Everyday Virucidal Disinfectant Portfolio



PVA Hygiene provides an innovative and sustainable method of cleaning. As the UK's leading manufacturer of water-soluble cleaning products, we cover all areas of commercial cleaning. Over 24 years, we have developed a system using pre-dosed sachets that is straightforward to implement and balances environment diligence with commercial demands. Based in the South West of England, we distribute globally.



This portfolio contains documents relating to PVA Hygiene's EVERYDAY VIRUCIDAL DISINFECTANT.

This unique formulation is contained within a PVA-OH film that dissolves at the point of use. The sachets are dry, compact and light, they reduce storage space and transportation costs, and heavily reduce the environmental implications often associated with delivering cleaning supplies. The sachets are packed in planet friendly packaging, that can either be composted or recycled, helping you to eliminate single-use plastic from your current cleaning procedure.



CONTENTS:

- 1) Technical Data Sheet.
- 2) Use Solution Health and Safety Summary.
- 3) Efficacy Data Reports.
- 4) Product Safety Data Sheet.







PRODUCT DESCRIPTION

Everyday Virucidal Detergent Sanitiser is based on PVA Hygiene's unique Aqua-Dis PDCS9 technology. Sachets contain a blend of biodegradable chelates, together with biodegradable surfactants and a cationic disinfectant. The product is designed for routine cleaning and disinfection of surfaces. Everyday Virucidal Disinfectant is safe for use on normal materials of construction, and when used as directed this product conforms to EN1276, EN1650, EN13697 and EN14476 (Enveloped Viruses).

Sachets are supplied in the following Pack Sizes:-

ſ	Pack Size	Pack Size Sachet Type		Outer Packaging		
ſ	20 * 15g	PVA-OH	DZ4:20	Pouch		

- Supplied in convenient PVA-OH water soluble sachets within a compostable container.
- Broad Spectrum Bacterial and Virucidal Activity.
- Phosphate Free.
- Perfume Free.

INSTRUCTIONS FOR USE

For general cleaning, remove any gross debris from the surface, place one sachet into the empty trigger spray bottle, and fill with water to the 750ml mark. Replace the trigger head and shake until the sachet has dissolved (note warm water will aid the rate of dissolution but is not essential). Spray the solution onto the surface and wipe clean. For disinfection apply a second spray to the clean surface and allow to air dry over 5 minutes.

Once made, in use solutions are expected to have a shelf life of at least a week.

TECHNICAL DATA SUMMARY

Appearance	White Powder
Odour	Non distinct (Perfume free)
Foam	Low
pH of use solution	10 - 11
Storage Temperature Range	0°C to +40°C
Shelf Life of Sachet	Minimum of 2 years under normal conditions of dry storage.

PVA Hygiene, Unit 6, Havyat Road Business Park, Havyat Road, Bristol, BS40 5PA. Tel: +44 (0) 1934 862859 Email: sales@pva-hygiene.co.uk



EFFICACY DETAILS

Test	Complia	ance Conditions	Organism Type/Compliance			
	Time /	Minimum				
	Minutes	Concentration				
EN1276	5	(1 sachet /750ml)	Claim supported by standard organisms of:-			
			Pseudomonas aeruginosa.			
			Escherichia coli.			
			Enterococcus hirae.			
			Staphylococcus aureus.			
EN1650	5	(1 sachet /750ml)	Claim supported by standard organisms of:-			
			Candida Albicans			
EN13697	5	(1 sachet /750ml)	Claim supported by standard organisms of:-			
			Pseudomonas aeruginosa.			
			Escherichia coli.			
			Enterococcus hirae.			
			Staphylococcus aureus.			
EN13697	1	(1 sachet / 750ml)	Streptococcus pyogenes (Strep A)			
EN14476	5	(1 sachet / 750ml)	Vaccinia virus VR-1508.			

EMERGENCY DETAILS

For accident, emergency and health & safety information refer to the Safety Data Sheet for this product.

This product is registered with the UK National Poisons Information Service.

Office Hours Emergency Number +44 (0) 1934 862859

Outside Office Hours: - +44 (0)7967 149256 (This is for health, safety and environmental emergencies only, it is not for general enquires or ordering).

DISCLAIMER

Whilst every effort is made to ensure that the information given in this product information sheet is accurate it is given without guarantee, since the conditions of use are beyond our control.



EVERYDAY VIRUCIDAL DISINFECTANT USE SOLUTION HEATH AND SAFETY SUMMARY

Issue Date 20/05/2023 Version 2.0

IDENTIFICATION OF TH	IDENTIFICATION OF THE MATERIAL						
Product Name Everyday Virucidal Disinfectant use solution							
Main Use Cleaning and Disinfecting Hard Surfaces and Floors							
Uses Advised Against	Not for Direct Oral Consumption						
	Keep Out of Reach of Children						
	Do Not Mix with other Chemicals/Detergents.						
Manufacturer	PVA Hygiene, Unit 6 Havyat Business Park						
	Havyat Road, Bristol, BS40 5PA						
Telephone	+44 (0) 1934 862859						

PHYSICAL AND CHEMICAL F	PROPERTIES
Appearance	Clear Liquid
Colour	Colourless
рН	10-11.0

CLASSIFICATION, PPE	CLASSIFICATION, PPE, FIRST AID AND DISPOSAL						
Health	In use solutions of this product have no Health Classifications						
Physical	In use solutions of this product have no Physical Classifications						
Environmental	In use solutions are classified as H412 Harmful to Aquatic Life with Long						
	Term Effects						
PPE	No PPE is mandated for this product at use strength. However, we						
	suggest gloves for general hygiene.						
First Aid	EYES:-						
	May cause reddening, discomfort and blurred vision						
	Rinse with Plenty of Water.						
	SKIN:-						
	Repeated extended contact may result in skin dryness.						
	Use a suitable re-moisturising cream and get medical attention if						
	symptoms persist.						
	INHALATION:-						
	Unlikely.						
	INGESTION:-						
	A soapy taste may be reported, together with irritation to mouth						
	and GI Tract rinse mouth thoroughly.						
	If concerned seek medical advice						
	Show the label or Safety Data sheet to the Physician.						
Disposal	Solutions can be disposed to normal sewers and septic tanks.						

PVA Hygiene, Unit 6, Havyat Road Business Park, Havyat Road, Bristol, BS40 5PA. Tel: +44 (0) 1934 862859 Email: sales@pva-hygiene.co.uk

BS EN 1276:2019





Company Name:	PVA Hygiene Ltd
Contact Name:	Jim Taylour
Contact Email:	technical@pva-hygiene.co.uk
Purchase Order No:	ТВС
Report Date:	28/06/2021
Melbec Ref Number: No. of Samples:	28489 1
Name of Test Product: Batch Number:	PDCCS9 A Surface Disinfectant N/A



BS EN 1276:2019

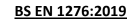


Sample Details:

	Product storage condition Appearance of the produ Appearance of the produ Appearance of product w Active substance and con Product dilution preparat	ns: ct (as supplied): ct (after dilution): ith interfering substance and test organism: centration: ion trations:	PVA Hygiene Ltd Ambient White Sachet Cloudy Liquid Cloudy Liquid Benzalkonium Chloride Volume/Volume 1 x 10g Sachet dissolved into 600ml of water (Volume of water adjusted from 750ml to allow for dilution during test)				
	Diluent used to dilute pro	oduct:	Synthetic Hard Water				
	Incubation temperature:		35 °C to 38 °C				
	The test product was in s	atisfactory condition for testing when receive	ed.				
	Date product received:	07/06/21	Test Date:	17/06/21			
Experime	ntal Conditions: Interfering substance: Test temperature: Contact time: Test organisms:	Bovine Albumin (clean 0.3g/l) 18 °C to 25 °C 5 minutes Pseudomonas aeruginosa ATCC 15442 Staphylococcus aureus ATCC 6538 Escherichia coli ATCC 10536 Enterococcus hirae ATCC 10541					

Deviations:

EN1276 states incubation temperature of 36±1°C or 37±1°C. Melbec Microbiology Ltd method states 35°C - 38°C.







Requirements of the Standard:

The test product shall demonstrate at least a 5 decimal logarithm (lg) reduction when tested in accordance with this standard under simulated clean or dirty conditions.

Conclusion:

For the product PDCCS9 A Surface Disinfectant, [Batch code: N/A] the log reduction requirements as specified in EN 1276:2019 (5 lg within the relevant contact time) were met in clean conditions with a contact time of 5 minutes.

Report authorised by:

354

Name: Position: Date:

Dawn Mellors **Technical Director** 28/06/2021

All samples are tested as received and the condition on receipt is deemed to be satisfactory for testing unless client is informed otherwise. If an unsatisfactory sample is received and tested on instruction from the client comments are included on the report detailing this information. Results given for this may be invalid. Results detailed above relate only to the samples tested. Sample description and batch references stated are as provided by the customer. This test report shall not be reproduced except in full without the approval of Melbec Microbiology Ltd.

Test Report for a General-Purpose Disinfectant Product

BS EN 1276:2019





Test Results:

Neutralisation Method Used:

Membrane filtration

Rinsing Liquid Used: N7

Test Report for a General-Purpose Disinfectant Product BS EN 1276:2019



Pseudomonas aeruginosa ATCC

1544	2		Validation and controls							Melbec Ref No 2848	
Validation suspension (Nv ₀)			Experimental conditions control (A)			Neutra	lizer contro	ol (B)	Method validation (C) 10g in 750ml water		
Vc 1	72	x =	Vc 1	45	x =	Vc 1	43	x =	Vc 1	44	x =
Vc 2	66	69	Vc 2	43	44	Vc 2	38	40.5	Vc 2	32	38
30 ≤ <i>x</i>	$30 \le \overline{x}$ of $Nv_0 \le 160$? Yes		\overline{x} of A is $\ge 0.5 \text{ x } \overline{x}$ of Nv_0 ? Yes			\overline{x} of B is $\ge 0.5 \times \overline{x}$ of Nv_0 ? Yes			\overline{x} of C is $\ge 0.5 \times \overline{x}$ of Nv _o ? Yes		

Test suspension

	N	<i>Vc</i> 1	Vc 2	X m 4.65E+08 ; lg N = 8.67
Test suspension (<i>N</i> and <i>N</i> ₀):	10 -6	>330	>330	$N_0 = N/10$; lg $N_0 = 7.67$
(/• and /• ₀).	10 ⁻⁷	53	40	$7.17 \le \lg N_0 \le 7.70$? Yes
				\overline{X} quotient = >5 and <15? N/A

Bactericidal activity results

Conc. of the active (%)	Vc 1	Vc 2	$Na = \overline{X} x 10$	lg Na	lg N _ =	R 7.67	Contact time	Result
1 x 10g Sachet dissolved into	<14	<14	1.40E+02	<2.15	/v ₀ –	>5.52	5 minutes	Pass
600ml of water								

Test Report for a General-Purpose Disinfectant Product



BS EN 1276:2019



Staphylococcus aureus ATCC

6538 Validation and controls									Melbec Ref No 2848			
Validation suspension (<i>Nv</i> ₀)			Experimental conditions control (A)			Neutra	lizer contro	ol (B)	Method validation (C) 10g in 750ml water			
Vc 1	83	x =	<i>Vc</i> 1	105	x =	<i>Vc</i> 1	73	x =	Vc 1	82	x =	
Vc 2	74	78.5	Vc 2	87	96	Vc 2	72	72.5	Vc 2	75	78.5	
30 ≤ X	$30 \le \overline{x}$ of $Nv_0 \le 160$? Yes		\overline{X} of A is \ge	\overline{x} of A is $\ge 0.5 \times \overline{x}$ of Nv_0 ? Yes			\overline{X} of B is $\ge 0.5 \times \overline{X}$ of Nv_0 ? Yes			\overline{x} of C is $\ge 0.5 \times \overline{x}$ of Nv ₀ ? Yes		

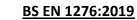
Test suspension

	Ν	<i>Vc</i> 1	Vc 2	X m 3.60E+08 ; lg N = 8.56
Test suspension (<i>N</i> and <i>N</i> ₀):	10 ⁻⁶	>330	>330	N ₀ = N/10 ; lg N ₀ = 7.56
(/• and /• ₀).	10 ⁻⁷	37	35	$7.17 \le \lg N_0 \le 7.70$? Yes
				\overline{X} quotient = >5 and <15? N/A

Bactericidal activity results

Conc. of the active (%)	Vc 1	Vc 2	$Na = \overline{X} x10$	lg Na	lg N ₀ =	R 7.56	Contact time	Result
1 x 10g Sachet dissolved into	<14	<14	1.40E+02	<2.15		>5.41	5 minutes	Pass
600ml of water								

Test Report for a General-Purpose Disinfectant Product







Escherichia coli ATCC 10536

		Validation and controls							Melbec R	ef No	28489
Validat	tion suspe (<i>Nv</i> ₀)	ension	-	ental cond ntrol (A)	litions	Neutra	lizer contro	ol (B)		od validatior g in 750ml wa	``'
<i>Vc</i> 1	104	x =	Vc 1	87	x =	Vc 1	76	x =	Vc 1	74	x =
Vc 2	95	99.5	Vc 2	67	77	Vc 2	66	71	Vc 2	71	72.5
30 ≤ X	c of Nv ₀ s Yes	≤ 160?	\overline{X} of A is \ge	0.5 x x o Yes	of Nv ₀ ?	\overline{X} of B is	≥ 0.5 x X o Yes	of Nv ₀ ?	\overline{x} of C	is ≥ 0.5 x <i>X</i> o Yes	f Nv _o ?

Test suspension

	Ν	<i>Vc</i> 1	Vc 2	X m 4.95E+08 ; lg N = 8.69
Test suspension (<i>N</i> and <i>N</i> ₀):	10 ⁻⁶	>330	>330	N ₀ = N/10 ; lg N ₀ = 7.69
(N and N ₀).	10 ⁻⁷	52	47	$7.17 \le \lg N_0 \le 7.70$? Yes
				\overline{X} quotient = >5 and <15? N/A

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Bactericidal activity results

Conc. of the active (%)	Vc 1	Vc 2	$Na = \overline{X} x 10$	lg Na	lg. N ₀ =	R 7.69	Contact time	Result
1 x 10g Sachet dissolved into	<14	<14	1.40E+02	<2.15		>5.55	5 minutes	Pass
600ml of water								

Test Report for a General-Purpose Disinfectant Product BS EN 1276:2019





Enterococcus hirae ATCC 10541

	Validation and controls							Melbec R	ef No	28489	
Validat	tion suspe (Nv ₀)	ension		ental conc ntrol (A)	litions	Neutra	lizer contro	ol (B)		nod validatior g in 750ml wa	
Vc 1	115	x =	Vc 1	85	x =	<i>Vc</i> 1	99	x =	Vc 1	92	x =
Vc 2	76	95.5	Vc 2	64	74.5	Vc 2	82	90.5	Vc 2	81	86.5
30 ≤ X	c of Nv ₀ s Yes	≤ 160?	\overline{X} of A is \ge	0.5 x X o Yes	of Nv ₀ ?	\overline{x} of B is	≥ 0.5 x X o Yes	of Nv ₀ ?	\overline{x} of C	is ≥ 0.5 x <i>X</i> o Yes	f Nv _o ?

Test suspension

	Ν	<i>Vc</i> 1	<i>Vc</i> 2	X m 4.00E+08 ; lg N = 8.60
Test suspension (N and N ₀):	10 ⁻⁶	>330	>330	N ₀ = N/10 ; lg N ₀ = 7.60
(/• and /• ₀).	10 ⁻⁷	42	38	$7.17 \le \lg N_0 \le 7.70$? Yes
				\overline{X} quotient = >5 and <15? N/A

Bactericidal activity results

Conc. of the active (%)	Vc 1	Vc 2	$Na = \overline{X} x 10$	lg Na	ן N ₀ =	R 7.60	Contact time	Result
1 x 10g Sachet dissolved into	<14	<14	1.40E+02	<2.15		>5.46	5 minutes	Pass
600ml of water								

Test Report General-Purpose Disinfectant Product



BS EN 1650:2019

Company Name:	PVA Hygiene Ltd
Contact Name:	Jim Taylour
Purchase Order No:	ТВС
Report Date:	28/06/2021
Melbec Ref Number: No. of Samples:	28490 1
Name of Test Product: Batch Number:	PDCCS9 A Surface Disinfectant N/A

Test Report General-Purpose Disinfectant Product BS EN 1650:2019



Sample Details:

Manufacture / Supplier:	PVA Hygiene Ltd
Product storage conditions:	Ambient
Appearance of the product (as supplied):	Sachet
Appearance of the product (after dilution):	Cloudy liquid
Appearance of product with interfering substance and test organism:	Cloudy liquid
Active substance and concentration:	Benzalkonium Chloride
Product dilutions/concentrations:	1 Sachet in 600ml Hard Water
	(Volume of water adjusted from 750ml to allow for
	dilution during test)
Diluent used to dilute product:	Synthetic Hard Water

The test product was in satisfactory condition for testing when received.Date product received:07/06/21Test Date:17/06/21

Experimental Conditions:

Interfering substance:	Bovine Albumin (clean 0.3g/l)
Test temperature:	18 to 25 °C
Contact time:	5 Minutes
Test organisms:	Candida albicans ATCC 10231

Incubation temperature: $30^{\circ}C + - 1^{\circ}C$

Requirements of the Standard:

The test product shall demonstrate at least a 4 decimal logarithm (lg) reduction when tested in accordance with this standard under simulated dirty conditions.



Test Report General-Purpose Disinfectant Product BS EN 1650:2019

Conclusion:

For the product PDCCS9 A Surface Disinfectant, [Batch code N/A] the log reduction requirements as specified in EN 1650:2019 (4 lg within the relevant contact time) were met in clean conditions with a contact time of 5 minutes for Candida albicans.

Report authorised by:

Name: Position: Date: Dawn Mellors Technical Director 28/06/2021 <u>Test Report General-Purpose Disinfectant Product</u>



BS EN 1650:2019

Test Results:

Neutralisation Method Used:

Membrane filtration

Rinsing Liquid Used: N7



Test Report General-Purpose Disinfectant Product BS EN 1650:2019

Candida	albicans	ATCC 10231
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	Validation and controls									ef No	28490	
Valida	ation susp (<i>Nv</i> ₀)	ension		ental conc introl (A)	litions	Neutralizer control (B)		Method validation (C) Product conc: 1 Sachet ir			600ml Hard Water	
<i>Vc</i> 1	82	x =	<i>Vc</i> 1	50	x =	Vc 1	57	x =	Vc 1	58	x =	
Vc 2	79	80.5	Vc 2	49	49.5	Vc 2	45	51	Vc 2	52	55	
30 ≤	\overline{X} of Nv_0 : Yes	≤ 160?	\overline{X} of A is \ge	2 0.5 x X o Yes	of Nv ₀ ?	\overline{x} of B is	\overline{X} of B is $\ge 0.5 \times \overline{X}$ of Nv_0 ? Yes		\overline{x} of C	is ≥ 0.5 x X o Yes	of Nv _o ?	

Test suspension and test		N	<i>Vc</i> 1	Vc 2	X m 4.30E+07 ; lg N = 7.63
	Test suspension (N and N _o):	10 -5	>330	>330	$N_0 = N/10$; $\lg N_0 = 6.63$
	(10 and 10 0).	10 ⁻⁶	50	36	$6.17 \le \lg N_0 \le 6.70$? Yes
					x quotient = >5 and <15? N/A

Conc. of the active (%)	10 ^{-X}	Vc 1	Vc 2	$Na = \overline{X}$	la Ma	lg	R	Contact	Result	
	10	VCI	VCZ	Nu – X	lgNa	N ₀ =	6.63	time	Result	
1 Sachet in 600ml Hard Water	-1	<14	<14	1.40E+02	<2.15		>4.49	5 Minutes	Pass	



Company Name:	PVA Hygiene Ltd
Contact Name:	Jim Taylor
Contact Email:	technical@pva-hygiene.co.uk
Purchase Order No:	1554
Report Date:	16/03/2021
Melbec Ref Number: No. of Samples:	23556 1
Name of Test Product: Batch Number:	PDCCS9 A Surface Disinfectant N/A

MTF 5.10.94 Issue 1

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Sample Details:

Manufacture / Supplier:	PVA Hygiene Ltd
Product storage conditions:	Ambient
Appearance of the product (as supplied):	Clear colourless
Appearance of the product (after dilution):	Clear colourless
Active substance and concentration:	Benzalkonium Chloride
Product dilutions/concentrations:	Sachet in 750ml of water
Diluent used to dilute product:	Synthetic Hard Water
Incubation temperature:	36 [°] C±1 [°] C (24h)

The test product was in	satisfactory o	condition for testing when received.	
Date product received:	21/12/20	Test Date:	07/01/21

Experimental Conditions:

Interfering substance:	Bovine Albumin (clean 0.3g/l)
Test temperature:	18 to 25 degrees
Contact time:	5 Minutes
Test organisms:	Pseudomonas aeruginosa ATCC 15442
	Staphylococcus aureus ATCC 6538
	Escherichia coli ATCC 10536
	Enterococcus hirae ATCC 10541

Requirements of the Standard:

The test product shall demonstrate at least a 4 decimal logarithm (lg) reduction for bacteria and a at least a 3 decimal logarithm (lg) reduction for fungi when tested in accordance with this standard under simulated clean or dirty conditions. MTF 5.10.94 Issue 1



Conclusion:

The test product has met the requirements as specified in EN13697 for Pseudomonas aeruginosa, Staphylococcus aureus, Escherichia coli and Enterococcus hirae in clean conditions with a contact time of 5 minutes.

Testing carried out by:

Report authorised by:

Name:Danika WeatherburnPosition:Laboratory Manager

SMS

Name:Dawn MellorsPosition:Technical DirectorDate:16/03/2021

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Test Results:

Neutralisation Method Used:

Dilution neutralisation by pour plate

Neutraliser used N1

Viable Counts (Nc, Nd & Nts)

Nc is the mean log number of organisms per test surface of the water control at the end of the contact time Nd is the mean log number of organisms per test surface of the disinfectant test at the end of the contact time Nts is the mean number of organisms remaining on the test surface at the end of the test.

NC is the neutraliser control NT is the method validation

Log Reduction:

Log reduction (R) = LogNc - LogNd

MTF 5.10.94 Issue 1



Bacterial or Fungal Test Suspension (N) (cfu/disc)

		monas ae ATCC 1544	-	Staphylococcus aureus ATCC 6538			
Count	-7	>330	>330	-6	>330	>330	
Count	-8	47	42	-7	34	29	
Weighted Mean		4.45E+0	9	3.15E+08			
Lg		9.65			8.50		
6.57 <n<7.10< td=""><td colspan="4">- 6.90</td><td></td></n<7.10<>	- 6.90						
7.57 <n<8.10< td=""><td></td><td>8.05</td><td></td><td colspan="3">-</td></n<8.10<>		8.05		-			

	Esche	erichia col 10536	i ATCC	Enterococcus hirae ATCC 10541			
Count	-7	>330	>330	-6	>330	>330	
Count	-8	45	42	-7	34	31	
Weighted Mean		4.35E+0	9	3.25E+08			
Lg		9.64			8.51		
6.57 <n<7.10< td=""><td colspan="4">- 6.91</td><td>6.91</td><td></td></n<7.10<>	- 6.91				6.91		
7.57 <n<8.10< td=""><td></td><td>8.04</td><td></td><td colspan="4">-</td></n<8.10<>		8.04		-			

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Validation and Controls (Counts on Test Surfaces)

		Pseudom	onas aeru	ginosa A	TCC 15442	2	Staphylococcus aureus ATCC 6538					
	NT			NC		NT			NC			
Count	-2	>330	>330	-2	>330	>330	-3	>330	>330	-3	>330	>330
Count	-3	137	124	-3	180	150	-4	96	76	-4	103	93
Weighted Mean		1.31E+0	6	1.65E+06			8.60E+06			9.80E+06		
Lg		6.12		6.22			6.93			6.99		
NC - Nc (Not > +/- 0.3lg)	-		-0.19			-			-0.09			
NT - Nc (Not > +/- 0.3lg)		-0.29			-		-0.14			-		

		Esch	erichia co	li ATCC 1	0536		Enterococcus hirae ATCC 10541					
		NT NC				NT			NC			
Count	-3	>330	>330	-3	>330	>330	-3	>330	>330	-3	>330	>330
Count	-4	41	41	-4	38	26	-4	69	47	-4	55	42
Weighted Mean		4.10E+0	6	3.20E+06			5.80E+06			4.85E+06		
Lg	6.61		6.51			6.76			6.69			
NC - Nc (Not > +/- 0.3lg)	-		0.16			-			0.14			
NT - Nc (Not > +/- 0.3lg)		0.27		-			0.21			-		

MTF 5.10.94 Issue 1



Determination of Microbicidal Activity (Nd) and Water Control (Nc) (Count/Test Surface)

Pseudomonas aeruginosa ATCC 15442

10 [×]	Water Co	ntrol (Nc)	Test Proce	Test Procedure (Nd)			
10			Sachet in 75	0ml of water			
N	-	-	0 0				
-1	-	-	-	-			
-3	266	225	-				
-4	42	33		-			
Mean	2.57	E+06		-			
Lg	6.4	41	<0	.10			
Nts (count remaining on disc)	>1	00	()			
Log Reduction (R)			>6	.31			

Staphylococcus aureus ATCC 6538

10 [×]	Water Co	ontrol (Nc) Test Procedure		edure (Nd)
10			Sachet in 75	0ml of water
N	-	-	48	43
-1	-		0 0	
-3	>330	>330	-	
-4	120	120	-	
Mean	1.20	E+07	4.55E+02	
Lg	7.08		2.66	
Nts (count remaining on disc)	>100		()
Log Reduction (R)			4.	42

MTF 5.10.94 Issue 1

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Determination of Microbicidal Activity (Nd) and Water Control (Nc) (Count/Test Surface)

Escherichia coli ATCC 10536

10 [×]	Water Control (Nc)		Test Procedure (Nd)	
10			R	ſU
N	-	-	0	0
-1	-		-	-
-3	>330	>330	-	
-4	25	19	-	
Mean	2.20	E+06	-	
Lg	6.34 <0.10		.10	
Nts (count remaining on disc)	57		()
Log Reduction (R)			>6	.24

Enterococcus hirae ATCC 10541

10 [×]	Water Control (Nc)		Test Procedure (Nd)	
10			R	ſU
N	-	-	36	30
-1	-			-
-3	>330	>330	-	
-4	39	32	-	
Mean	3.55E+06		3.30E+02	
Lg	6.55 2.52		52	
Nts (count remaining on disc)	>100		()
Log Reduction (R)			4.	03

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Note:

Viable counts of 1-14 (below the lower limit) are expressed as <1.4 x 10² (<2.15 Log)

Viable counts of 0 are expressed as < 0.10 Log

Viable counts >330 for bacteria and yeasts and >165 for mould (higher than the upper limit) are expressed as > 3.3×10^5 (> $5.52 \log$) or > 1.65×10^5 (> $5.22 \log$) Nts counts of >100 are expressed as >100

Method Verification:

For Each Test:	
The mean counts used for calculation of N, Nc, Nd, NC and NT are between 14 and 330 for bacteria and yeasts and 14 and 165 for moulds	Yes
6.57≤N≤7.10 for bacteria in dirty conditions and clean conditions (except Pseudomonas aeruginosa) and for Candida albicans in clean conditions	Yes
7.57≤N≤8.10 for Pseudomonas aeruginosa in clean conditions	Yes
5.57≤N≤6.10 for Candida albicans in dirty conditions and Aspergillus brasiliensis	N/A
NC-Nc is not > ± 0.3 log	Yes
NT-Nc is not > ± 0.3 log	Yes
Nts is <100 cfu for active concentrations	Yes
Weighted mean quotient for N is 5≤N≤15	Yes
Nc is sufficiently high to demonstrate a 4 lg reduction for bacteria and a 3 lg reduction for fungi	Yes

The sample detailed in this report will be retained for 1 month after report date, unless otherwise requested.

The results on this report refer to the items tested only.

Sample description (name of product) and batch references (batch number) stated are as provided by the customer.

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End of test report

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<u>Test Report for</u> BS EN 13697:2015+A1:2019



Chemical disinfectants and antiseptics- Quantitative non-porous surface test for the evaluation of bacterial and/or fungicidal activity of chemical disinfectants used in food, industrial, domestic and institutional areas-Tested method and requirements without mechanical action (phase 2, step 2)

Company Name:	PVA Hygiene
Contact Name:	Jim Taylour
Contact Email:	technical@pvahygiene.com
Purchase Order No:	2076
Report Date:	23/12/2022
Melbec Ref Number: No. of Samples:	49508.2 1
Name of Test Product: Batch Number:	Everyday Virucidal Surface Disinfectant PDC N/A





Sample Details:

Manufacture / Supplier:	· PVA Hygiene
Product storage conditions:	Ambient - Keep powder samples dark and dry
Appearance of the product (as supplied):	White poweder
Appearance of the product (after dilution):	N/A
Active substance and concentration:	· Benzalkonium chloride
Product dilutions/concentrations:	2% wt/v solution
Diluent used to dilute product:	Synthetic Hard Water
Incubation temperature:	Bacteria: 35 to 38°C for 48+6h:
Product preparation:	Dissolve 1 x 15g sachet in 750ml water

The test product was in a	satisfactory condition f	for testing when received.	
Date product received:	14/12/22	Test Date:	21/12/22

Experimental Conditions:

Interfering substance:	Bovine Albumin (clean 0.3g/l)	
Test temperature:	19 to 21 °C	2% wt/v solution
Contact time:	1 minutes	
Test organisms:	Streptococcus pyogenes	

Deviations:

Page 2 of 7

EN13697 states incubation temperature of 36±1°C or 37±1°C. Melbec Microbiology Ltd method states 35°C - 38°C. The standard specifies testing at three product concentrations. Client requested one concentration only hence the methodology is based on EN13697.

The test organism was specified by the client.





Requirements of the Standard:

The test product shall demonstrate at least a 4 decimal logarithm (lg) reduction for bacteria and at least a 3 decimal logarithm (lg) reduction for fungi when tested in accordance with this standard under simulated clean or dirty conditions.

Conclusion:

For the product Everyday Virucidal Surface Disinfectant PDC, [Batch code: N/A] the log reduction requirements as specified in EN 13697:2015 (4 lg for bacteria within the relevant contact time) were met in clean conditions with a contact time of 1 minutes for Streptococcus pvogenes. pH of product, bacteria and interfering substance = 10.87

Report authorised by:

Name: Position: Date: Dawn Mellors Technical Director 23/12/2022

MTF 5.10.128 i4





Test Results:

Neutralisation Method Used:

Dilution neutralisation by pour plate

Neutraliser used N1

Viable Counts (Nc, Nd & Nts)

Nc is the mean log number of organisms per test surface of the water control at the end of the contact time Nd is the mean log number of organisms per test surface of the disinfectant test at the end of the contact time Nts is the mean number of organisms remaining on the test surface at the end of the test. NC is the neutraliser control NT is the method validation

Log Reduction:

Log reduction (R) = LogNc - LogNd





Bacterial or Fungal Test Suspension (N) (cfu/disc)

	Streptococcus pyogenes		
Count	-6	>330	>330
Count	-7	37	33
Weighted Mean	3.50E+08		
Lg		8.54	
6.57 <n<7.10< td=""><td></td><td>6.94</td><td></td></n<7.10<>		6.94	

Validation and Controls (Counts on Test Surfaces)

		Streptococcus pyogenes				
	NT NC					
Count	-3	>330	>330	-3	>330	>330
Count	-4	61	54	-4	91	68
Weighted Mean	5.75E+06 7.95E+06			6		
Lg		6.76 6			6.90	
NC - Nc (Not > +/- 0.3lg)	-			0.00		
NT - Nc (Not > +/- 0.3lg)	-0.14				-	





Determination of Microbicidal Activity (Nd) and Water Control (Nc) (Count/Test Surface)

Streptococcus pyogenes

10 [×]	Water Control (Nc)		Test Procedure (Nd) 2 %	
			2	%
Ν	-	-	0	0
-1	-	-	-	-
-2	-	-	-	-
-3	>330	>330	-	
-4	80	80	-	
Mean	8.00	E+06	-	
Lg	6.90		< 0.10	
Nts (count remaining on disc)	>1	00	0	
Log Reduction (R)			> 6.80	
			PA	SS





Note:

Viable counts of 1-14 (below the lower limit) are expressed as $<1.4 \times 10^2$ (<2.15 Log)

Viable counts of 0 are expressed as < 0.10 Log

Viable counts >330 for bacteria and yeasts and >165 for mould (higher than the upper limit) are expressed as > 3.3×10^5 (>5.52 log) or > 1.65×10^5 (>5.22 log) Nts counts of >100 are expressed as >100

Method Verification:

For Each Test:	
The mean counts used for calculation of N, Nc, Nd, NC and NT are between 14 and 330 for bacteria and yeasts and 14 and 165 for moulds	Yes
6.57≤N≤7.10 for bacteria in dirty conditions and clean conditions (except Pseudomonas aeruginosa) and for Candida albicans in clean conditions	Yes
7.57≤N≤8.10 for Pseudomonas aeruginosa in clean conditions	N/A
5.57≤N≤6.10 for Candida albicans in dirty conditions and Aspergillus brasiliensis in clean or dirty conditions	N/A
NC-Nc is not > ± 0.3 log	Yes
NT-Nc is not > ± 0.3 log	Yes
Nts is <100 cfu for active concentrations	Yes
Weighted mean quotient for N is 5≤N≤15	Yes
Nc is sufficiently high to demonstrate a 4 lg reduction for bacteria and a 3 lg reduction for fungi	Yes

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Sample description (name of product) and batch references (batch number) stated are as provided by the customer.

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End of test report



Company Name:	PVA Hygiene	
Contact Name:	Jim Taylour	
Contact Email:	technical@pva-hygiene.co.uk	
Purchase Order No:	1612	
Report Date:	20/05/2021	
Melbec Ref Number:	27538	
Name of Test Product:	Sachets PDCSS9A	
Batch Number:	n/a	



Test Report for Virology BS EN 14476:2013+A2:2019

Sample Details:

Product storage condition Product appearance: Active substance and co Product dilution prepara Product dilutions/conce	ncentration: tion: ntrations:	Ambient Sachets dissolved ADBAC Volume/Volume 15g, 12g, and 10g	sachets dissolved in 750 ml of water.
Cytotoxicity Reduction n	nethod:	MicroSpin S 400 H	R columns and Large volume plating
Incubation temperature			
The test product was in	satisfactory condition for testing whe	en received.	
Date product received:	30/04/21	Test Date:	14/05/21
Experimental Conditions: Interfering substance:	Bovine Albumin (clean 0.3g/l)		
Test temperature: Contact time:	20 +/- 1 °C 5 minutes		
Test organisms: Cell line identification: Cell culture media:	Vaccinia virus VR-1508 (Modified BHK-21 Clone 13 Dulbeco's minimum essential me		Serum

Requirements of the Standard:

The test product shall demonstrate at least a 4 decimal logarithm (lg) reduction when tested in accordance with this standard under simulated clean or dirty conditions.



Test Report for Virology BS EN 14476:2013+A2:2019

Conclusion:

For the product Sachets PDCSS9A, [Batch code: n/a] the log reduction requirements as specified in BS EN 14476:2013+A2:2019 (4 lg within the relevant contact time) were met in clean conditions with a contact time of 5 minutes for the 15g sachet.

Report authorised by:

SM

Name: Position: Date:

Dawn Mellors Technical Director 20/05/2021

All samples are tested as received and the condition on receipt is deemed to be satisfactory for testing unless client is informed otherwise. If an unsatisfactory sample is received and tested on instruction from the client comments are included on the report detailing this information. Results given for this may be invalid. Results detailed above relate only to the samples tested. Sample description and batch references stated are as provided by the customer. This test report shall not be reproduced except in full without the approval of Melbec Microbiology Ltd.



Test Report for Virology BS EN 14476:2013+A2:2019

Method

Test procedure

To determine the virucidal activity of the product, test virus is exposed to product dilutions for the required contact time and subsequently, the product is neutralised. The solution is then serially diluted and titrated on cell monolayers. The surviving virus tissue culture infective dose (TCID₅₀) is determined by the appearance of cytopathic effect (CPE) on the cells and is calculated using the Spearman-Kärber calculation.

Several controls are run alongside each test to validate the assay.

Titration of Virus control: The titration of the virus test suspension is determined at the start of the test and at the end of the test to determine its infectivity.

Reference for Virus Inactivation control: Formaldehyde is used instead of the test product, at 2 time points to demonstrate that the virus remains resistant to biocidal action at known concentrations.

Efficiency of Suppression: The test product is neutralised during the test, prior to the addition of test virus. Recovery of the test virus at it's original titre demonstrates effective product neutralisation.

Interference control: Cell are incubated with the test product for 1 hour and subsequently the test virus is added. Recovery of the test virus at it's original titre demonstrates that the presence of the product does prevent infection of the cells by the test virus, and thus does not interfere with quantification of virucidal activity.

Cytotoxicity: Both the product and formaldehyde are incubated with cells, without the addition of test virus, to determine if any morphological changes occur that may mirror CPE normally caused by virus. This ensures any CPE seen is a result of residual virus and not the product.



Test Report for Virology BS EN 14476:2013+A2:2019

Vaccinia virus VR-1508 (Modified Vaccinia Ankara)

	Test Results				
Contact time	5 minutes	Raw data	log TCID ₅₀ /ml	Log reduction	
Product	t (15g)	000000	3.50	4.83	
Product (12g)		666000	5.50	2.83	
Product (10g)		666600	6.50	1.83	
Virus Test	Start	06666660	8.33		
Suspension	Finish	06666640	640 8.33		

Inactivation control (0.7% Formaldehyde)				
Contact time Raw data $\begin{bmatrix} \log \\ TCID_{50}/ml \end{bmatrix}$ Log reduction				
15 mins	064400	5.17	3.17	

Formaldehyde cytotoxicity		
Raw data	000000	
Level of cytotoxicity	3.50	

Product neutralisation				
Raw data	Raw data log TCID ₅₀ /ml			
06666640	8.17	0.17		
Product cyto				
Raw data	Level of cytotoxicity			
0000000	3.50			

Product interference				
	Raw data log TCID ₅₀ /ml		Log reduction	
PBS	06666660	8.50	-0.17	
Test product	06666640	8.17		
Difference		0.33		

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Verification of the methodology

Result Summary	Log of TCID50	Average	Log Reduction	Criteria	met/not met
Titration of Virus Control (Start)	8.50	8.33			
Titration of Virus Control (End)	8.17	0.55			
Product (15g)	3.50		4.83	Log Reduction >= 4 Log	Met
Product (12g)	5.50		2.83	-	-
Product (10g)	6.50		1.83	Log Reduction <= 4 Log	Met
Reference test for virus inactivation (15 mins)	5.17		3.17	2.0>=Log reduction=<4.0	Met
Efficiency of Suppression	8.17		0.17	<=0.5 log of Average	Met
Inactivation Control (Product)	8.17		0.17	<=1.0 log of Average	Met
Inactivation Control (PBS)	8.50		-0.17	<=0.5 log of Average	N/A
Product Cytotoxicity	3.50				N/A

1) The titre of the test suspension is at least 10^8 TCID50 /ml or is sufficiently high to at least enable a titre reduction of 4 lg to verify the method: detectable titre reduction shall be at least 4 lg.

2) The difference between the logarithmic titre of the virus control and the logarithmic titre of the test organism in the reference inactivation test should be between -2,0 and <= -4,0 after 15 mins for the Vaccinia virus VR-1508 (Modified Vaccinia Ankara).

3) Cytotoxicity of the product test solution should not affect cell morphology and growth or susceptibility for the test organism in the dilutions of the test mixtures which are necessary to demonstrate a 4 lg reduction of the virus.

4) The product should not interfere with susceptibility of the cells to the test organism, the difference in the titre of the test suspension and the recovered titre of the interference control should be <1lg.

5) Control of efficiency for suppression of product activity (the difference to the test suspension shall be \leq 0,5 lg).

6) At least one concentration per test shall demonstrate a 4 lg or more reduction and at least one concentration shall demonstrate a lg reduction of less than 4.



Safety Data Sheet

According to GB and EU REACH and CLP Regulations Issue date: 20/03/2023 Revision date: 20/03/2023 Supersedes version of: 29/10/2021 Version: 1.2

SECTION 1: Identification of the substance/mixture and of the company/undertaking

1.1. Product identifier	
Product form Product name UFI Product code	 Mixture EVERYDAY VIRUCIDAL DISINFECTANT 0RXA-G6YM-430P-5GIM ((UFI Code for EU use)) DZ4:20, DZ4:2, DZ4:8
1.2. Relevant identified uses of the subst	ance or mixture and uses advised against
1.2.1. Relevant identified uses Main use category Use of the substance/mixture	: Professional use,Consumer use : DISINFECTANT/DETERGENT
1.2.2. Uses advised against Restrictions on use	: Not for Oral Consumption, Not for Direct Application to Food Stuffs
1.3. Details of the supplier of the safety of	lata sheet
Manufacturer PVA HYGIENE UNIT 6 Havyat Business Park Havyat Road BS40 5PA Bristol – United Kingdom T +44 (0)1934 862 859 sales@pva-hygiene.co.uk	
1.4. Emergency telephone number	

Emergency number

: 01934 862859 (Office Hours). For Immediate first aid advice in the UK call 111 This product is registered with NPIS in the UK.

SECTION 2: Hazards identification	
2.1. Classification of the substance or mixture	
Classification according to Regulation (EC) No. 1272/2008 [CLP] and	nd GB CLP Regulations
Skin corrosion/irritation, Category 1, Sub-Category 1B	H314
Serious eye damage/eye irritation, Category 1	H318
Hazardous to the aquatic environment – Acute Hazard, Category 1	H400

Hazardous to the aquatic environment – Acute Hazard, Category 1H400Hazardous to the aquatic environment – Chronic Hazard, Category 2H411Full text of H- and EUH-statements: see section 16H411

Adverse physicochemical, human health and environmental effects

In Use Solutions are Un-Classified for Physical and Health hazards.

2.2. Label elements

Labelling according to Regulation (EC) No	5. 1272/2008 [CLP]
Hazard pictograms (CLP)	
	GHS05 GHS09
Signal word (CLP)	: Danger
Contains	: Alkyl (C12-14) Dimethylbenzylammonium Choride; Alcohols C9-11, Ethoxylated
Hazard statements (CLP)	: H314 - Causes severe skin burns and eye damage.
	H410 - Very toxic to aquatic life with long lasting effects.
Precautionary statements (CLP)	: P264 - Wash hands thoroughly after handling.
	P273 - Avoid release to the environment.
	P280 - Wear eye protection, protective gloves.

Safety Data Sheet

According to GB and EU REACH and CLP Regulations

P301+P330+P331 - IF SWALLOWED: Rinse mouth. Do NOT induce vomiting.
P303+P361+P353 - IF ON SKIN (or hair): Take off immediately all contaminated clothing.
Rinse skin with water or shower.
P305+P351+P338 - IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.
P402+P404 - Store in a dry place. Store in a closed container.
P501 - Dispose of contents to national regulations.

2.3. Other hazards

This product does not contain any substances classifed as PBT

This product does not contain any substances clasified as vPvB.

Contains no PBT/vPvB substances ≥ 0.1% assessed in accordance with REACH Annex XIII

SECTION 3: Composition/information on ingredients

3.1. Substances

Not applicable

3.2. Mixtures

Name	Product identifier	%	Classification according to Regulation (EC) No. 1272/2008 [CLP] and GB CLP Regulations
sodium carbonate	CAS-No.: 497-19-8 EC-No.: 207-838-8 EC Index-No.: 011-005-00-2 REACH-no: 01-2119485498- 19	≥ 60 - < 70	Eye Irrit. 2, H319
Alkyl (C12-14) Dimethylbenzylammonium Choride	CAS-No.: 85409-22-9 EC-No.: 287-089-1 REACH-no: 01-2120754638- 42	≥ 15 – < 20	Skin Corr. 1B, H314 Eye Dam. 1, H318 Aquatic Acute 1, H400 (M=10) Aquatic Chronic 1, H410
Citric Acid Mono Hydrate	CAS-No.: 5949-29-1 EC-No.: 691-328-9 REACH-no: 01-2119457026- 42	≥5-<8	Eye Irrit. 2, H319
Alcohols C9-11, Ethoxylated	CAS-No.: 68439-46-3	≥ 0.5 – < 1.5	Acute Tox. 4 (Oral), H302 Eye Dam. 1, H318 Aquatic Chronic 2, H411
Benzododecinium Chloride	CAS-No.: 139-07-1 EC-No.: 205-351-5 REACH-no: 01-2120831693- 52_XXX	≥ 0.5 – < 1.5	Acute Tox. 4 (Oral), H302 Skin Corr. 1B, H314 Aquatic Acute 1, H400 (M=10) Aquatic Chronic 1, H410
Cetalkonium Chloride	-	≥ 0.1 – < 0.5	Acute Tox. 4 (Oral), H302 Acute Tox. 4 (Dermal), H312 Skin Corr. 1B, H314 Eye Dam. 1, H318 Aquatic Acute 1, H400

Full text of H- and EUH-statements: see section 16

Safety Data Sheet

According to GB and EU REACH and CLP Regulations

SECTION 4: First aid measures	
4.1. Description of first aid measures	5
First-aid measures general	: If medical advice is needed, have product container or label at hand. For immediate First Aid advice in the UK, dial 111. When it is safe to do so, remove the victim immediately fro the source of exposure. However, consideration should be given as to whether moving the victim will cause further injury.
First-aid measures after inhalation	 Remove person to fresh air and keep comfortable for breathing. If unconscious place in recovery position and seek medical advice.
First-aid measures after skin contact	: Wash skin with plenty of water. Take off contaminated clothing. If skin irritation occurs: Get medical advice/attention.
First-aid measures after eye contact	: Rinse cautiously with water for several minutes. Remove contact lenses, if present and east to do. Continue rinsing. If eye irritation persists: Get medical advice/attention.
First-aid measures after ingestion	: Do not induce vomiting. Rinse mouth thoroughly with water. Get medical attention. If unconscious place in recovery position and seek medical advice.
4.2. Most important symptoms and e	ffects, both acute and delayed
Symptoms/effects	 Neat product is corrosive to skin and eyes. Diluted product is Unclassified for health hazards.
Symptoms/effects after inhalation	: Unlikely route of exposure, but inhalation of dilute solution droplets may result in a sore throat.
Symptoms/effects after skin contact	: Causes severe burns.
Symptoms/effects after eye contact	: Causes serious eye burns.
Symptoms/effects after ingestion	: Unlikely route of exposure without deliberate abuse. If sachets are swallowed they may swell and could block the throat and GI tract. If Powder is ingested, irritation and burning t the mouth and GI tract may occur, a soapy taste may be reported. Ingestion of diluted solution is unlikely to cause long term harm, but a soapy taste may be reported together with mild irritation to the lips, throat and GI tract.

4.3. Indication of any immediate medical attention and special treatment needed

Rinse with plenty of water. Check for abrasion to the surface of the eye from powder particles.

SECTION 5: Firefighting measures			
5.1. Extinguishing media			
Suitable extinguishing media Unsuitable extinguishing media	: Use extinguishing agent suitable for surrounding fire. : Water.		
5.2. Special hazards arising from the substance or mixture			
Fire hazard Hazardous decomposition products in case of fire	The product is not flammable.On heating, irritating fumes may be produced.		
5.3. Advice for firefighters			
Protection during firefighting	: Do not attempt to take action without suitable protective equipment. Self-contained breathing apparatus. Complete protective clothing.		

SECTION 6: Accidental release measures			
6.1. Personal precautions, protective e	equipment and emergency procedures		
6.1.1. For non-emergency personnel			
Emergency procedures	: Ventilate spillage area. Avoid contact with skin and eyes.		
6.1.2. For emergency responders			
Protective equipment	: Do not attempt to take action without suitable protective equipment. For further information refer to section 8: "Exposure controls/personal protection".		

Safety Data Sheet

According to GB and EU REACH and CLP Regulations

6.2. Environmental precautions

Normal use solutions can be disposed to sewers and septic tanks. Large scale spillages or uncontrolled discharges into water systems must be reported to the relevant Environment Agency.

6.3. Methods and material for containment and cleaning up

Methods for cleaning up

: Collect and place spillage in suitable containers. Seal the containers and apply labelling to identify the material and hazards. For disposal see section 13 of this SDS. Dispose of via an authorised person/ licensed waste disposal contractor or by other suitable waste treatment techniques.

6.4. Reference to other sections

For further information refer to section 13. See sections 2,8,12,13 &14.

SECTION 7: Handling and storage		
7.1. Precautions for safe handling		
Precautions for safe handling Hygiene measures	Carefully comply with the instructions for use. Avoid contact with eyes.Always wash hands after handling the product.	
7.2. Conditions for safe storage, including any incompatibilities		
Storage conditions	: Store in a dry place. Store in a closed container.	
7.3. Specific end use(s)		

DISINFECTANT/DETERGENT.

SECTION 8: Exposure controls/personal protection

8.1. Control parameters

8.1.1 National occupational exposure and biological limit values

EVERYDAY VIRUCIDAL DISINFECTANT	
United Kingdom - Occupational Exposure Limits	
Remark	No exposure limits known for ingredients.

8.1.2. Recommended monitoring procedures

No additional information available

8.1.3. Air contaminants formed

No additional information available

8.1.4. DNEL and PNEC

No additional information available

8.1.5. Control banding

No additional information available

8.2. Exposure controls

8.2.1. Appropriate engineering controls

Appropriate engineering controls:

Ensure good ventilation of the work station.

8.2.2. Personal protection equipment

Personal protective equipment: Gloves. Safety glasses.

Safety Data Sheet

According to GB and EU REACH and CLP Regulations

Personal protective equipment symbol(s):



8.2.2.1. Eye and face protection

Eye protection:

Safety glasses. In normal use eye protection is not required. During manufacture and packing operations, eye protection is recommended. Refer to EN166 to select appropriate level of protection.

8.2.2.2. Skin protection

Hand protection:

During normal use gloves are not required. During manufacture and packing operations, the use of gloves with a breakthrough time >60 minutes is recommended. Refer to EN374 to select appropriate level of protection. Rubber and PVC gloves are recommended. NOTE:- Use of gloves is a good general hygiene practice.

8.2.2.3. Respiratory protection

Respiratory protection:

In case of insufficient ventilation, wear suitable respiratory equipment. Note:- This would be very unusual in normal use.

8.2.2.4. Thermal hazards

No additional information available

8.2.3. Environmental exposure controls

Environmental exposure controls:

Avoid release to the environment.

Other information:

The PPE indicated in this SDS is not a COSHH assessment. It represents the PPE that should be considered for the neat product at all stages of the products life cycle, including manufacture, packing, distribution, use and disposal. Use solutions are unclassified, but for these we recommend use of gloves as minimum PPE.

SECTION 9: Physical and chemical properties

9.1. Information on basic physical and chemical properties

— •••••••••		
Physical state	: So	
Appearance	: Po	wder.
Colour	: wh	nite.
Odour	: od	ourless.
Odour threshold	: No	o data available
рН	: No	o data available
pH solution	: 10	– 11 @1%
Relative evaporation rate (butylacetate=1)	: No	ot applicable.
Melting point	: No	ot applicable
Freezing point	: No	ot applicable
Boiling point	: No	ot applicable
Flash point	: No	ot applicable
Auto-ignition temperature	: No	ot applicable
Decomposition temperature	: No	ot applicable
Flammability (solid, gas)	: No	on flammable.
Vapour pressure	: No	ot applicable
Relative vapour density at 20°C	: No	ot applicable
Relative density	: 0.8	3 – 0.9
Solubility	: Co	mpletely soluble in water.
Partition coefficient n-octanol/water (Log Pow)	: No	data available
Viscosity, kinematic	: No	ot applicable
Viscosity, dynamic	: No	data available
Explosive properties	: Pro	oduct is not explosive.
Oxidising properties		t oxidising.
Explosive limits		ot applicable
•		

Safety Data Sheet

According to GB and EU REACH and CLP Regulations

9.2. Other information

VOC content

: Contains no VOC material.

SECTION 10: Stability and reactivity

10.1. Reactivity

The product is non-reactive under normal conditions of use, storage and transport.

10.2. Chemical stability

Stable under normal conditions.

10.3. Possibility of hazardous reactions

No dangerous reactions known under normal conditions of use.

10.4. Conditions to avoid

Store away from moisture in a closed container. Protect from sunlight.

10.5. Incompatible materials

Strong acids. Oxidizing agent. Do not mix with Bleach or products containing Sodium Hypochlorite, this could result in dangerous heating of the solution.

10.6. Hazardous decomposition products

Under normal conditions of storage and use, hazardous decomposition products should not be produced.

SECTION 11: Toxicological information		
11.1 Information on toxicological effects		
Acute toxicity (dermal) :	Not classified Not classified Not classified	
Alkyl (C12-14) Dimethylbenzylammonium Cho	oride (85409-22-9)	
LD50 oral rat	≈ 344 ml/kg	
LD50 dermal rat	> 2000 ml/kg	
Benzododecinium Chloride (139-07-1)	·	
ATE CLP (oral)	500 mg/kg bodyweight	
Cetalkonium Chloride		
ATE CLP (oral)	500 mg/kg bodyweight	
ATE CLP (dermal)	1100 mg/kg bodyweight	
Alcohols C9-11, Ethoxylated (68439-46-3)		
LD50 oral rat	300 – 2000 ml/kg	
LD50 dermal rat	> 2000 ml/kg	
ATE CLP (oral)	500 mg/kg bodyweight	
Skin corrosion/irritation :	Causes severe skin burns.	
Serious eye damage/irritation :	Causes serious eye damage.	
Respiratory or skin sensitisation :	Not classified	
Germ cell mutagenicity :	Not classified	
Carcinogenicity :	This mixture is not classified as a carcinogen.	

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Reproductive toxicity	This mixture has no reproductive/foetal harm classifications and is not expected to be a risk to expectant mothers.
STOT-single exposure	Not classified
STOT-repeated exposure	Not classified
Aspiration hazard	Not classified
EVERYDAY VIRUCIDAL DISINFECTANT	
Viscosity, kinematic	Not applicable
sodium carbonate (497-19-8)	
Viscosity, kinematic	Not applicable

SECTION 12: Ecological information

12.1. Toxicity

Ecology - general : Hazardous to the aquatic environment, short-term : (acute) Hazardous to the aquatic environment, long-term : (chronic) Not rapidly degradable	Normal use solutions of this product are not classified for environmental harm. Very toxic to aquatic life. Toxic to aquatic life with long lasting effects.
Alkyl (C12-14) Dimethylbenzylammonium Ch	oride (85409-22-9)
LC50 - Fish [1]	≈ 0.791 ml/l Rainbow Trout
EC50 - Crustacea [1]	≈ 0.0164 ml/l Water flea
EC50 72h - Algae [1]	≈ 0.00785 mg/l Green Algae
Alcohols C9-11, Ethoxylated (68439-46-3)	
LC50 - Fish [1]	1 – 10 mg/l
EC50 - Crustacea [1]	1 – 10 g/l
EC50 72h - Algae [1]	1 – 10 mg/l
12.2. Persistence and degradability	
EVERYDAY VIRUCIDAL DISINFECTANT	
Persistence and degradability	The Surfactants and Chelants used in this mixture are Biodegradable.
12.3. Bioaccumulative potential	
EVERYDAY VIRUCIDAL DISINFECTANT	
Bioaccumulative potential	Not expected to Bioaccumulate.
12.4. Mobility in soil	
EVERYDAY VIRUCIDAL DISINFECTANT	
Additional information	soluble in water
12.5. Results of PBT and vPvB assessment	

EVERYDAY VIRUCIDAL DISINFECTANT

This product does not contain any substances classifed as PBT This product does not contain any substances clasified as vPvB.

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12.6. Other adverse effects

No additional information available

SECTION 13: Disposal considerations

13.1. Waste treatment methods

Waste treatment methods

Sewage disposal recommendations

Disposal of this product must comply with local and national environmental legislation.Small volumes of use solution can be disposed of to sewage drains.

SECTION 14: Transport information

ADR	IMDG	ΙΑΤΑ	ADN	RID
14.1. UN number	l	I		I
UN 1759	UN 1759	UN 1759	UN 1759	UN 1759
14.2. UN proper shippin	g name			-
CORROSIVE SOLID, N.O.S. (Alkyl (C12-14) Dimethylbenzylammonium Choride ; Alcohols C9-11, Ethoxylated)	CORROSIVE SOLID, N.O.S. (Alkyl (C12-14) Dimethylbenzylammonium Choride ; Alcohols C9-11, Ethoxylated)	Corrosive solid, n.o.s. (Alkyl (C12-14) Dimethylbenzylammonium Choride ; Alcohols C9-11, Ethoxylated)	CORROSIVE SOLID, N.O.S. (Alkyl (C12-14) Dimethylbenzylammonium Choride ; Alcohols C9-11, Ethoxylated)	CORROSIVE SOLID, N.O.S. (Alkyl (C12-14) Dimethylbenzylammoniuu Choride ; Alcohols C9-11 Ethoxylated)
Transport document descr	iption	I		I
UN 1759 CORROSIVE SOLID, N.O.S. (Alkyl (C12- 14) Dimethylbenzylammonium Choride ; Alcohols C9-11, Ethoxylated), 8, II, (E)	UN 1759 CORROSIVE SOLID, N.O.S. (Alkyl (C12- 14) Dimethylbenzylammonium Choride ; Alcohols C9-11, Ethoxylated), 8, II	UN 1759 Corrosive solid, n.o.s. (Alkyl (C12-14) Dimethylbenzylammonium Choride ; Alcohols C9-11, Ethoxylated), 8, II	UN 1759 CORROSIVE SOLID, N.O.S. (Alkyl (C12- 14) Dimethylbenzylammonium Choride ; Alcohols C9-11, Ethoxylated), 8, II	UN 1759 CORROSIVE SOLID, N.O.S. (Alkyl (C1 14) Dimethylbenzylammoniur Choride ; Alcohols C9-11 Ethoxylated), 8, II
14.3. Transport hazard o	class(es)			
8	8	8	8	8
14.4. Packing group	1	Ι		Ι
II	П	II	II	II
14.5. Environmental haz	ards			
Dangerous for the environment: Yes	Dangerous for the environment: Yes Marine pollutant: Yes	Dangerous for the environment: Yes	Dangerous for the environment: Yes	Dangerous for the environment: Yes
•	•	$\frac{1}{1}$ (quantity of liquids \leq 5 litres of the state of the ADR regulation, se	•/	he environmentally
No supplementary informatio	on available			
4.6. Special precaution	s for user			
overland transport				
Classification code (ADR)	: C1	0 4		

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According to GB and EU REACH and CLP Regulations

According to GB and EU REACH and CLP Regulations	
Limited quantities (ADR)	: 1kg
Excepted quantities (ADR)	: E2
Packing instructions (ADR)	: P002, IBC08
Special packing provisions (ADR)	: B4
Mixed packing provisions (ADR)	: MP10
Portable tank and bulk container instructions (ADR)	
Portable tank and bulk container special provisions	: TP33
(ADR)	
Tank code (ADR)	: SGAN, L4BN
Vehicle for tank carriage	: AT
Transport category (ADR)	: 2
Special provisions for carriage - Packages (ADR)	: V11
Hazard identification number (Kemler No.)	: 80
Orange plates	
	80
	1750
	1759
Tunnel restriction code (ADR)	: E
EAC code	: 2X
Transport by sea	
Special provisions (IMDG)	: 274
Limited quantities (IMDG)	: 1 kg
Excepted quantities (IMDG)	: E2
Packing instructions (IMDG)	: P002
IBC packing instructions (IMDG)	: IBC08
IBC special provisions (IMDG)	: B21, B4
Tank instructions (IMDG)	: T3
Tank special provisions (IMDG)	: TP33
EmS-No. (Fire)	: F-A
EmS-No. (Spillage)	: S-B
Stowage category (IMDG)	: A
Properties and observations (IMDG)	: Causes burns to skin, eyes and mucous membranes.
Air transport	
PCA Excepted quantities (IATA)	: E2
PCA Limited quantities (IATA)	: Y844
PCA limited quantity max net quantity (IATA)	: 5kg
PCA packing instructions (IATA)	: 859
PCA max net quantity (IATA)	: 15kg
CAO packing instructions (IATA)	: 863
CAO max net quantity (IATA)	: 50kg
Special provisions (IATA)	: A3, A803
ERG code (IATA)	: 8L
, , , , , , , , , , , , , , , , , , ,	
Inland waterway transport	
Classification code (ADN)	: C10
Special provisions (ADN)	: 274
Limited quantities (ADN)	: 1 kg
Excepted quantities (ADN)	: E2
Equipment required (ADN)	: PP, EP
Number of blue cones/lights (ADN)	: 0
,	
Rail transport	
Classification code (RID)	: C10
Special provisions (RID)	: 274
Limited quantities (RID)	: 1kg
Excepted quantities (RID)	: E2
Packing instructions (RID)	: P002, IBC08
Special packing provisions (RID)	: B4
Mixed packing provisions (RID)	: MP10
Portable tank and bulk container instructions (RID)	: T3

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Portable tank and bulk container special provisions (RID)	:	TP33
Tank codes for RID tanks (RID) Transport category (RID)		SGAN, L4BN 2
Special provisions for carriage – Packages (RID) Colis express (express parcels) (RID)	-	W11 CE10
Hazard identification number (RID)	:	80

14.7. Transport in bulk according to Annex II of Marpol and the IBC Code

Not applicable

SECTION 15: Regulatory information

15.1. Safety, health and environmental regulations/legislation specific for the substance or mixture

15.1.1. EU-Regulations

REACH Annex XVII (Restriction List)

Contains no substance(s) listed on REACH Annex XVII (Restriction Conditions)

REACH Annex XIV (Authorisation List)

Contains no substance(s) listed on REACH Annex XIV (Authorisation List)

REACH Candidate List (SVHC)

Contains no substance(s) listed on the REACH Candidate List

PIC Regulation (Prior Informed Consent)

Contains no substance(s) listed on the PIC list (Regulation EU 649/2012 concerning the export and import of hazardous chemicals)

POP Regulation (Persistent Organic Pollutants)

Contains no substance(s) listed on the POP list (Regulation EU 2019/1021 on persistent organic pollutants)

Ozone Regulation (1005/2009)

Contains no substance(s) listed on the Ozone Depletion list (Regulation EU 1005/2009 on substances that deplete the ozone layer)

VOC Directive (2004/42)

VOC content

: Contains no VOC material.

Explosives Precursors Regulation (2019/1148)

Contains no substance(s) listed on the Explosives Precursors list (Regulation EU 2019/1148 on the marketing and use of explosives precursors)

Drug Precursors Regulation (273/2004)

Contains no substance(s) listed on the Drug Precursors list (Regulation EC 273/2004 on the manufacture and the placing on market of certain substances used in the illicit manufacture of narcotic drugs and psychotropic substances)

15.1.2. National regulations

GB REACH and CLP regulations. UK HSE EH40 Publication.

15.2. Chemical safety assessment

No chemical safety assessment has been carried out

SECTION 16: Other information

Indication of changes:

Inclusion of EU UFI code and additional comments in section 7.

Abbreviations and acronyms:	
ADN European Agreement concerning the International Carriage of Dangerous Goods by Inland Waterways	
ADR European Agreement concerning the International Carriage of Dangerous Goods by Road	

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According to GB and EU REACH and CLP Regulations

Abbreviations and acronyms:		
ATE	Acute Toxicity Estimate	
BCF	Bioconcentration factor	
BLV	Biological limit value	
BOD	Biochemical oxygen demand (BOD)	
COD	Chemical oxygen demand (COD)	
DMEL	Derived Minimal Effect level	
DNEL	Derived-No Effect Level	
EC-No.	European Community number	
EC50	Median effective concentration	
EN	European Standard	
IARC	International Agency for Research on Cancer	
ΙΑΤΑ	International Air Transport Association	
IMDG	International Maritime Dangerous Goods	
LC50	Median lethal concentration	
LD50	Median lethal dose	
LOAEL	Lowest Observed Adverse Effect Level	
NOAEC	No-Observed Adverse Effect Concentration	
NOAEL	No-Observed Adverse Effect Level	
NOEC	No-Observed Effect Concentration	
OECD	Organisation for Economic Co-operation and Development	
OEL	Occupational Exposure Limit	
РВТ	Persistent Bioaccumulative Toxic	
PNEC	Predicted No-Effect Concentration	
RID	Regulations concerning the International Carriage of Dangerous Goods by Rail	
SDS	Safety Data Sheet	
STP	Sewage treatment plant	
ThOD	Theoretical oxygen demand (ThOD)	
TLM	Median Tolerance Limit	
VOC	Volatile Organic Compounds	
CAS-No.	Chemical Abstract Service number	
N.O.S.	Not Otherwise Specified	
vPvB	Very Persistent and Very Bioaccumulative	
ED	Endocrine disrupting properties	

Full text of H- and EUH-statements:		
Acute Tox. 4 (Dermal)	Acute toxicity (dermal), Category 4	
Acute Tox. 4 (Oral)	Acute toxicity (oral), Category 4	
Aquatic Acute 1	Hazardous to the aquatic environment – Acute Hazard, Category 1	
Aquatic Chronic 1	Hazardous to the aquatic environment – Chronic Hazard, Category 1	

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According to GB and EU REACH and CLP Regulations

Full text of H- and EUH-statements:	
Aquatic Chronic 2	Hazardous to the aquatic environment – Chronic Hazard, Category 2
Eye Dam. 1	Serious eye damage/eye irritation, Category 1
Eye Irrit. 2	Serious eye damage/eye irritation, Category 2
H302	Harmful if swallowed.
H312	Harmful in contact with skin.
H314	Causes severe skin burns and eye damage.
H318	Causes serious eye damage.
H319	Causes serious eye irritation.
H400	Very toxic to aquatic life.
H410	Very toxic to aquatic life with long lasting effects.
H411	Toxic to aquatic life with long lasting effects.
Skin Corr. 1B	Skin corrosion/irritation, Category 1, Sub-Category 1B

Safety Data Sheet (SDS), EU

This information is based on our current knowledge and is intended to describe the product for the purposes of health, safety and environmental requirements only. It should not therefore be construed as guaranteeing any specific property of the product.