysorbate-80 5 g/L,							
	glycine 1 g/L, 1-histidine 1 g/L and saponim 30 g/L.							
D) INFORMATION ABOUT SAMPLE RECEPTION								
Date of reception of the sample	24.03.2021							
Date of reception of order with test conditions	14.04.2021: 3% concentration							
-	05.05.2021: 80% concentration							
Aspect of the received product	Blue liquid in plastic package.							
E) EXPERIMENTAL CONDITIONS								
Assay period	2021/04/12 to 2021/05/24 (including prior preparation of the							
	strains)							
Solvent of the product used in the assay	Sterile hard water							
Product concentrations for the assay	Pure (80%), 3% and 0.1%							
Aspect of the dilutions of the product	Pure (80%) Blue liquid; 3% and 0.1% transparent							
Contact time	60 seconds							
Assay temperature	20°C ± 1°C							
Interfering substance	Bovine albumin 0.3 g/L							
Stability of the mixture (interfering substance and product	stable							
diluted in sterile hard water) Temperature of incubation	26°C + 1°C							
Temperature of incubation	36°C ± 1°C Mysels actorium guium (ATCC 15760)							
Identification of the origin of viral stains and number of passes	Mycobacterium avium (ATCC 15769) Mycobacterium terrae (CECT 3028 – ATCC 15755)							
	Mycobacterium terrae (CECT 3028 = ATCC 15755)							

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Date: 09.06.2021

Authorized by: Mariana Ilinca, Manager of Microbiological laboratory

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Results of the assay

Control and validation assays......
 Evaluation of mycobactericidal activity...
 See tables 1, 2, 4 and 5
 See tables 3 and 6.

 Number of replicates for each assay microorganism.....

1.

Special remarks

- All controls and validation were between the basic limits.
- One concentration of the sample at least showed a log reduction less than 4 log.
- One concentration of the sample at least showed a log reduction higher than 4 log.
- The highest concentration that can be assayed in the test (80%) is due to the mixtures to perform the assay.
- When the client requested to perform the test at 80% concentration, the test had been started, using hard water.

Conclusion

The product **DEZINFECTANT UNIVERSAL** "BIO-DEZ", batch not indicated, when tested pure (80%), shows mycobactericidal activity after 60 seconds at 20°C under clean conditions (bovine albumin 0.3 g/L), against *Mycobacterium avium* (ATCC 15769) and *Mycobacterium terrae* (CECT 3028 = ATCC 15755), when tested as required by **UNE-EN 14348: 2005** Standard with deviations from the standard since the dilutions of the product, ready to use, have been prepared in sterile hard water instead of in sterile distilled water. The client informed us that the product was ready to use once the test have been started.

Note: The results obtained correspond to the sample received in the laboratory.

Quality Assurance Review:

The assay development and the results obtained have been supervised by the Director of the study.

The Quality Assurance Director has inspected the development of the assay, proving that has been realized following the proper procedure and using the adequate media, materials and reagents, following as well the Good Laboratory Practices (GLPs) and the final report contains the primary data obtained.

Reference:

 UNE-EN 14348: 2005. Chemical disinfectants and antiseptics. Quantitative suspension test for the evaluation of mycobactericidal activity of chemical disinfectants in the medical area including instrument disinfectants. Test methods and requirements (phase 2, step 1). AENOR.

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Table 1.-Assay with Mycobacterium avium (ATCC 15769): Validation and controls.

Suspension of validation (Nv ₀)		Control of experimental conditions (A)			Neutralizer (B)			Validation of the method (C) with sample concentration: Pure (80%)				
Vc1	135	X=	Vc_1	125	V- 122	Vc_1	121	X= 124	Vc_1	130	X=	
Vc_2	124	129.5	Vc_2	119	X= 122	Vc_2	127		Vc_2	119	124.5	
30 ≤	$30 \le x \text{ of } Nv_0 \le 160$?			$x \text{ of } A \text{ is } \ge 0.5 \text{ x X of}$			$x \text{ of } B \text{ is } \ge 0.5 \text{ x of }$			$x \text{ of } C \text{ is } \ge 0.5 \text{ X of}$		
	Yes			Nv ₀ ? Yes			Nν ₀ ? Yes			Nv ₀ ? Yes		

Table 2.- Assay with Mycobacterium avium (ATCC 15769): Suspension of the assay.

Suspension of	N	Vc1	Vc2	$X_{wm} = 4.85 \times 10^9 = 1g = 9.69$
the assay (N y	10-7	>660	>660	$N_0 = N/10 = 1g = 8.69$
N ₀)	10-8	51	46	$8.17 \le N_0 \le 8.70$? Yes

Table 3.- Assay with Mycobacterium avium (ATCC 15769).

Concentrations of the sample (%)	Dilutions	Vc ₁	Vc2	$Lg Na = lg$ $(X \times 10 \text{ o}$ $Xwm \times 10)$	LgR (lgN ₀ = 8.69)	Time of contact (seconds)	
	10 ⁰	<14	<14				
Dame (900/)	10 ⁻¹	<14	<14	<2.15	>6.54	60	
Pure (80%)	10-2	<14	<14				
	10-3	<14	<14				
	10°	>660	>660			60	
20/	10-1	>660	>660	>6.82	<1.87		
3%	10-2	>660	>660				
	10-3	>660	>660				
	10 ⁰	>660	>660				
0.10/	10-1	>660	>660	>6.82	-1.07	60	
0.1%	10-2	>660	>660	70.82	<1.87		
	10 ⁻³	>660	>660				

Observations:

$$Nv_0$$
: 74 + 61; 59 + 65;

B:
$$63 + 58$$
; $71 + 57$;

Na Pure (80%)
$$10^{\circ}$$
: $0 + 0$; $0 + 0$;

$$3\% 10^{-3}$$
: >330 + >330; >330 + >330;

$$0.1\% 10^{-3}$$
; >330 + >330; >330 + >330;

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Table 4.-Assay with *Mycobacterium terrae* (CECT 3028 = ATCC 15755): Validation and controls.

S	Suspension of validation (Nv ₀)		Control of experimental conditions (A)			Neutralizer (B)			Validation of the method (C) with sample concentration: Pure (80%)			
Vc1	53	X= 55	Vc_1	54	X= 52	Vc_1	50	X= 49	Vc_1	49	X= 47	
Vc_2	57	A= 33	Vc_2		A- 32	Vc_2	48	A= 49	Vc_2	45	A- 4/	
30 ≤	$30 \le x \text{ of } Nv_0 \le 160$?			$x \text{ of } A \text{ is } \ge 0.5 \text{ x X of}$			$x \text{ of } B \text{ is } \ge 0.5 \text{ x of }$			$x \text{ of } C \text{ is } \ge 0.5 \text{ X of }$		
	Yes			Nv ₀ ? Yes			Nv ₀ ? Yes			Nv ₀ ? Yes		

Table 5.- Assay with *Mycobacterium terrae* (CECT 3028 = ATCC-15755): Suspension of the assay.

Suspension of	N	Vc1	Vc_2	$X_{wm} = 2.19 \times 10^9 = 1g = 9.34$
the assay (N y	10-7	227	211	$N_0 = N/10 = 1g = 8.34$
N_0)	10-8	21	22	$8.17 \le N_0 \le 8.70$? Yes

Table 6.- Assay with Mycobacterium terrae (CECT 3028 = ATCC 15755).

Concentrations of the sample (%)	Dilutions	Vc ₁	Vc2	Lg Na = lg (X x 10 o Xwm x 10)	LgR (lgN ₀ = 8.34)	Time of contact (seconds)	
	10 ⁰	<14	<14				
Dama (900/)	10-1	<14	<14	<2.15	>6.19	60	
Pure (80%)	10-2	<14	<14				
	10-3	<14	<14				
	10°	>660	>660	>6.92		60	
20/	10-1	>660	>660		<1.52		
3%	10-2	>660	>660	>6.82			
	10-3	>660	>660				
	10 ⁰	>660	>660				
	10-1	>660	>660	>6.92	-1.50	60	
0.1%	10-2	>660	>660	>6.82	<1.52		
	10-3	>660	>660				

Observations:

Explanations:

Vc: Counts per mL

Xwm: ponderated mean of X

X: Values of Vc_1 and Vc_2 (1. + 2. duplicates); R: reduction (LgR = lgN₀ - lgNa)

R: reduction (LgR = $\lg N_0 - \lg Na$)

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