mgsLABORATORIES

Doc No.

TRA-2012-080-02

Title	Microbiological Analys Quantitative carrier tes instruments used in the (Phase 2 / Step 2)	st for the evalua	tion of bacte		ity for
Product	Clinell Universal Sanitising Wipes	MGS No	02645	SO No	3269



a) Identification of test laboratory:

Test laboratory

MGS Laboratories Ltd Unit 14, Newlands Drive

Poyle 14 Horton Road Poyle Berkshire SL3 0DX

b) Identification of the Customer:

Customer Name

Gama Healthcare Ltd

Unit 2

Customer Address

Brent Cross Gardens,

Brent Cross , London , NW4 3RJ

c) Identification of the sample:

Name of product

Clinell Universal Sanitising Wipes

Batch number (and expiry date if

available)

UB1205 Exp Feb 17

Manufacturer

Gama Healthcare Ltd

Date of delivery

23 Mar 12

Storage conditions

Room temperature and darkness

Product diluent recommended by the

manufacturer for use

N/A

Active substance(s) and their

concentration(s) (optional)

Not stated

Appearance of the product

Wet wipe

d) Test method and its validation:

MGS procedure reference

WIN-1000.060-05

Method

Dilution neutralisation

Neutraliser

Tryptone soya broth 30g/l, lecithin 30g/l, polysorbate 20 100g/l, Catalase

0.25g/l

Details of validation of the neutraliser

Neutraliser validation performed according to 5.5.2 in EN 14561:2006

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Doc No. Microbiological Services and Consultancy

TRA-2012-080-02

Microbiological Analysis Based on EN 14561 (2006) Quantitative carrier test for the evaluation of bactericidal activity for Title instruments used in the medical area. (Phase 2 / Step 2)

Clinell Universal Product Sanitising Wipes

MGS No 02645 SO No

3269



e) Experimental conditions:

Period of analysis

29 Mar 12 - 13 Apr 12

Product diluent used during the test

N/A

Product test concentrations

Ready to Use (RTU)

Product preparation

Solution was squeezed out of the wipes and used for testing

Appearance of product dilutions

N/A

Contact time

60 seconds ± 10s

Test temperature range

20°C ± 2°C

Interfering substance

3.0ml/l sheep erythrocytes and 3.0g/l Bovine albumin

Stability and appearance of the

mixture

Precipitate absent throughout test

Temperature of incubation

36°C ± 2°C

Identification of the bacterial strains

used

Pseudomonas aeruginosa ATCC 15442 Staphylococcus aureus

Enterococcus hirae

ATCC 6538 **NCIMB 8192**

f) Results:

Test results

1) Controls and validation

2) Evaluation of bactericidal activity

g) Conclusion:

Based on EN 14561 (2006), the batch UB1205 of the product Clinell Universal Sanitising Wipes, when tested at RTU, possesses bactericidal activity in 60 seconds at 20°C under dirty conditions for the referenced

strains of P. aeruginosa, S. aureus and E. hirae.

h) Deviations:

None

Re-issued By:

Name: Mrs Emma Newton BSc (Hons)

Approved by:

Name: Mrs Kim Morwood BSc (Hons) CBiol MiBiol

Position: Quality Manager

Position: Technical Director

Date:

19April

Date: 19 APRIZ

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	LABORATORI Microbiological Services and Consu		Doc No.	TRA-2	012-080-02	<u></u>
Title	Microbiological Analys Quantitative carrier tes instruments used in the (Phase 2 / Step 2)	st for the evalua	tion of bacteric	idal activ	rity for	UKAS TESTING
Product	Clinell Universal Sanitising Wipes	MGS No	02645	SO No	3269	4393

The MGS procedure WIN-1000.060 is a laboratory method based on the EN 14561 (2006) standard; the minor deviations from the standard, which do not affect the overall results, are detailed below:

- The carriers are sterilised in an autoclave rather than by dry heat.
- EN 14561 states an allowed tolerance of 36°C ±1°C or 37°C±1°C, MGS laboratories equipment is validated to ±2°C therefore MGS procedures state ±2°C.
- Organisms are prepared by swabbing plates and adding to 9ml diluent to form a suspension, rather than adding loopfuls of organism to 10ml diluent with beads, shaking for 3 minutes, aspirating and adding to a new container. Swabbing forms a smooth suspension removing the need to shake with beads.
- The laboratory is regulated at 20°C; therefore for testing at 20°C a temperature controlled chamber is not used.
- Hard water is prepared as stated in EN 1276:2009, solution A is autoclaved, Solution B is filter sterilised so SHW does not require sterilisation.
- Sterilisation of BSA is performed as stated in EN 1276:2009 0.22µm filters are used to filter sterilize BSA and hard water, not 0.45µm filters.
- Once inoculated surfaces are dried in a validated manner rather than at 36°C±1°C.
- Neutraliser is prepared at 8ml, 9ml and 10ml taking into account required concentrations so that water does not have to be added to 8ml/9ml for Test, Nw and NTV aliquots.
- Neutralisation is not proved prior to the test, but is validated in the test.
- Dilutions of controls plated differ, if the plated dilutions were not correct and the plates had <1cfu
 or >300cfu the test would be invalid.
- Plates are incubated for the full time rather than performing an interim read; in addition the incubation period may be extended to a maximum of 4 days due to business hours.
- Any part of the method may be altered to meet customer requirements; MGS does not insist on testing the standard conditions or three concentrations of product with replicates of the limiting organism

mgsLABORATORIES Doc No. TRA-2012-080-02 Microbiological Services and Consultancy Microbiological Analysis Based on EN 14561 (2006) Quantitative carrier test for the evaluation of bactericidal activity for Title instruments used in the medical area. (Phase 2 / Step 2) **Clinell Universal** 4393 SO No 3269 Product MGS No 02645 **Sanitising Wipes**

Draduot botob number LIB1205		
Product batch number: UB1205		
Appearance of product: Clear colourless solution	X	
Number of plates: 1/ml	Pour plate	Spread plate
Neutraliser: Tryptone soya broth 30g/l, lecithin 30g/l, p	olysorbate 20 100g/l, catalas	se 0.25g/l
Actual test temperature: 20°C		
Interfering substances: 3.0g/l Bovine albumin and 3ml/	/I sheep erythrocytes	
Test organism: P. aeruginosa ATCC 15442	2	
Drying time on carrier: 15 minutes (not >60 minutes)		
Incubation temperature: 36°C ± 2°C	P	
Date of Test: 11 Apr 12	1	
	Signature:	

Validation and Controls

Validation suspension (Nv ₀)		Experimental Conditions Control (A)			Neutraliser Control (B)			Method Validation (C) Prod conc: 1 wipe in 75ml water			
Vc1	38	x =40	Vc1	59	χ =56	Vc1	61	x =70	Vc1	91	χ =75
Vc2	42		Vc2	52		Vc2	78		Vc2	59	
		χ of A Yes	χ of A is $\geq 0.5 \times \chi$ of Nv ₀ ?		χ of B is ≥ 0.5 x χ of Nv ₀ ? Yes χ No			χ of C is \geq 0.5 x χ of Nv ₀ ? Yes χ No			

Test suspension, water control and test

Test suspension (N and N ₀):	N	Vc1	Vc2	x wm = 202 x 10'; lg N =	9.31		
	10" 178	223	$N_0 = N/10$; $\log N_0 = 8.31$ $8.17 \le \log N_0 \le 8.70$?	Yes	Х	No	
	10 ⁻⁸	25	19	0.17 = 19 N ₀ = 0.70?	165	^	140

Water Control Nw)	Nw	Vc1	Vc2	$\chi \times 10 = 157 \times 10^{5}$; Ig Nw = 7.20 7.15 \le Ig Nw = 7.20 \le (IgN-1.3)?
(,,,,,	10-4	163	150	
	10 ⁻⁵	12	16	Yes X No

Conc of the product	Dilution Step	Vc1	Vc2	Na = χ x10	lg Na Lg(χ)	Ig R (IgNw = 7.20)	Contact time
	10 ⁰	<14	<14	-110	-0.45		60 sec
DTU	10-1	<14	<14				
RTU	10-2	<14	<14	<140	<2.15	>5.05	
	10-3	<14	<14	1			

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Product batch number: UB1205		
Appearance of product: Clear colourless solution	X	
Number of plates: 1/ml	Pour plate	Spread plate
Neutraliser: Tryptone soya broth 30g/l, lecithin 30g/l,	polysorbate 20 100g/l, catalas	se 0.25g/l
Actual test temperature: 20°C		Visit 61.51 (1) — VI
Interfering substances: 3.0g/l Bovine albumin and 3r	ml/l sheep erythrocytes	
Test organism: S. aureus ATCC 6538		
Drying time on carrier: 15 minutes (not >60 minutes)		
Incubation temperature: 36°C ± 2°C	/)	
Date of Test: 11 Apr 12		
Person responsible: Janice Wong	Signature:	

Validation and Controls

Validation suspension (Nv ₀)			imental tions C	ontrol (A)	Neutraliser Control (B)			Method Validation (C) Prod conc: RTU			
Vc1	105	χ = 96	Vc1	123	x = 109	Vc1	101	χ =98	Vc1	114	x =107
Vc2	86		Vc2	94		Vc2 95		Vc2	99		
$30 \le \chi$ of Nv ₀ ≤ 160 ? χ of A is $\ge 0.5 \times \chi$ of Nv ₀ ? Yes χ No Yes χ No		χ of B is ≥ 0.5 x χ of Nv ₀ ? Yes χ No			χ of C is $\geq 0.5 \times \chi$ of Nv ₀ ? Yes χ No						

Test suspension, water control and test

Test suspension (N and N ₀):	N	Vc1	Vc2	$\chi = 50 \times 10^8$; $\lg N = 9.70$ $N_0 = N/10$; $\lg N_0 = 8.70$			
	10-7	>330	>330	$8.17 \le \lg N_0 \le 8.70$?	Yes	Х	No
	10 ⁻⁸	50	50	0.17 = 19 No = 0.70?	Les	^	L

Water Control (Nw)	Nw	Vc1	Vc2	$\chi \times 10 = 123 \times 10^{6}$; Ig Nw = 8.0 7.15 \le Ig Nw = 8.09 \le (IgN-1.3)?
,,	10-4	>330	>330	7.10 2 lg (tw = 0.00 2 (lg (t=1.0))
	10 ⁻⁵	125	121	Yes X No

Conc of the product	Dilution Step	Vc1	Vc2	Na = χ x10	lg Na Lg(χ)	Ig R (IgNw = 8.09)	Contact time
	10 ⁰	25	29				60 sec
DTU	10-1	<14	<14				
RTU	10-2	<14	<14	270	2.43	5.66	
	10-3	<14	<14			L	

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mgsLABORATORIES Doc No. TRA-2012-080-02 Microbiological Services and Consultancy Microbiological Analysis Based on EN 14561 (2006) Quantitative carrier test for the evaluation of bactericidal activity for Title instruments used in the medical area. (Phase 2 / Step 2) Clinell Universal SO No 3269 MGS No 02645 Product **Sanitising Wipes**

Product batch number: UB1205	·	<u> </u>
Appearance of product: Clear colourless solution	Pour plate X	
Number of plates: 1/ml	Pour plate	Spread plate
Neutraliser: Tryptone soya broth 30g/l, lecithin 30g/l, pol	lysorbate 20 100g/l, catala	se 0.25g/l
Actual test temperature: 20°C		
Interfering substances: 3.0g/l Bovine albumin and 3ml/l	sheep erythrocytes	
Test organism: E. hirae NCIMB 8192	500 F - 1440	
Drying time on carrier: 15 minutes (not >60 minutes)		
Incubation temperature: 36°C ± 2°C	ž.	

Validation and Controls

Person responsible: Janice Wong

Date of Test: 11 Apr 12

Valida (Nv₀)	ation su	spension	Experimental Conditions Control (A)		Neutraliser Control (B)			Method Validation (C) Prod conc: RTU			
Vc1	93	x =79	Vc1	83	χ =83	Vc1	79	χ =73	Vc1	60	χ =69
Vc2	65	10000	Vc2	83	177.0	Vc2	66	1000	Vc2	77	
30 ≤ x Yes	of Nv ₀	≤ 160? No	χ of A is $\geq 0.5 \times \chi$ of Nv ₀ ? Yes χ No			χ of B is \geq 0.5 x χ of Nv ₀ ? Yes χ No			χ of C is \geq 0.5 x χ of Nv ₀ ? Yes χ No		

Signature:

Test suspension, water control and test

Test suspension (N and N ₀):	N	Vc1	Vc2	$\chi = 38 \times 10^8$; Ig N = 9.58 $N_0 = N/10$; Ig $N_0 = 8.58$			
	10-7	>330	>330	$- \frac{10_0 - 10_110_1 \text{ ig N}_0 - 8.38}{8.17 \le \text{Ig N}_0 \le 8.70?}$	Yes	X	No
	10 ⁻⁸	39	37		103	^	140

Nater Control Nw)	Nw	Vc1	Vc2	$\chi \times 10 = 46 \times 10^{5}$; Ig Nw = 7.66 7.15 \le Ig Nw = 7.66 \le (IgN-1.3)
(,	10-4	>330	>330	7.10 2 ig (till = 7.50 2 (ig)(* 1.5)
	10 ⁻⁵	45	47	Yes X No

Conc of the product	Dilution Step	Vc1	Vc2	Na = χ x10	lg Na Lg(x)	Ig R (IgNw = 7.66)	Contact
RTU	10°	<14	<14	<140		>5.51	60 sec
	10-1	<14	<14		<2.15		
	10-2	<14	<14				
	10-3	<14	<14				

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Title	Microbiological Analysis Based on EN 14561 (2006) Quantitative carrier test for the evaluation of bactericidal activity for instruments used in the medical area. (Phase 2 / Step 2)					UKAS TESTING
Product	Clinell Universal Sanitising Wipes	MGS No	02645	SO No	3269	4393

Explanations:

Vc = count per ml

χ = average of Vc1 and Vc2 (1. + 2. duplicate)

 χ wm = weighed mean of χ

Na = number of survivors in the test mixture

Nv = number of cells in the validation suspension

 $Nv_0 = Nv/10$

 $R = reduction (lg R = lgN_w - lgNa)$

If Na<140, IgR = > [IgNw - 2.15]

If Na > X x 10, $IgR = \langle [IgN_w - Ig X] (X = upper limit for Vc)$

All test results have an associated uncertainty of measurement; for this test the expanded uncertainty is based on the estimated uncertainty multiplied by a coverage factor K=2 providing a level of confidence of approximately 95%. The uncertainty evaluation has been assessed in accordance with MGS laboratories' UKAS Accreditation and is available on request.