Regulation (EU) No 528/2012 concerning the making available on the market and use of biocidal products

PRODUCT ASSESSMENT REPORT OF A BIOCIDAL PRODUCT FAMILY FOR UNION AUTHORISATION APPLICATIONS

(submitted by the evaluating Competent Authority)



Contec IPA Product Family

Product type(s) 2 & 4

Propan-2-ol

Case Number in R4BP: BC-LA025582-58

Evaluating Competent Authority: UK CA

Date: 12/03/2019

Table of Contents

1	CONCLUS	SION	5
	1.1 SUM	MARY OF DECISIONS AND RESTRICTIONS	5
	1.1.1	Usage area	5
	1.1.2	Pest and application rate	
	1.1.3	Active substance details	
	1.1.4	Comparative assessment and authorisation	
		SSARY ISSUES ACCOUNTED FOR IN THE PRODUCT LABEL	
		JIREMENT FOR FURTHER INFORMATION	
2	ASSESSM	ENT REPORT	7
	2.1 SUM	MARY OF THE PRODUCT ASSESSMENT	7
	2.1.1	Administrative information	
	2.1.1.1	Identifier of the product family	
	2.1.1.2	Authorisation holder	
	2.1.1.3	Manufacturer(s) of the products of the family	7
	2.1.1.4	Manufacturer(s) of the active substance(s)	8
	2.1.2	Product family composition and formulation	9
	2.1.2.1	Identity of the active substance	9
	2.1.2.2	Candidate(s) for substitution	
	2.1.2.3	Qualitative and quantitative information on the composition of the biocidal product family	
	2.1.2.4	Information on technical equivalence	
	2.1.2.5	Information on the substance(s) of concern	
	2.1.2.6	Type of formulation	
	2.1.3	Hazard and precautionary statements	
	2.1.4	Authorised use(s)	
	2.1.4.1	Meta SPC 1 – Contec IPA Liquid Products	
	2.1.4.2	Meta SPC 2 – Contec IPA Wipes	
	2.1.5	General directions for use	
	2.1.5.1	Instructions for use	
	2.1.5.2	Risk mitigation measuresParticulars of likely direct or indirect effects, first aid instructions and emergency measures to protect the	
	2.1.5.3	ment	
	2.1.5.4	Instructions for safe disposal of the product and its packaging	
	2.1.5.4	Conditions of storage and shelf-life of the product under normal conditions of storage	
	2.1.6	Other information	
	2.1.7	Packaging of the biocidal product family	
	2.1.7	Documentation	
	2.1.8.1	Data submitted in relation to product application	
	2.1.8.2	Access to documentation	
	2.1.8.3	Similar conditions of use	
		SSMENT OF THE BIOCIDAL PRODUCT FAMILY	-
	2.2.1	Intended use(s) as applied for by the applicant	
	2.2.2	Physical, chemical and technical properties	
	2.2.3	Physical hazards and respective characteristics	
	2.2.4	Methods for detection and identification	
	2.2.5	Efficacy against target organisms	
	2.2.5.1	Function and field of use	
	2.2.5.1	Organisms to be controlled and products, organisms or objects to be protected	
	2.2.5.3	Effects on target organisms, including unacceptable suffering	
	2.2.5.4	Mode of action, including time delay	
	2.2.5.5	Efficacy data	
	2.2.5.6	Occurrence of resistance and resistance management	
	2.2.5.7	Known limitations	
	2.2.5.8	Evaluation of the label claims	63
	2.2.5.9	Relevant information if the product is intended to be authorised for use with other biocidal product(s)	63

2.2.6	Risk assessment for human health	64
2.2.6		
2.2.6	.2 Exposure assessment	67
2.2.6	.3 Risk characterisation for human health	81
2.2.7	Risk assessment for animal health	84
2.2.8	Risk assessment for the environment	84
2.2.8		
2.2.8	.2 Exposure assessment	88
2.2.8	.3 Risk characterisation	97
2.2.9	Measures to protect man, animals and the environment	102
2.2.10	Assessment of a combination of biocidal products	102
2.2.11	Comparative assessment	102
ANNEX	ES	103
3.1 LIS	ST OF STUDIES FOR THE BIOCIDAL PRODUCT FAMILY	103
3.2 Ot	JTPUT TABLES FROM EXPOSURE ASSESSMENT TOOLS	115
3.2.1	Human Health Exposure Output Tables	115
3.2.2	Environmental Exposure Output Tables	121
3.3 NE	EW INFORMATION ON THE ACTIVE SUBSTANCE	130
	2.2.6 2.2.7 2.2.8 2.2.8 2.2.8 2.2.9 2.2.10 2.2.11 ANNEX 3.1 LIS 3.2 Or 3.2.1 3.2.2 3.3 Ni 3.4 Re 3.5 Su 3.6 Co	2.2.6.1 Assessment of effects on Human Health 2.2.6.2 Exposure assessment 2.2.6.3 Risk characterisation for human health 2.2.7 Risk assessment for animal health 2.2.8 Risk assessment for the environment 2.2.8.1 Effects assessment on the environment 2.2.8.2 Exposure assessment 2.2.8.3 Risk characterisation 2.2.9 Measures to protect man, animals and the environment 2.2.10 Assessment of a combination of biocidal products 2.2.11 Comparative assessment ANNEXES 3.1 LIST OF STUDIES FOR THE BIOCIDAL PRODUCT FAMILY 3.2 OUTPUT TABLES FROM EXPOSURE ASSESSMENT TOOLS 3.2.1 Human Health Exposure Output Tables 3.2.2 Environmental Exposure Output Tables 3.3 NEW INFORMATION ON THE ACTIVE SUBSTANCE 3.4 RESIDUE BEHAVIOUR 3.5 SUMMARIES OF THE EFFICACY STUDIES 3.6 CONFIDENTIAL ANNEX

Changes history table

Applica tion type	ref MS/ eCA	Case number in the refMS	Decision date	Assessment carried out (i.e. first authorisation / amendment / renewal)	Chapte r/ page
	UK		18.12.2019	Initial assessment	
UA-APP	DE	BC- LA025582- 58	02.03.2022	Post Authorisation Requirements: long-term storage stability	1.3 / p. 6 2.2.2 / p. 32, 33, 35 3.1 / p. 115

1 CONCLUSION

The outcome of the assessment for Contec IPA Product Family is specified in the BPC opinion following discussions at the BPC-29 meeting of the Biocidal Products Committee (BPC). The BPC opinion is available from the ECHA website.

1.1 Summary of decisions and restrictions

It is concluded after evaluation that sufficient data have been provided to verify the outcome and conclusions, and permit authorisation of the biocidal product subject to the following conditions:

1.1.1 Usage area

User	Usage Area
Professional	Indoor – hard, non-porous surfaces in cleanrooms for biotechnology, pharmaceutical, manufacture of medical devices, healthcare industries and other critical life science applications; industrial food and feed preparation areas

1.1.2 Pest and application rate

Authorisation is granted for use against bacteria, mycobacteria and yeast.

Application rate:

Meta SPC 1 – Contec IPA Liquid Products: 50 ml product per m² of surface ensuring a 1 minute contact time for bacteria, myobacteria and yeast

Meta SPC 2 – Contec IPA Wipes: Two wipes per m² of surface ensuring a 1-minute contact time for bacteria and myobacteria and a contact time of 3 minutes for yeast.

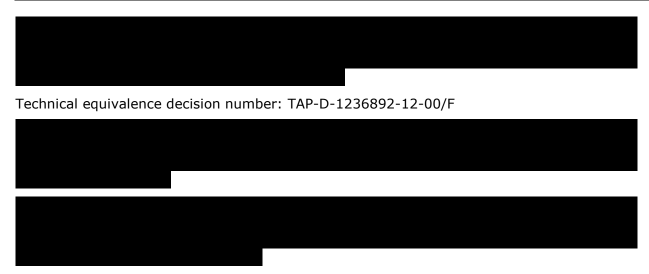
1.1.3 Active substance details

The concentration of the active substance propan-2-ol in the biocidal product is 70 % v/v (62.9 % w/w).

The minimum purity of the active substance propan-2-ol is 99 % w/w.



Technical equivalence decision number: TAP-D-1236889-07-00/F



1.1.4 Comparative assessment and authorisation

A comparative assessment is not required since propan-2-ol is not considered a candidate for exclusion in accordance with Article 5(1) or substitution in accordance with Article 10(1) of EU Regulation 528/2012.

1.2 Necessary issues accounted for in the product label

Wash hands and exposed skin before meals and after use.

Store in a cool, dry, well-ventilated place in original container.

For heavily soiled surfaces, cleaning prior to disinfection is required.

1.3 Requirement for further information

Storage stability data (Wipes):

Data showing satisfactory chemical and physical properties for the product family and their retention after ambient storage in the commercial packaging (PET/Foil/PE sachet and HDPE canister wipe packs) for the required shelf life must be provided. The specification proposed and properties tested should be in accordance with "Guidance on information requirements" (ECHA, Nov. 2014). All relevant properties should be determined prior to and after storage.

Addendum:

The required data on long-term storage stability for the product family have been submitted and assessed. The studies were considered acceptable.

2 ASSESSMENT REPORT

2.1 Summary of the product assessment

2.1.1 Administrative information

2.1.1.1 Identifier of the product family

Identifier	Country (if relevant)
Contec IPA Product Family	EU

2.1.1.2 Authorisation holder

Name and address of the	Name	Contec Cleanroom (UK) Limited
authorisation holder	Address	1 Park Row, Leeds, LS1 5AB, UK
Pre-submission phase started on	22 nd Decer	mber 2015
Pre-submission phase concluded on	18 th Februa	ary 2016
Authorisation number		
Date of the authorisation		
Expiry date of the authorisation		

2.1.1.3 Manufacturer(s) of the products of the family

Name of manufacturer	Contec Inc.
Address of manufacturer	525 Locust Grove, Spartanburg, SC 29303, USA
Location of manufacturing sites	525 Locust Grove, Spartanburg, SC 29303, USA

Name of manufacturer	Contec Cleanroom Technology (Suzhou) Co., Ltd. China
Address of manufacturer	17 Longyun Road, Suzhou Industrial Park, Suzhou, 215024, China
Location of manufacturing sites	17 Longyun Road, Suzhou Industrial Park, Suzhou, 215024, China

Name of manufacturer	Contec Cleanroom (UK) Ltd
	Unit 6A Wansbeck Business Park, Rotary Parkway, Ashington, Northumberland, NE63 8QW, UK
<u> </u>	Unit 6A Wansbeck Business Park, Rotary Parkway, Ashington, Northumberland, NE63 8QW, UK

Name of manufacturer	Flexible Medical Packaging
Address of manufacturer	Unit 8 Hightown, White Cross Industrial Estate, Lancaster, Lancashire, LA1 4XS
Location of manufacturing sites	Unit 8 Hightown, White Cross Industrial Estate, Lancaster, Lancashire, LA1 4XS

2.1.1.4 Manufacturer(s) of the active substance(s)

Active substance	Propan-2-ol
Name of supplier	Brenntag GmbH
Address of supplier	Stinnes Platz 1 45472 – Mülheim an der Ruhr Germany
Name of manufacturer	ExxonMobil
Address of manufacturer	Polderdijkweg 3B, B-2030 Antwerpen, Belgium
Location of manufacturing sites	Baton Rouge Chemical Plant (BRCP), Exxon Mobil Chemical Plant, 4999 Scenic Highway, Baton Rouge, Louisiana, 70897, USA

Active substance	Propan-2-ol
Name of supplier	Brenntag GmbH
Address of supplier	Stinnes Platz 1 45472 – Mülheim an der Ruhr Germany
Name of manufacturer	Shell Nederland Raffinaderij B.V.
Address of manufacturer	PO Box 2334, 3000 CH, Rotterdam, Netherlands
Location of manufacturing sites	Haven 3222, Vondelingenweg 601, 3196 KK, Vondelingenplaat, Netherlands

2.1.2 Product family composition and formulation

NB: the full composition of the product according to Annex III Title 1 should be provided in the confidential annex.

Does the product have the same identity and composition as the product evaluated in connection with the approval for listing of the active substance(s) on the Union list of approved active substances under Regulation No. 528/2012?

Yes \square This was a dummy product therefore new data have been submitted. No \square

2.1.2.1 Identity of the active substance

Main constituent(s)						
ISO name	Propan-2-ol					
IUPAC or EC name	Propan-2-ol					
EC number	200-661-7					
CAS number	67-63-0					
Index number in Annex VI of CLP	603-117-00-0					
Minimum purity / content	990 g/kg					
Structural formula	$HO \longrightarrow CH_3$					

2.1.2.2 Candidate(s) for substitution

The active substance is not a candidate for substitution.

2.1.2.3 Qualitative and quantitative information on the composition of the biocidal product family

Common name	IUPAC	Function	CAS number	EC number			Content (% w/w)	
	name		number		Min	Max	Min	Max
Propan-2-ol	Propan-2-ol	Active	67-63-0	200-661-7	70	70	62.9*	62.9*
		substance		200-001-7	(pure)	(pure)	(pure)	(pure)

^{*}Technical material content = 62.9 to 63.5% (Water content adjusted)

The full formulation composition details are contained within the Confidential Annex Section 3.6.1.

2.1.2.4 Information on technical equivalence



Technical equivalence decision number: TAP-D-1236889-07-00/F



Technical equivalence decision number: TAP-D-1236892-12-00/F

2.1.2.5 Information on the substance(s) of concern

No substances of concern have been identified in the product family.

2.1.2.6 Type of formulation

meta SPC 1: AL (any other liquid) - RTU solution/trigger spray meta SPC 2: AL (any other liquid) - RTU wipe

2.1.3 Hazard and precautionary statements¹

Classification and labelling of the products of the family according to the Regulation (EC) 1272/2008

Classification					
Hazard category	Flam. Liquid 2				
	Eye Irritant 2				
	STOT SE 3				
Hazard statement	H225: Highly flammable liquid and vapour.				
	H319: Causes serious eye irritation.				
	H336: May cause drowsiness or dizziness.				
Labelling					
Signal words	Danger				
Hazard statements	H225: Highly flammable liquid and vapour.				
	H319: Causes serious eye irritation				
	H336: May cause drowsiness or dizziness				
Precautionary statements	P101: If medical advice is needed, have product container or label at hand.				
	P102: Keep out of reach of children.				
	P210: Keep away from heat/sparks/open flames/hot surfaces. – No smoking.				
	P233: keep container tightly closed.				

¹ For micro-organisms based products: indication on the need for the biocidal product to carry the biohazard sign specified in Annex II to Directive 2000/54/EC (Biological Agents at Work).

10

P261: Avoid breathing vapours.

P264: Wash hands thoroughly after handling.

P271: use only outdoors or in a well-ventilated area.

P280: Wear protective gloves/eye protection/face protection

P303+P361+P353: IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water.

P304+P340: IF INHALED: Remove person to fresh air and keep comfortable for breathing.

P305+P351+P338: IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.

P312: Call a POISON CENTER/ doctor/.../if you feel unwell.

P337 + P313 If eye irritation persists: Get medical advice/attention

P370+P378: In case of fire: Use alcohol-resistant foam to extinguish.

P403+P235: Store in a well-ventilated place. Keep cool.

P405: Store locked up

P501: Dispose of contents/container in accordance with local regulations.

Note

Supplementary Hazard Information

EUH066: Repeated exposure may cause skin dryness or cracking.

Both the <u>assessment report</u> and the <u>Biocidal Products</u> <u>Committee (BPC) opinion</u> on propan-2-ol propose this additional label phrase based on local skin effects and reactions that have been described for human individuals exposed to formulations containing propan-2-ol or to propan-2-ol dilutions. As propan-2-ol is present at a concentration of 70 % in the formulation, it is the opinion of the UK CA that the additional labelling phrase should be applied as a precautionary measure.

The wipe itself is an inert polymer matrix and does not affect the classification. The user will be exposed directly to the liquid on the inert matrix.

P240, P241, P242, and P243 are not considered necessary as the products are not so volatile as to generate a hazardous atmosphere. See Section 2.2.3.

2.1.4 Authorised use(s)

2.1.4.1 Meta SPC 1 – Contec IPA Liquid Products

2.1.4.1.1 Use description

Table 1. Use # 1 - Professional use

Product Type	PT02 – Disinfectants and algaecides not intended for direct application to humans or animals PT04 – Food and feed area
Where relevant, an exact description of the authorised use	Disinfectant for use against bacteria, mycobacteria and yeast on hard, non-porous surfaces' surfaces in cleanrooms for biotechnology, pharmaceutical, manufacture of medical devices, healthcare industries and other critical life science applications; industrial food and feed preparation areas. Acceptable use temperature: room temperature (20±2°C)
Target organism (including development stage)	Bacteria Mycobacteria Yeast
Field of use	Indoor
Application method(s)	Spraying Wiping
Application rate(s) and frequency	50 ml product per m² of surface. 1-minute contact time for bacteria, mycobacteria and yeast
Category(ies) of users	Professional
Pack sizes and packaging material	HDPE trigger spray bottle – 0.5 – 1L HDPE capped refill bottle – 5L

2.1.4.1.1.1 Use-specific instructions for use

Use at room temperature. For visibly soiled surfaces, cleaning prior to disinfection is required.

Apply product to a suitable quality cleanroom wipe. Ensure the wipe is sufficiently and uniformly saturated before wiping the surface to be cleaned.

Ensure the surface is uniformly covered with the solvent then wipe to dry with a sterile cleanroom wipe.

Contact times: Spraying 1 minute for bacteria, mycobacteria and yeast

Wiping 1 minute for bacteria and mycobacteria 3 minutes for yeast.

Used wipes must be disposed of in a closed container.

2.1.4.1.1.2 Use-specific risk mitigation measures

Wash hands and exposed skin before meals and after use.

Avoid contact with eyes

2.1.4.1.1.3 Where specific to the use, the particulars of likely direct or indirect effects, first aid instructions and emergency measures to protect the environment

Likely direct or indirect adverse effects:

Headache, vertigo, hallucinations, respiratory depression, CNS depression or coma.

Severe irritation of the eyes and or ocular damage.

Nausea, vomiting, diarrhea and hemorrhagic gastritis.

Pulmonary aspiration hazard may induce pneumonitis, hypotension and hypoglycemia.

First aid measures:

Relocate the individual from the exposure source and remove any contaminated/spattered clothing articles.

Eye contact: IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing. If eye irritation persists: Get medical advice/attention

Skin contact; wash affected area with plenty of water and soap, No scrubbing.

In case of ingestion; Do NOT induce vomiting and never give anything by mouth to an impaired or unconscious individual; if the individual is unconscious place individual in left sideways (recovery) position with the head lowered and the knees bent.

Keep the individual calm and at rest, conserve body temperature and control breathing. If necessary check for pulse and initiate artificial respiration.

Take the individual to a healthcare center and bring packaging or label whenever possible.

NEVER LEAVE THE AFFECTED INDIVIDUAL UNATTENDED!

Advice for medical and healthcare personnel:

Monitor vital signs and provide symptomatic and supportive treatment.

Evaluate endoscopic procedure in case of ingestion.

Monitor glycaemia and ketones.

Ipecac use is contraindicated.

WHEN ASKING FOR MEDICAL ADVICE KEEP PACKAGING OR LABEL AT HAND AND CALL YOUR LOCAL POISON CONTROL CENTER

[INSERT LOCAL NUMBER HERE].

2.1.4.1.1.4 Where specific to the use, the instructions for safe disposal of the product and its packaging

Dispose of contents/container in accordance with local regulations.

Do not re-use empty container for any other purpose.

2.1.4.1.1.5 Where specific to the use, the conditions of storage and shelf-life of the product under normal conditions of storage

Store in a cool, dry, well-ventilated place in original container.

Keep away from sources of ignition.

Keep away from direct sunlight.

Keep container tightly closed.

Shelf-life: 2 years

2.1.4.2 Meta SPC 2 – Contec IPA Wipes

2.1.4.2.1 Use description

Table 2. Use # 1 - Professional use

Product Type	PT02 – Disinfectants and algaecides not intended for direct application to humans or animals PT04 – Food and feed area						
Where relevant, an exact description of the authorised use	Disinfectant for use against bacteria, mycobacteria and yeast on hard, non-porous surfaces' surfaces in cleanrooms for biotechnology, pharmaceutical, manufacture of medical levices, healthcare industries and other critical life science applications; industrial food and feed preparation areas						
Target organism (including development stage)	acteria lycobacteria east						
Field of use	Indoor						
Application method(s)	Wiping						
Application rate(s) and frequency	1-minute contact time for bacteria and mycobacteria 3-minute contact time for yeast						
Category(ies) of users	Professional						
Pack sizes and packaging material	Impregnated 100 % polypropylene wipes in: - HDPE canister with HDPE cap – 150 wipes (1.7 or 2.15 L) - PET/PE packet sealed with PET/PE flow wrap – 30, 40 or 50 wipes						
	 Impregnated 100 % polyester wipes in: HDPE canister with HDPE cap - 100 wipes (2.25 L) PET/PE packet sealed with PET/PE flow wrap - 20 wipes Impregnated 100 % knitted polyester wipes in: 						
	- PET/PE packet sealed with PET/PE flow wrap - 8, 10,						

20, 30 or 50 wipes

Impregnated 55 % cellulose / 45% polyester wipes in:

- HDPE canister with HDPE cap 100 wipes (2.25 L)
- PET/PE packet sealed with PET/PE flow wrap 24, 30,
 50 or 75 wipes

Impregnated 50 % rayon / 50% polyester wipes in:

- HDPE canister with HDPE cap – 700 wipes (11.4 L)

4.1.4.2.1.1 Use-specific instructions for use

Use at room temperature. For visibly soiled surfaces, cleaning prior to disinfection is required.

Ensure the surface is uniformly covered with the solvent then wipe to dry with a cleanroom wipe.

Used wipes must be disposed in a closed container

4.1.4.2.1.2 Use-specific risk mitigation measures

Wash hands and exposed skin before meals and after use.

Avoid contact with eyes

4.1.4.2.1.3 Where specific to the use, the particulars of likely direct or indirect effects, first aid instructions and emergency measures to protect the environment

Likely direct or indirect adverse effects:

Headache, vertigo, hallucinations, respiratory depression, CNS depression or coma.

Severe irritation of the eyes and or ocular damage.

Nausea, vomiting, diarrhea and hemorrhagic gastritis.

Pulmonary aspiration hazard may induce pneumonitis, hypotension and hypoglycemia.

First aid measures:

Relocate the individual from the exposure source and remove any contaminated/spattered clothing articles.

Eye contact: IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing. If eye irritation persists: Get medical advice/attention

Skin contact; wash affected area with plenty of water and soap, No scrubbing.

In case of ingestion; Do NOT induce vomiting and never give anything by mouth to an impaired or unconscious individual; if the individual is unconscious place individual in left sideways (recovery) position with the head lowered and the knees bent.

Keep the individual calm and at rest, conserve body temperature and control breathing. If

necessary check for pulse and initiate artificial respiration.

Take the individual to a healthcare center and bring packaging or label whenever possible.

NEVER LEAVE THE AFFECTED INDIVIDUAL UNATTENDED!

Advice for medical and healthcare personnel:

Monitor vital signs and provide symptomatic and supportive treatment.

Evaluate endoscopic procedure in case of ingestion.

Monitor glycaemia and ketones.

Ipecac use is contraindicated.

WHEN ASKING FOR MEDICAL ADVICE KEEP PACKAGING OR LABEL AT HAND AND CALL YOUR LOCAL POISON CONTROL CENTER

[INSERT LOCAL NUMBER HERE].

4.1.4.2.1.4 Where specific to the use, the instructions for safe disposal of the product and its packaging

Dispose of contents/container in accordance with local regulations.

Do not re-use empty container for any other purpose.

4.1.4.2.1.5 Where specific to the use, the conditions of storage and shelf-life of the product under normal conditions of storage

Store in a cool, dry, well-ventilated place in original container.

Keep away from sources of ignition.

Keep away from direct sunlight.

Keep container tightly closed.

Shelf-life: 2 years

2.1.5 General directions for use

2.1.5.1 Instructions for use

See section 2.1.4.1.1.1 or 2.1.4.2.1.1

2.1.5.2 Risk mitigation measures

See section 2.1.4.1.1.2 or 2.1.4.2.1.2

2.1.5.3 Particulars of likely direct or indirect effects, first aid instructions and emergency measures to protect the environment

See section 2.1.4.1.1.3 or 2.1.4.2.1.3

2.1.5.4 Instructions for safe disposal of the product and its packaging

See section 2.1.4.1.1.4 or 2.1.4.2.1.4

2.1.5.5 Conditions of storage and shelf-life of the product under normal conditions of storage

See section 2.1.4.1.1.5 or 2.1.4.2.1.5

2.1.6 Other information

Polypropylene, polyester, knitted polyester, 55 % cellulose / 45% polyester or 50 % rayon / 50% polyester wipes, 34-240 gsm, containing 5 – 38 ml product (2.75 – 20.9 g propan-2-ol)

The product contains propan-2-ol (CAS No: 67-63-0), for which an European reference value of 52.6 ppm for the professional user was agreed and used for the risk assessment of this product.

2.1.7 Packaging of the biocidal product family

Type of packaging	Size/volume of the packaging	Material of the packaging	Type and material of closure(s)	Intended user	Compatibility of the product with the proposed packaging materials
META-SPC	1				Acceptable. There were no
Bottle	0.5 – 5 L	HDPE	Capped	Professional	adverse interactions between the product and the HDPE trigger spray packaging for 18 weeks at 30°C. Packaging suitability will also be assessed following long term storage.
META-SPC	2		Acceptable. There were no		
Wipes - 10	0% polypropylene		adverse interactions between the liquid formulation and the		
Packet	30, 40 or 50 wipes	PET/PE	Flow wrap - PET/PE	Professional	HDPE packaging for 18 weeks at 30°C. These data can be
Canister	150 wipes, 1.7 or 2.15 L	HDPE	Cap - HDPE	Professional	extrapolated to support the impregnated wipe products. The formulation can be
Wipes - 10	0% polyester				considered similar to an
Packet	20 wipes	PET/PE	Flow wrap - PET/PE	Professional	aqueous based formulation due to the main component being an aliphatic alcohol;
Canister	100 wipes, 2.25 L	HDPE	Cap - HDPE	Professional	therefore extrapolation
Wipes - 10	0% knitted polyester				between packaging types is acceptable. In addition
Packet	8, 10, 20, 30 or 50 wipes	PET/PE	Flow wrap - PET/PE	Professional	propan-2-ol is known to have good resistance to a range of

Wipes - 55°	% cellulose / 45% po	lyester				
Packet	24, 30, 50 or 75 wipes	PET/PE	Flow wrap - PET/PE	Professional		
Canister	100 wipes, 2.25 L	HDPE	Cap - HDPE Professional			
Wipes – 50% rayon / 50% polyester						
Canister	700 wipes, 11.4 L	HDPE	Cap - HDPE	Professional		

The specification of the wipes is as follows:

Polypropylene wipes

Product size manufactured			Mass of active substance in product manufactured (g) ¹			Additional information		

¹ Density of propan-2-ol: 0.785 g/ml

Polyester wipes

Product size manufactured	Mass of active substance in product manufactured (g) ¹	Additional information	

¹ Density of propan-2-ol: 0.785 g/ml

Knitted polyester wipes

Product size manufactured			bstance in ctured (g) ¹	Additional information	

¹ Density of propan-2-ol: 0.785 g/ml

Cellulose/polyester wipes

Product size manufactured		Mass of active substance in product manufactured (g) ¹				Additional information	

¹ Density of propan-2-ol: 0.785 g/ml

Rayon/polyester wipes

Product size manufactured			Mass of act product ma		,	Additional information

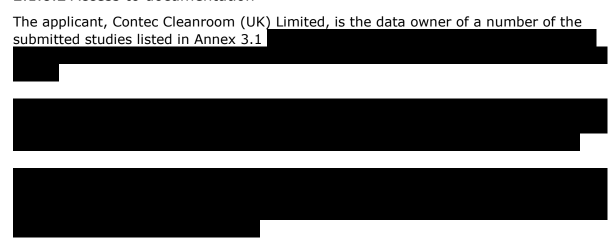
¹ Density of propan-2-ol: 0.785 g/ml

2.1.8 Documentation

2.1.8.1 Data submitted in relation to product application

No new data on the active substance has been submitted as part of this product family application. Please see Annex 3.1 for the list of studies used to support the product family.

2.1.8.2 Access to documentation



2.1.8.3 Similar conditions of use

The outcome of the Union Authorisation Pre-Submission Phase was communicated to the applicant, Contec Cleanroom (UK) Limited, by ECHA in communication number D(2016)0696 dated 18^{th} February 2016 and states the following:

The biocidal product family Contec IPA Product Family is deemed to be eligible for Union authorisation.

Reasons

Based on the information provided by the applicant, it appears that the application could meet the basic requirements of Article 42(1) of the Biocidal Products Regulation.

No objections were raised from either the Commission or the Member States Competent Authorities (MSCAs) as regards the eligibility of the prospective application for Union authorisation on the grounds that the biocidal product family Contec IPA Product Family falls outside of the scope of the Biocidal Products Regulation, or had been attributed the wrong product type, or that it would have non-similar conditions of use across the Union.

This document can be found in section 13 of the IUCLID dossier.

2.2 Assessment of the biocidal product family

2.2.1 Intended use(s) as applied for by the applicant

Table 3. Intended use # 1 - Ready to use liquid (Meta SPC 1 Contec IPA Liquid Products)

Product Type(s)	PT2: Disinfectants and algaecides not intended for direct application to humans or animals PT4: Disinfectants for food and feed area				
Where relevant, an exact description of the authorised use	General Disinfection of hard surfaces, Industrial and Institutional Areas.				
Target organism (including development stage)	Bacteria, yeast and mycobacteria				
Field of use	Disinfectant for use in cleanrooms for biotechnology, pharmaceutical, manufacture of medical devices, healthcare industries and other critical life science applications; industrial food and feed preparation areas				
Application method(s)	RTU solution				
Application rate(s) and frequency	For refill of the trigg	er spray			
Category(ies) of user(s)	Professional users				
Pack sizes and packaging material	Product name	Product code	Size/material		
	Contec Sterile 70% IPA	SBC570I	5L capped HDPE bottle (sterile)		
	Contec 70% IPA	FBC570I	5L capped HDPE bottle (non-sterile)		

Table 2. Intended use # 2 - Ready to use liquid trigger spray (Meta SPC 1 Contec IPA Liquid Products)

Product Type	PT2: Disinfectants and algaecides not intended for direct application to humans or animals PT4: Disinfectants for food and feed area
Where relevant, an exact description of the authorised use	General Disinfection of hard surfaces, Industrial and Institutional Areas.
Target organism(s) (including development stage)	Bacteria, yeast and mycobacteria
Field(s) of use	Disinfectant for use in cleanrooms for biotechnology, pharmaceutical, manufacture of medical devices, healthcare industries and other critical life science applications; industrial food and feed preparation areas.
Application method(s)	RTU spray
Application rate(s) and frequency	50 mL/m ² . As required and according to use instructions

Category(ies) of users	Professional users						
Pack sizes and packaging material	Product name	Product code	Size/material				
	Contec Sterile 70% IPA		1L HDPE trigger spray (sterile)				
	Contec Sterile 70% IPA		0.5L HDPE trigger spray(sterile)				
	Contec 70% IPA	FBT170I	1L HDPE trigger spray(non-sterile)				

Table 3. Intended use # 3 - Ready to use wipes (Meta SPC 2 Contec IPA Wipes)

Product Type	PT2: Disinfectants and algaecides not intended for direct application to humans or animals PT4: Disinfectants for food and feed area
Where relevant, an exact description of the authorised use	General Disinfection of hard surfaces, Industrial and Institutional Areas.
Target organism(s) (including development stage)	Bacteria, yeast and mycobacteria
Field(s) of use	Disinfectant for use in cleanrooms for biotechnology, pharmaceutical, manufacture of medical devices, healthcare industries and other critical life science applications; industrial food and feed preparation areas.
Application method(s)	RTU wipe
Application rate(s) and frequency	As required and according to use instructions. 5 mL – 26 mL liquid per wipe (depending on specific product – see below).
Category(ies) of users	Professional users
Pack sizes and packaging material	100% polypropylene wipes PROSAT MBPP PS-850 50 wipes (200 x 200mm): PET/PE laminate flow wrap, non-sterile, PROSAT MBPP PS-840IR 40 wipes (200 x 200mm): PET/PE laminate flow wrap, sterile, PROSAT MBPP PS-911 30 wipes (230 x 280mm): PET/PE laminate flow wrap, non-sterile, PROSAT MBPP PS-911EB 30 wipes (230 x 280mm): PET/PE laminate flow wrap, sterile, PROSAT MBPP PS-LPP-7030 50 wipes (410 x 430mm): PET/PE laminate flow wrap, non-sterile, PROSAT MBPP PS-LPP-7030IR 50 wipes (410 x 430mm): PET/PE laminate flow wrap, sterile, PROSAT MBPP PSPP0039 50 wipes (230 x 280mm): PET/PE laminate flow wrap, sterile, PROSAT MBPP PSPP0039 50 wipes (230 x 280mm): PET/PE laminate flow wrap, sterile, PROSAT MBPP PSPP0043

50 wipes (230 x 280mm): PET/PE laminate flow wrap, non-sterile, SATWipes MBPP SWPP0003
150 wipes (150 x 230mm):
1.7L HDPE canister, non-sterile,

Table 4. Intended use # 4 - Ready to use wipes (Meta SPC 2 Contec IPA Wipes)

Product Type	PT2: Disinfectants and algaecides not intended for direct application to humans or animals
Where relevant, an exact description of the authorised use	General Disinfection of hard surfaces, Industrial and Institutional Areas.
Target organism(s) (including development stage)	Bacteria, yeast and mycobacteria
Field(s) of use	Disinfectant for use in cleanrooms for biotechnology, pharmaceutical, manufacture of medical devices, healthcare industries and other critical life science applications; industrial food and feed preparation areas
Application method(s)	RTU wipe
Application rate(s) and frequency	As required and according to use instructions. 3 mL – 38 mL liquid per wipe (depending on specific product – see below).
Category(ies) of users	Professional users
Pack sizes and packaging material	100% polyester wipes PROSAT Delta PS-7030IR 20 wipes (230 x 230mm), PET/PE laminate flow wrap, sterile, SATWipes nonwoven polyester SAT-C3-7030 100 wipes (150 x 230mm), PET/PE laminate flow wrap, during storage, loaded into 2.25L HDPE canister during use, non-sterile, 100% knitted polyester wipes Spec-Wipe 4 115-0039 30 wipes (230 x 230mm), PET/PE laminate flow wrap, non- sterile, PROSAT Polynit Heatseal PS-HS9-7030 30 wipes (230 x 230mm), PET/PE laminate flow wrap, non- sterile, PROSAT EasyReach PSME0001 8 wipes (190 x 70mm), PET/PE laminate flow wrap, sterile, PROSAT Polynit Heatseal PSPS0047 30 wipes (300 x 300mm), PET/PE laminate flow wrap, non- sterile, PROSAT Polynit Heatseal PSPS0076 20 wipes (230 x 230mm), PET/PE laminate flow wrap, sterile, PROSAT Polynit Heatseal PSPS0091 20 wipes (300 x 300mm), PET/PE laminate flow wrap, sterile, PROSAT Polynit Heatseal PSPS0091 20 wipes (300 x 300mm), PET/PE laminate flow wrap, sterile, PROSAT Polynit Heatseal PSPS0091

10 wipes (230 x 230mm), PET/PE laminate flow wrap, sterile, PROSAT Polynit Heat eal PSWE0002 30 wipes (300 x 300mm), PET/PE laminate flow wrap, sterile, PROSAT Polynit Heatseal PSWE0003 10 wipes (300 x 300mm), PET/PE laminate flow wrap, sterile, 55% cellulose/45% polyester wipes Spec-Wipe 3 115-0034 75 wipes (230 x 230mm), PET/PE laminate flow wrap, nonsterile, PROSAT Sigma PSC20001 24 wipes (230 x 280mm), PET/PE laminate flow wrap, sterile, PROSAT Sigma PSC20002 24 wipes (230 x 280mm), PET/PE laminate flow wrap, nonsterile, PROSAT Theta PSC20005 50 wipes (230 x 280mm), PET/PE laminate flow wrap, sterile, PROSAT Theta PSC20006 50 wipes (230 x 280mm), PET/PE laminate flow wrap, nonsterile, PROSAT Theta PSC20009 30 wipes (230 x 230mm), PET/PE laminate flow wrap, sterile, PROSAT Theta PSC20010 50 wipes (230 x 230mm), PET/PE laminate flow wrap, nonsterile, PROSAT Theta PSCP0001 50 wipes (230 x 280mm), PET/PE laminate flow wrap, sterile, PROSAT Theta PSCS1010 50 wipes (230 x 230mm): PET/PE laminate flow wrap, nonsterile, PROSAT Theta PSCS1010IR 50 wipes (230 x 230mm): PET/PE laminate flow wrap sterile, SATWipes polyester/cellulose SAT-C1-7030 100 wipes (150 x 230mm), PET/PE laminate flow wrap, during storage, loaded into 2.25L HDPE canister during use, non-sterile, 50% rayon/50% polyester; 80% rayon/20% polyester wipes SATWipes polyester/rayon SWNW0013 700 wipes (175 x 230mm), 11.4L HDPE canister, non-sterile,

PT 2 & 4

2.2.2 Physical, chemical and technical properties

Contec IPA Product Family is a family of ready to use any other liquid products. Meta-SPC 1 contains liquid products supplied to the user as pouring bottles (for refill) and trigger sprays; and meta-SPC 2 contains liquid products supplied to the user as impregnated wipes. The liquid formulation in the products of meta SPC 1 is identical to the solution impregnated on the wipe material of the products in meta SPC 2; full details are provided in the Confidential Annex Section 3.6.1.

A full data package has been submitted and evaluated for the liquid formulation alone, however only an accelerated storage stability study entitled 'Cosmetic Regulation Stability Test Report' was submitted for the wipes. This report lacks the details required for biocidal product authorisation, nevertheless the data have been reported. No further data have been requested at this point as for the majority of properties it is possible to extrapolate data from the liquid product.

This was the representative formulation considered at active substance approval. However, as this was a 'model formulation' no physical, chemical or storage stability data were evaluated. The physical, chemical and storage stability data submitted to support this product family application are summarised in the following table.

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
Physical state at 20 °C and 101.3 kPa	Visual	Propan-2-ol 70 % liquid	A free flowing, uniform colourless liquid.	2016	- Acceptable
Colour at 20 °C and 101.3 kPa	assessment	Propan-2-ol 70 % wipes	Alcohol solution on a solid support, white wipe.	2016	
Odour at 20 °C and		Propan-2-ol 70 % liquid	Alcohol-like	2016	- Acceptable
101.3 kPa	Assessment	Propan-2-ol 70 % wipes	Alcohol	2016	
Acidity / alkalinity	CIPAC MT 75.3	Propan-2-ol 70 % liquid	pH = 6.52 (at 20 °C, neat)	2016	Acceptable. For the wipe products the experimental details (e.g. method and temperature) have not been provided. However as the liquid formulation of the

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
	No specified method	Propan-2-ol 70 % wipes	pH = 5.8	2016	impregnated wipes is identical to the liquid formulation alone, tested in Miller, R; 2016, it is possible to extrapolate the accelerated storage data from the liquid formulation to the impregnated wipes. Therefore further data has not been requested. As pH is within the range 4 – 10 acidity or alkalinity data are not required.
Relative density	OECD 109	Propan-2-ol 70 % liquid	0.87334 g/mL at 20°C 0.85928 g/mL at 40°C	2016	Acceptable, these data can also be extrapolated to the wipe products.
	CIPAC MT 46.3	Propan-2-ol 70 % liquid	Liquid samples were stored in HDPE containers for 18 weeks at 30 °C.		Acceptable storage period and temperature.
	Visual assessment		Appearance Initial: A free flowing, uniform colourless liquid. After: No change.		Acceptable, no change in product appearance.
	Weighing method		Weight change -3.24 g		Acceptable.
Storage stability test –	Validated GC-MSD method		Active substance content Initial: 69.67 % v/v After: 69.03 % v/v		Acceptable, a < 1 % decrease in active substance content was noted after storage.
accelerated storage 18 weeks at 30°C	CIPAC MT 75.3		pH (neat) Initial: 6.52 After: 6.07 pH (1 % dilution) Initial: 6.86 After: 6.84	2016	Acceptable.
	OECD 109		Density (at 20°C) Initial: 0.87334 g/mL After: 0.87244 g/mL		Acceptable.
	In-house method		Sprayability Initial: 11.1926 g		Acceptable, the mean trigger spray weight remains constant after accelerated storage.

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
			After: 11.4390 g		
Storage stability test – accelerated storage 12 weeks at 45°C	No specified methods	Propan-2-ol 70 % wipes	Polyester wipe samples were stored in PP sealed packs (50 wipes) for 12 weeks at 45 °C Appearance Initial: White wipe with alcohol odour. After: No change. pH Initial: 5.8 After: 6.2 Pack performance Initial: Peel back After: Satisfactory, no change. Pack weight change -4.3 g (Weight of pack = 226g) = -1.9%	2016	Acceptable storage temperature and period, these conditions exceed those specified in the BPR guidance. The limited data provided show acceptable results, however active substance content has not been determined. As the liquid formulation of the impregnated wipes is identical to the liquid formulation alone, tested in Miller, R; 2016, it is possible to extrapolate the accelerated storage data from the liquid formulation to the impregnated wipes. In addition, the % weight change for the 50 wipe pack is -1.9%, the product is 62.9% active substance and therefore even if all the loss (2%) was the active substance this would still be less than allowed 10%. Therefore the most pragmatic way forward, based on the data available on the liquid in the wipes, is to allow authorisation of the wipes (efficacy data indicate wipes are effective) and request ambient storage stability data on the wipes addressing active substance content, appearance and pH. The eCA considers this to be the best way forward, allowing authorisation of the wipes, with a safe guard of a data requirement.
Storage stability test – long term storage at ambient temperature and at 30°C	Visual assessment container	Propan-2-ol 70 % liquid	Ambient The container remained stable and unchanged after 24 months storage 30°C	DNA 3465	Acceptable, no change in the container after 24 months storage

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
for 24 months, in 0.75 litre HDPE trigger packs (sales			The container remained stable and unchanged after 24 months storage		
pack)	Visual assessment product		Ambient Initial = Free flowing uniform colourless liquid 6 months = Free flowing uniform colourless liquid with no sedimentation 12 months = Free flowing uniform colourless liquid with no sedimentation 24 months = Free flowing uniform colourless liquid with no sedimentation 30°C Initial = Free flowing uniform colourless liquid 6 months = Free flowing uniform colourless liquid with no sedimentation 12 months = Free flowing uniform colourless liquid with no sedimentation 24 months = Free flowing uniform colourless liquid with no sedimentation 24 months = Free flowing uniform colourless liquid with no sedimentation		Acceptable, no change in product appearance.
	Weight change (Weighing method) Active substance		Ambient 12 months = -0.04% 24 months = -0.08% 30°C 12 months = -0.2% 24 months = -0.4% Ambient Initial = 69.67%		Acceptable. Acceptable, a <3% decrease in active
					Acceptable, a <3% decrease in active substance content was noted after stora

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
	(Validated GC-MSD method)		12 months = 69.21% 24 months = 68.76%		
			30°C Initial = 69.67% 6 months = 68.46% 12 months = 68.37%		
			24 months = 68.01% Ambient (neat) Initial = 6.52 24 months = 6.08		
			Ambient (1% dilution) Initial = 6.86 24 months = 6.57		
	pH (CIPAC MT 75.3)		30°C (neat) Initial =6.52 6 months = 6.44 12 months = 6.46		Acceptable.
			24 months = 6.57 30°C (1% dilution) Initial = 6.86 6 months = 7 12 months = 6.99		
	Acidity (CIPAC MT 191)		24 months = 6.74 Ambient Initial = 0.0012% 24 months = 0.0012% 30°C		Acceptable.

Property Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
		Initial = 0.0012% 24 months = 0.0011%		
Sprayabili (In-house	- I	Spray Weight (10 sprays) Ambient Initial = 11.2g 6 months = 10.9g 12 months = 11.4g 24 months = 11.3-11.4g 30°C Initial = 11.2g 12 months = 11.1g 24 months = 11.2g Spray diameter Ambient Initial = 7.4x11.8cm (egg shape pattern) 24 months = 7.4x11cm to 8.7x12 cm (egg shape pattern) 30°C Initial = 7.4x11.8cm (egg shape pattern) 24 months = 8.2x11.6cm to (egg shape pattern) Nozzle Blockage Initial = No blockage of the nozzle occurred during use 24 months (part used pack) = No blockage of the nozzle occurred during use		Acceptable
Density		Ambient		Acceptable.

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
	(OECD 109)		Initial:0.8733g/ml at 20°C 24 months:0.8729g/ml at 20°C 30°C Initial = 0.8733g/ml at 20°C 6 months= 0.8735g/ml at 20°C		
			12 months = 0.873g/ml at 20°C 24 months:0.8731g/ml at 20°C Ambient		
	Surface Tension (EC A5)		Initial = 24.22 mNm ² at 20°C 22.59 mNm ² at 25°C 24 months = 23.87 mNm ² at 20°C 23.73 mNm ² at 25°C		Acceptable.
	Viscosity (OECD 114)		Ambient Initial = 0.1796 cm²/s at 20°C		Acceptable.
Storage stability test – long term storage at ambient temperature in the dark for 24 months (wipes) in two different sales packs	Active substance content: GC Analysis (FID)	Packaging A: Cellulose/Polyester nonwoven wipes (9"x11"), presaturated with Propan-2-ol 70 % (50 wipes in resealable pouch) Batch No.: 684804-0-0-1	Active substance content (% v/v) Packaging A: t = 0: 69.71 t = 12 months: 70.46 (+1.1 %) t = 24 months: 72.80 (+4.4 %) Packaging B: t = 0: 67.63 t = 12 months: 68.67 (+1.5 %) t = 24 months: 66.82 (-1.2 %)	2021 Report No.: 2019/121AM 2021 Report No.: 2019/122AM	DE CA (August 2021): Long term storage stability data for wipes (Meta-SPC 2) was submitted as postauthorisation requirement. The data is acceptable.

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
	pH: CIPAC MT 75.3	Packaging B: Cellulose/Polyester nonwoven wipes (6"x9"), presaturated with Propan-2-ol 70 % (100 wipes in foil package) Batch No.: 695202-0-0-1	pH (1 % aqueous solution, 20 °C) Packaging A: t = 0: 6.13 t = 12 months: 5.78 t = 24 months: 6.15 Packaging B: t = 0: 6.27 t = 12 months: 6.23 t = 24 months: 6.07		
	Weight loss and visual appearance		Weight loss Packaging A: -4.24 % after 24 months Packaging B: -0.03 % after 24 months No change in appearance of the product or the packaging observed during storage in any case.		
Storage stability test - low temperature stability test for liquids	CIPAC MT 39.3	Propan-2-ol 70 % liquid	Post seven days storage at 0°C ±2.0°C and 3 hours at room temperature (and all interim time points) the product remained a clear colourless solution with no signs of separation into oil, cream, sediment or crystallisation.	2016	Acceptable.
	No specified method	Propan-2-ol 70 % wipes	Limited freeze thaw data have been provided stating no change in product appearance after 12 weeks storage.	2016	As these data do not state the exact test protocol they are not suitable to support low temperature stability. As the liquid formulation of the impregnated wipes is identical to the liquid formulation alone, tested in Miller, R; 2016, it is possible to extrapolate the low temperature storage data from the liquid formulation to the

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments	
					impregnated wipes.	
Effects on content of the active substance and technical characteristics of the biocidal product - light	-	-	Propan-2-ol does not absorb ultraviolet radiation (No absorption > 290 nm). Consequently photolysis could not be a route of degradation of propan-2-ol, i.e. the stability of a propan-2-ol product will not be affected by luminous intensity.	Case	Acceptable.	
Effects on content of the active substance and technical characteristics of the biocidal product – temperature and humidity	-	-	-	-	See section 'Storage stability test – accelerated storage'.	
Effects on content of the active substance and technical characteristics of the biocidal product - reactivity towards container material	-	Propan-2-ol 70 % liquid/wipes	There was no significant change in container appearance and container weight following storage at 30°C for 18 weeks for the liquid products and 12 weeks at 45°C for the wipe products. This determination following ambient storage remains in progress.	2016	Acceptable based on the accelerated storage stability data. Further consideration to be made once the ambient temperature storage stability study is available.	
Particle size distribution, content of dust/fines, attrition, friability	CIPAC MT187 (Laser diffraction method)	Propan-2-ol 70 % liquid	MMAD = 267 μm	2019 No.6412	Acceptable, MMAD greater than 50 μm	
Physical compatibility	-	-	Not applicable as the products are not designed to be used in combination with other products and no incompatibilities have been identified.	Case	Acceptable.	
Chemical compatibility	-	-	Not applicable as the products are not designed to be used in combination with other products and no incompatibilities have been identified.	Case	Acceptable.	
Surface tension	EC A.5	Propan-2-ol 70 %	24.224 mN/m at 20°C		Acceptable.	

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
		liquid	22.592 mN/m at 25°C	2016	
Viscosity	OCED 114	Propan-2-ol 70 % liquid	Dynamic viscosity at 20°C 12.25-19.87 mPas (30-100 R.P.M.) Mean = 15.69 mPas Kinematic viscosity at 20°C 0.1796 cm²/s Dynamic viscosity at 40°C 10.11-14.14 mPas (50-100 R.P.M.) Mean = 11.80 mPas Kinematic viscosity at 40°C 0.1374 cm²/s	2016	Acceptable.
Sprayability	In-house method	Propan-2-ol 70 % liquid	The propan-2-ol product produced a mean trigger spray weight of 11.1926g from 10 sprays.	2016	Acceptable.
Spray diameter	In-house method	Propan-2-ol 70 % liquid	The propan-2-ol product produced a completely wet egg shaped central area with average dimensions of 5.6cm x 9.6cm and a circular shaped outer droplet area with average dimensions of 7.4cm x 11.8cm.		Acceptable.

Conclusion on the physical, chemical and technical properties of the product family

The physical, chemical and technical properties for Contec IPA Product Family are acceptable for the liquid formulation supplied to the user as trigger spray and impregnated wipe products. For the majority of properties data on the liquid formulation alone (meta-SPC 1) can be extrapolated to the impregnated wipes (meta-SPC 2) as the liquid formulations are identical. Therefore, the data provided are sufficient to support the product family. Two 24 month ambient storage stability studies on the wipe product were submitted post authorisation.

Accelerated and ambient temperature storage stability data for the liquid formulation alone were acceptable after 18 weeks at 30°C and 24 months at ambient temperature (and 30°C), therefore a shelf life of 2 years is supported for the product family. This was confirmed by long term storage stability data on the wipe product in 2021. The product labels state 'Store in a cool place.' therefore it is unlikely the products would be exposed to temperatures above 30°C. A low temperature storage stability study showed no significant change in the liquid product following storage at 0°C for seven days.

2.2.3 Physical hazards and respective characteristics

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
Explosives	Expert statement	-	The major constituent of the product is propan-2-ol, which is the active ingredient and is not classified under CLP regulation (EC) No 1272/2008 for explosivity and oxidising. None of the components are explosive or oxidizing and it is therefore concluded that the solution is unlikely to undergo rapid decomposition with the evolution of gases or release of heat and, therefore does not present a risk of explosion. It is also important to note that the product as a whole would not be expected to behave differently with regards to explosivity compared with individual constituents when combined.	2016	Acceptable. The product family is not classified as explosive.
Flammable liquids	EC A.9	Propan-2-ol 70 % liquid	The product had a flash point of 21.0°C and is classified as H225: Flammable liquid 2.	2016	Acceptable. The product family is classified as a category 2 flammable liquid (H225).
Self-reactive substances and mixtures	Expert statement	-	None of the individual constituents of the solution are classified as explosive, oxidising, or are organic peroxides. The chemical structures of all of the constituents were examined for the presence of characteristic groups associated with self-reactivity, none were present and therefore the product would not be expected to be susceptible to rapid, exothermic chemical reaction. The major constituent of the solution is propan-2-ol, which is the active ingredient and is only classified under CLP regulation (EC) No 1272/2008 for the physical chemical property of flammability. The composition of the formulation is fully known (see confidential annex). When mixed, no self-reacting properties would be expected. The product as a whole would not be expected to behave differently with regards to this CLP	2016	Acceptable. The product family is not classified as self-reactive.

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments	
			endpoint compared with the individual constituents. It is concluded that solution should not be considered for classification for self-reactivity.			
			Self-heating is generally a property of the substance when in bulk. As the preparation is a liquid, it should not be considered for the classification of self-heating as this property is generally only applicable to solids.			
Self-heating substances and mixtures	Expert statement	-	The composition of the formulation is fully known (see confidential annex). Propan-2-ol is not classified for any physical hazard under CLP regulation (EC) No 1272/2008, apart from flammability. The product is a liquid at room temperature and the boiling point is between 80°C and 100°C. As the melting point is below 160°C, this product should not be considered under this endpoint according to BPR guidance. In the absence of test results from screening procedures, the self-heating property has been assessed theoretically based on the characteristics of the components. It is	2016	Acceptable. The product family is not classified as self-heating.	
			concluded that the formulation does not contain any substances that contribute to the properties of self-heating, and therefore it would not be considered for the classification of self-heating material.			
Oxidising liquids	s Expert statement	na namae i ' i-	The major component of the solution is propan-2-ol, which is an alcohol and the active substance within the formulation. It is not classified under CLP regulation (EC) No 1272/2008 for oxidising properties and does not contain any oxidizing functional groups. It is classified for flammability under EU CLP but not for any other physical chemical CLP endpoint.	2016	Acceptable. The product family is not classified as	
			It can be concluded that the preparation itself will not be oxidising and will therefore be incapable of reacting exothermically with combustible materials. Additionally, as all of the constituents only contain oxygen that is chemically bonded to carbon or hydrogen, and no halogens are present,		oxidising.	

Property	Guideline and Method	Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
			according to BPR guidance the oxidising hazard class is not applicable to this formulation.		
Corrosive to metals	Expert statement	-	For a substance to be classified as Corrosive to Metals under CLP, it must corrode steel or aluminium at a rate of 6.5 mm per year or greater at 55°C. A material would have to be very corrosive to corrode metal at this rate, which would not be expected to apply to this product. Additionally the formulation would not be expected to have an extreme pH – the pH would be expected to be between 6.5 and 7.5, and the pKa of propan-2-ol in water is ≈ 17. The classification of the constituents indicates they do not possess general hazards related to the property of corrosivity. The major component of the solution is propan-2-ol, which functions as the active substance forming the bulk of the product. No functional groups present on any of the constituents of the solution would be considered strongly acidic or basic, and no halogens are present. The product as a whole would not be expected to behave differently with regards to this CLP endpoint compared with the combined individual constituents. It is therefore concluded that the formulation would not be considered for classification as a substance with "Corrosive to Metals" properties.	2016	Acceptable. The product is not classified as being corrosive to metals.
	OPPTS 830.6320	Propan-2-ol 70 % liquid	The test material had no apparent changes take place in the corrosivity to metals test when in contact with aluminium and HDPE. The test material, when in contact with zinc produced a dull white coating on the surface which was easily removed with wet and dry sandpaper. There was an average weight loss of 2.9 mg and is not considered significant. The test material, when in contact with copper dulled the surface which was easily removed with wet and dry sandpaper. There were no signs of corrosion to the copper and there was no significant weight change. No significant changes were observed to occur during this	2016	

		Purity of the test substance (% v/v)	Results	Reference	UK CA Comments
			corrosivity to metals test so the substance is not considered corrosive to metals.		
Auto-ignition temperatures of products (liquids and gases)	EC A.15	Propan-2-ol 70 % liquid	The product did not auto-ignite below 400°C and is therefore considered not highly auto-flammable.	2016	Acceptable. The product is not classified as auto-flammable.

Conclusion on the physical hazards and respective characteristics of the product family

Based on expert consideration of the composition, Contec IPA Product Family is considered not to be explosive, oxidising, self-reactive, self-heating or corrosive to metals. The flash point was measured to be 21.0°C therefore the product family is classified as category 2 flammable liquid (H225). The auto-ignition temperature >400°C.

2.2.4 Methods for detection and identification

Analytical methods for the active and impurities in the technical material

The sources of the active substance are considered technically equivalent to that considered for active substance approval. Letters of access to the list of endpoints agreed at active substance approval and the alternative active substance dossier (Annex II) have been provided where methods of analysis for the active substance and impurities have already been considered. Therefore, no further consideration is required at product authorisation.

Analytical methods for the active substance in the biocidal product family

Propan-2-ol content in the product family was determined by GC-MS as follows:

In duplicate approximately 0.05 g of liquid formulation were accurately weighed and transferred to a 50 mL volumetric flask and made to partial volume with deionised water containing 40 g/L sodium chloride. Each sample was sonicated for 5 minutes before being made to volume with deionised water containing 40 g/L sodium chloride. The samples were diluted 1:10 by transferring 1.0 mL of sample into a 20 mL headspace vial and making to 10 mL volume with deionised water containing 40 g/L sodium chloride. The samples were subsequently analysed using GC-MSD with Headspace Sampler. This gives a sample concentration of 0.1 mg/mL (100 mg/L) and therefore a propan-2-ol concentration of 70 mg/L.

The following spectroscopic conditions were noted:

Instrument: Shimadzu GC-MSD with HS-20 Headspace Sampler

Column: Rtx-5MS, $(30m \times 0.32mm \times 1.0\mu m)$

Temperatures:

Column: 30°C for 5 minutes, then 10°C/ minute to 200°C,

held for 3 minutes

Detector: Scan: 30 to 250 m/z

Carrier gas: Helium

Data Collection: GCMS Solutions

Retention Time: Approximately 5.4-5.7 minutes

Headspace Conditions:

UK CA

Cycle Time: 25 minutes

Shake Strength: 4/5

Oven Temperature: 70°C Loop Temperature: 150°C Transfer Line Temperature: 180°C

Analytical methods for the analysis of the product family as such including the active substance, impurities and residues

ΔηΖΙΝΤΑ	Analytical	Analytical Fortification range / Number of measurements	Linearity	Specificity	Recovery rate (%)			LOQ
	method				Range	Mean	RSD	
		70 mg/L (n=6) 5.0-200 mg/L	5.0-200 mg/L		96.3 - 104.0	100.6	2.7	
Propan-2-ol	GC-MS	5 mg/mL (n=6)	(7-285 % of nominal content) $R^2 = 0.9912$	No interference	98.36-102.9	100.4	1.499	5.0 %v/v

Precision: %RSD = 0.422 (at 69.45 % v/v, n=6)

Modified Horwitz = 1.41% - Method precision is less than the Horwitz value therefore the method is considered to be suitably precise.

Analytical methods for the monitoring of residues (soil, water, air, body fluids and tissues and food)

Methods of analysis for the determination of propan-2-ol residues in air have previously been evaluated at EU level. Methods for detection in body fluids and tissues methods are not required as the active substance is not considered toxic. Methods for detection in soil, water, food/feed of plant and animal origin are not available due to lack of exposure via the intended uses. Therefore, no further consideration is required at product authorisation.

While it is noted that the product family is intended for use as disinfectants for the treatment of food and feed area (PT4), residues of propan-2-ol are not expected within food or feed. The following case was made within the Assessment Report for propan-2-ol (PT4):

"Due to its high vapour pressure, the active substance evaporates completely within the time of application of the representative biocidal products, which are highly concentrated so that no transfer from treated hands or surfaces to food should occur. In the unlikely event that residue transfer does occur, the active substance will evaporate from the food before it is eaten."

Therefore, development of an analytical method for food or feedstuff is not necessary and scientifically not justified.

Conclusion on the methods for detection and identification of the product family

A GC-MS method has been developed and validated for the determination of the active substance in the biocidal product family. The method meets the EU criteria with respect to specificity, linearity, accuracy and precision as described in ECHA's 'Guidance on Regulation (EU) No 528/2012 concerning the making available on the market and use of biocidal products (BPR), Version 1.0 July 2013'.

A monitoring method for propan-2-ol residues in air has been evaluated and accepted at an EU level. For all other matrices monitoring methods are deemed scientifically unjustified due to a lack of exposure, this has been agreed at EU level.

2.2.5 Efficacy against target organisms

2.2.5.1 Function and field of use

Contec IPA Product Family is a family of PT 2 & 4 disinfectant products intended for professional indoor use. The product family includes two Meta SPCs, the first is a ready to use liquid to be applied by pouring (refill) or spraying, the second consists of ready to use wipes.

The active substance is propan-2-ol at a concentration of 70% v/v. The products of the family are intended for general disinfection of hard surfaces in industrial and institutional areas.

2.2.5.2 Organisms to be controlled and products, organisms or objects to be protected

The products of the family are intended to control bacteria, fungi and yeasts.

2.2.5.3 Effects on target organisms, including unacceptable suffering

The products of the family are sold as ready to use liquids or wipes to be applied as required.

For Meta SPC1 (liquid products) complete wetting of the surface to be treated is recommended. The specific application rate is 50 ml/m² and the products are to be applied as required.

For Meta SPC2 (wipes) the application rate is 2 wipes per square metre (between 5 and 38 ml of liquid per wipe, equivalent to between 2.7 and 20.9 g propan-2-ol) and complete wetting of the treated area is recommended. Typical use is one application every thirty minutes. On heavily soiled surfaces more than one wipe may be required.

The products work by denaturation, which leads to a loss of cellular activity, resulting in death of the cells.

No unacceptable suffering is foreseen as a result of the use of these products.

2.2.5.4 Mode of action, including time delay

The applicant has provided the following statement on the mode of action:

'Alcohols, such as propan-2-ol, exhibit an unspecific mode of action. It affects the cell membrane causing alteration of membrane fluidity and leakage, enters the cytoplasm and destroys the inner structure of the cell molecules and of the cytoplasm's proteins. It similarly interacts with corresponding viral structures. This process (referred to as denaturation) and the enzymes' coagulation leads to a loss of cellular activity resulting in the cell's death. There is no time delay to the toxic effect.'

The UK CA accepts the applicant's statement on the mode of action.

2.2.5.5 Efficacy data

Tests with 1-minute contact time or shorter have been highlighted.

Studies discussed during WG-EFF VII which were considered not valid/unacceptable by member states are greyed out.

		Experimental data on t	the efficacy of the biocidal product against tar	get organism(s)	
Function and field of use envisaged	Test substance	Test organism(s)	Test method/ Test system / concentrations applied / exposure time	Test results: effects	Reference
Disinfectant for use on hard surfaces in industrial and institutional areas.	Medipal Alcohol Wipes (liquid) 70% IPA	Bacteria: Pseudomonas aeruginosa Escherichia coli Staphylococcus aureus Enterococcus hirae	The EN 1276 standard protocol was followed. Phase 2, Step 1 test. Contact time: 1 minute. Test temperature: 20°C. Interfering substance: 3.0 g/l Bovine Serum Albumin (BSA) to represent dirty conditions. Concentration: Ready to use (RTU).	For all of the test organisms, a reduction greater than 5 log reduction (the pass criterion for the standard) was observed.	<mark>2014a</mark>
	Alcohol Spray (FH55) 70% IPA	Bacteria: Pseudomonas aeruginosa Escherichia coli Staphylococcus aureus Enterococcus hirae	The EN 1276 standard protocol was followed. Phase 2, Step 1 test. Contact time: 1 minute. Test temperature: 10°C. Interfering substance: 0.3 % Bovine Serum Albumin (BSA) to represent dirty conditions and 0.03 % Bovine Serum Albumin (BSA) to represent clean conditions. Concentrations: 10 %, 50 % and 80 %.	For all of the test organisms, a reduction greater than 5 log reduction (the pass criterion for the standard) was observed with the 80 % concentration.	2017a
	70% IPA	Bacteria:	The EN 13697 standard protocol was followed.	For all of the test organisms, a	

Wipes Solution	Pseudomonas aeruginosa Escherichia coli Staphylococcus aureus Enterococcus hirae	Phase 2, Step 2 test. Interfering substance: 3.0g/l bovine albumin (dirty) Contact time: 30 s Temperature: 20°C Concentration: Ready to use (RTU)	reduction greater than 4 log reduction (the pass criterion for the standard) was observed.	2013a
FH55 70% v/v IPA clear liquid	Bacteria: Pseudomonas aeruginosa Escherichia coli Staphylococcus aureus Enterococcus hirae	The EN 13697 standard protocol was followed. Phase 2, Step 2 test. Interfering substance: 0.3% bovine albumin (dirty) Contact time: 1 minute. Temperature: 4°C Concentration: Ready to use (RTU)	For all of the test organisms, a reduction greater than 4 log reduction (the pass criterion for the standard) was observed.	2016
	Bacteria: Staphylococcus aureus	The EN 16615 standard protocol was followed. Phase 2, Step 2 test. Concentrations tested: 100%, 0.1% Interfering substance: 0.3g/l bovine albumin (clean) Contact time: 1 minute Temperature: 20°C	For the test organism a reduction greater than 5 log reduction and less than 50 cfu on test fields 2 to 4 (the pass criterions for the standard) were observed.	2016a
Contec Meltblown Polypropylene	Bacteria: Staphylococcus aureus	The EN 16615 standard protocol was followed. Phase 2, Step 2 test.	For the test organism a reduction greater than 5 log reduction and less than 50 cfu	2016b

	70/30 IPA PS- 850		Concentrations tested: 100%, 0.1%. Interfering substance: 0.3 g/l bovine albumin (clean) Contact time: 1 minute Temperature: 20°C Concentrations tested: 100%, 0.1%	on test fields 2 to 4 (the pass criterions for the standard) were observed.	
A A	Alcohol Wipes	Bacteria: Pseudomonas aeruginosa Staphylococcus aureus Enterococcus hirae	The EN 13727 standard protocol was followed. Phase 2, Step 1 test. Contact time: 1 minute Test temperature: 20°C Interfering substance: 3.0 g/l Bovine Serum Albumin (BSA) and 3.0 ml/l sheep erythrocytes to represent dirty conditions. Concentration: Ready to use (RTU).	For all of the test organisms, a reduction greater than 5 log reduction (the pass criterion for the standard) was observed.	<mark>2014b</mark>
	70% IPA Wipes Solution	Bacteria: Pseudomonas aeruginosa Staphylococcus aureus Enterococcus hirae	The EN 14561 standard protocol was followed. Phase 2, Step 2 test. Contact time: 30 s. Test temperature: 20°C Interfering substance: 3.0 g/l Bovine Serum Albumin (BSA) and 3.0 ml/l sheep erythrocytes to represent dirty conditions. Concentration: Ready to use (RTU).	For all of the test organisms, a reduction greater than 5 log reduction (the pass criterion for the standard) was observed.	<mark>2013b</mark>
7	70% IPA	Bacteria:	The EN 14561 standard protocol was followed.	For both of the test organisms, a	

Wipes Solution	Staphylococcus aureus (Methicillin-resistant) Enterococcus faecium (Vancomycin-resistant)	Phase 2, Step 2 test. Contact time: 30 s. Test temperature: 20°C Interfering substance: 3.0 g/l Bovine Serum Albumin (BSA) and 3.0 ml/l sheep erythrocytes to represent dirty conditions. Concentration: Ready to use (RTU).	reduction greater than 5 log reduction (the pass criterion for the standard) was observed.	2013c
Medipal Alcohol Wipes (liquid) 70% IPA	Mycobacteria: Mycobacterium avium Mycobacterium terrae	The EN 14348 standard protocol was followed. Phase 2, Step 1 test. Contact time: 1 minute Test temperature: 20°C Interfering substance: 3.0 g/l Bovine Serum Albumin (BSA) and 3.0 ml/l sheep erythrocytes to represent dirty conditions. Concentration: Ready to use (RTU).	For both of the test organisms, a reduction greater than 4 log reduction (the pass criterion for the standard) was observed.	2014
70% IPA Wipes Solution	Mycobacteria: Mycobacterium avium Mycobacterium terrae	The EN 14563 standard protocol was followed. Phase 2, Step 2 test. Contact time: 1 minute Test temperature: 20°C Interfering substance: 0.3 g/l Bovine Serum Albumin (BSA) to represent clean conditions. Concentrations: 0.1 %, 50 % and 100 %.	For both of the test organisms, a reduction greater than 4 log reduction (the pass criterion for the standard) was observed with both the 50 % and 100 % concentrations.	2017
Medipal	Fungi:	The EN 1650 standard protocol was followed.	For C. albicans a reduction	

Alcohol Wipes (liquid) 70% IPA	Aspergillus niger Yeast: Candida albicans	Phase 2, Step 1 test. Contact time: 1 minute Test temperature: 20°C Interfering substance: 3.0 g/l Bovine Serum Albumin (BSA) to represent dirty conditions. Concentration: Ready to use (RTU).	greater than 4 log reduction (the pass criterion for the standard) was observed. For A. niger, the reduction (IgR < 3.19) was less than 4 log reduction (the pass criterion for the standard). Therefore, it failed the test and did not possess sufficient fungicidal activity against this microorganism.	2014c
Alcohol Spray (FH55) 70% IPA	Yeast: Candida albicans	The EN 1650 standard protocol was followed. Phase 2, Step 1 test. Contact time: 15 minutes. Test temperature: 10°C Interfering substance: 0.3 % Bovine Serum Albumin (BSA) to represent dirty conditions and 0.03 % Bovine Serum Albumin (BSA) to represent clean conditions. Concentrations: 10 %, 50 % and 80 %.	For <i>C. albicans</i> a reduction greater than 4 log reduction (the pass criterion for the standard) was observed.	2017c
PAL Clinical Wipes Solution 70% IPA	Fungi: Aspergillus niger Yeast: Candida albicans	The EN 13624 (surface and instrument disinfection in the medical area) standard protocol was followed. Phase 2, Step 1 test. Contact time: 60 minutes Test temperature: 20°C Interfering substance: 0.3 g/l Bovine Serum Albumin (BSA) to represent clean conditions.	For <i>C. albicans</i> a reduction greater than 4 log reduction (the pass criterion for the standard) was observed with both the 50 % and 80 % concentrations. For <i>A. niger</i> , the reductions observed were less than 4 log reduction (the pass criterion for the standard) at all concentrations tested. The log reduction at 80 % was 3.95. Therefore, it failed the test and	2006

		Concentrations: 25 %, 50 % and 80 % (v/v).	did not possess sufficient fungicidal activity against this microorganism.	
		The EN 13624 (surface and instrument disinfection in the medical area) standard protocol was followed.		
70 % IPA	Yeast:	Phase 2, Step 1 test. Contact time: 1 minute	For <i>C. albicans</i> a reduction greater than 4 log reduction (the	
Wipe Solutions	Candida albicans	Test temperature: 20°C	pass criterion for the standard) was observed with both the 50 % and 100 % concentrations.	2016c
		Interfering substance: 0.3 g/l Bovine Serum Albumin (BSA) to represent clean conditions.		
		Concentrations: 0.1 %, 50 % and 100 %.		
		The EN 16615 standard protocol was followed.	Against all bacteria species	
	Bacteria:	Phase 2, Step 2 test.	tested a ≥5.0 log reduction was observed together with ≤50 cfu	
Sample A	Pseudomonas aeruginosa Staphylococcus aureus	Interfering substance: 0.3g/l bovine albumin (clean)	in test fields 2 to 4, meeting the pass criterion of the standard.	
TX1130 wipes	Enterococcus hirae Yeast:	Contact time: 5 minutes	Against <i>C. albicans</i> a reduction of ≥4 was observed together	2015a
(IPA 70%)	Candida albicans	Temperature: 20°C	with ≤50 cfu in test fields 2 to 4, meeting the pass criterion of the	
		Concentrations tested: 100%, 0.1%	standard.	
	Bacteria:	The EN 16615 standard protocol was followed.	Against all bacteria species tested a ≥5.0 log reduction was	
Sample C CTB 40 wipes	Pseudomonas aeruginosa Staphylococcus aureus	Phase 2, Step 2 test.	observed together with ≤50 cfu in test fields 2 to 4, meeting the	
	Enterococcus hirae	Interfering substance: 0.3g/l bovine albumin (clean)	pass criterion of the standard.	2015b
(IPA 70%)	Yeast: Candida albicans	Contact time: 5 minutes	Against <i>C. albicans</i> a reduction of ≥ 4 was observed together with ≤ 50 cfu in test fields 2 to 4,	

		Temperature: 20°C	meeting the pass criterion of the standard.	
		Concentrations tested: 100%, 0.1%		
Contec 70% IPA ProSat Sterile Polynit Heatseal IPA 70% PSWE0001	Bacteria: Pseudomonas aeruginosa Staphylococcus aureus Enterococcus hirae Yeast: Candida albicans	The EN 16615 standard protocol was followed. Phase 2, Step 2 test. Interfering substance: 3.0g/l bovine albumin (dirty) Contact time: 1 minute (bacteria), 3 minutes (yeast) Temperature: 20°C Concentrations tested: 100%, 0.1%	Against all bacteria species tested a ≥5.0 log reduction was observed together with ≤50 cfu in test fields 2 to 4, meeting the pass criterion of the standard. Against <i>C. albicans</i> a reduction of ≥4 was observed together with ≤50 cfu in test fields 2 to 4, meeting the pass criterion of the standard.	<mark>2016d</mark>
Contec Prosat Sterile Cell/Poly nonwoven 70/30 IPA PSC 2001	Staphylococcus aureus Enterococcus hirae	The EN 16615 standard protocol was followed. Phase 2, Step 2 test. Concentrations tested: 100%, 0.1%. Interfering substance: 3.0g/l bovine albumin (dirty) Contact time: 1 minute (bacteria), 3 minutes (yeast) Temperature: 20°C	The log reduction was sufficient against <i>P. aeruginosa</i> and <i>E. hirae</i> . However, the test failed against bacteria based on <i>S. aureus</i> result, where a less than 5 log reduction was observed. Against <i>C. albicans</i> a reduction of ≥4 was observed together with ≤50 cfu in test fields 2 to 4, meeting the pass criterion of the standard.	<mark>2016e</mark>
Contec Meltblown Polypropylene 70/30 IPA PS- 850	Bacteria: Pseudomonas aeruginosa Staphylococcus aureus Enterococcus hirae Yeast: Candida albicans	The EN 16615 standard protocol was followed. Phase 2, Step 2 test. Concentrations tested: 100%, 0.1%. Interfering substance: 3.0g/l bovine albumin (dirty)	The log reduction was sufficient against P. aeruginosa and E. hirae. However, the test failed against bacteria based on S. aureus result, where a less than 5 log reduction was observed. Against <i>C. albicans</i> a reduction of ≥4 was observed together	<mark>2016f</mark>

		Contact time: 1 minute (bacteria), 3 minutes (yeast) Temperature: 20°C	with ≤50 cfu in test fields 2 to 4, meeting the pass criterion of the standard.	
70% IPA Wipes Solution	Yeast: Candida albicans	The EN 13697 standard protocol was followed. Phase 2, Step 2 test. Interfering substance: 0.3g/l bovine albumin (clean) Contact time: 1 minute Temperature: 20°C Concentrations tested: 100%, 50%, 0.1%	Against <i>C. albicans</i> a reduction of ≥3 was observed, meeting the pass criterion of the standard.	<mark>2016g</mark>
Alcohol Spray (FH55) 70% IPA	Yeast: Candida albicans	The EN 13697 standard protocol was followed. Phase 2, Step 2 test. Contact time: 15 minutes. Test temperature: 10°C Interfering substance: 0.3 % Bovine Serum Albumin (BSA) to represent dirty conditions and 0.03 % Bovine Serum Albumin (BSA) to represent clean conditions. Concentrations: 10 %, 50 % and 100 %.	Against <i>C. albicans</i> a reduction of ≥3 was observed both the 50 % and 100 % concentrations in dirty and clean conditions, meeting the pass criterion of the standard.	2017b
Contec Sterile 70% IPA	Bacteria: Pseudomonas aeruginosa Staphylococcus aureus Enterococcus hirae Escherichia coli Yeast: Candida albicans	The EN 13967 standard protocol was followed. Phase 2, Step 2 test. Interfering substance: 0.3g/l bovine albumin (BSA) to represent clean conditions. Contact time: 1 and 5 minutes to for bacteria; 3	Against all bacteria species tested a >4 log reduction was observed after both 1 and 5 minutes, meeting the pass criterion of the standard. Against <i>C. albicans</i> and <i>A. niger</i> a reduction of >3 was observed	<mark>2014a</mark>

		<mark>Fungi:</mark> Aspergillus niger	and 15 minutes for fungi and yeast. Temperature: 20°C Concentration: Ready to use (RTU).	after both 3 and 15 minutes, meeting the pass criterion of the standard.	
Contec (extrac wipe) 7 IPA	c IPA ct from 70%	Bacteria: Pseudomonas aeruginosa Staphylococcus aureus Enterococcus hirae Escherichia coli Yeast: Candida albicans Fungi: Aspergillus niger	The EN 13967 standard protocol was followed. Phase 2, Step 2 test. Interfering substance: 0.3g/l bovine albumin (BSA) to represent clean conditions. Contact time: 1 and 5 minutes to for bacteria; 3 and 15 minutes for fungi and yeast. Temperature: 20°C Concentration: Ready to use (RTU).	Against all bacteria species tested a >4 log reduction was observed after both 1 and 5 minutes, meeting the pass criterion of the standard. Against <i>C. albicans</i> and <i>A. niger</i> a reduction of >3 was observed after both 3 and 15 minutes, meeting the pass criterion of the standard.	<mark>2014b</mark>
70% IF Solutio		<mark>Yeast:</mark> Candida albicans	The EN 14562 (instrument disinfection in the medical area) standard protocol was followed. Phase 2, Step 2 test. Contact time: 1 minute Test temperature: 20°C Interfering substance: 0.3 g/l Bovine Serum Albumin (BSA) to represent clean conditions. Concentrations: 0.1 %, 50 % and 100 %.	For <i>C. albicans</i> a reduction greater than 4 log reduction (the pass criterion for the standard) was observed with both the 50 % and 100 % concentrations.	<mark>2016h</mark>
70% IF Wipes	PA	Viruses: Adenovirus 5 (ATCC VR-5/HeLa cells) Murine norovirus (s99/RAW 264.7 cells)	The EN 14476 (use in human medicine) standard protocol was followed. Phase 2, Step 1 test. Contact time: 1 minute	For Murine norovirus, the test product passed the test at 80 %, indicating that it possesses virucidal activity against this virus. For Adenovirus, the test product	<mark>2016</mark>

	Interfering substance: 0.3 g/l Bovine Serum	failed the test at 80 %, indicating that it does not possess sufficient virucidal activity against this virus.	
	Concentrations: 5.0 %, 50 % and 80 %.		

After the efficacy working group additional studies were provided to address issues raised. These are summarised below.

	Experimental data on the efficacy of the biocidal product against target organism(s)				
Function and field of use envisaged	Test substance	Test organism(s)	Test method/ Test system / concentrations applied / exposure time	Test results: effects	Reference
Spray Product for use against bacteria and yeast.	Contec Sterile IPA. (Spray product)	C. albicans	Laboratory Surface test – EN13697 Temperature: 18 – 25 °C Interfering substances: 0.3 g/L BSA (Clean conditions) Concentration tested: Ready to use product. Contact times: 1, 3 and 5 mins	The pass criteria for the test against yeast is ≥ 3 log reduction. The product demonstrated the following log reductions. 1 min: >4.85 3 mins: >4.90 5 mins: >4.49 The results therefore meet the pass criteria and support contact times of 1, 3 and 5 minutes against yeast. The controls were sufficient to validate	TRA-2018-326- 02_1 (to replace 2014a, 2014b 2016g)
Wipe product for use against bacteria and	Wipe solution on the standard wipe material	P. aeruginosa S. aureus E. hirae C. albicans	Laboratory Surface test (with mechanical action) – EN16615. Temperature: 20 °C	the test. The pass criteria for the test is ≥ 5 log reduction against bacteria and ≥ 4 log reduction against yeast.	J000973-1

yeast	(55% pulp, 45% PET Tork Premium Spezial Tucher)		Interfering substances: 0.3 g/L BSA (Clean conditions) Contact times: 1 min and 3 mins (yeast only)	The product demonstrated the following log reductions. 1 min: P. aeruginosa – 5 S. aureus – 5 E. hirae – > 5 C. albicans – 4 3 mins: C. albicans – 4 The results therefore meet the pass criteria and support contact times of 1 minute against bacteria yeast, as well as 3 minutes against yeast. The controls were sufficient to validate the test.	
	Prosat MBPP PS850 (100% polypropylene wipe)	S. aureus	Laboratory Surface test (with mechanical action) – EN16615. Temperature: 18-25 °C Interfering substances: 0.3 g/L BSA (Clean conditions) Contact times: 1 min for bacteria and 3 mins for yeast.	The pass criteria for the test is ≥ 5 log reduction against bacteria and ≥ 4 log reduction against yeast. The product demonstrated the following log reductions. 1 min: P. aeruginosa - 5.09 S. aureus - >5.32 E. hirae - >5.03 3 mins: C. albicans - >4.06 The results therefore meet the pass	TRA-2018-332- 01 (To replace 2016d)

Prosat Sigma PSC-20001 (55% cellulose/45% polyester wipe)	P. aeruginosa S. aureus E. hirae C. albicans	Laboratory Surface test (with mechanical action) – EN16615. Temperature: 18-25 °C Interfering substances: 0.3 g/L BSA (Clean conditions) Contact times: 1 min for bacteria and 3 mins for yeast.	criteria and support contact times of 1 minute against bacteria yeast, as well as 3 minutes against yeast. The controls were sufficient to validate the test. The pass criteria for the test is ≥ 5 log reduction against bacteria and ≥ 4 log reduction against yeast. The product demonstrated the following log reductions. 1 min: P. aeruginosa - 5.09 S. aureus - >5.32 E. hirae - >5.03 3 mins: C. albicans - >4.06 The results therefore meet the pass criteria and support contact times of 1 minute against bacteria yeast, as well as 3 minutes against yeast.	TRA-2018-333- 02 (to replace 2016a and 2016)
			The controls were sufficient to validate the test.	
Prosat Polynit Heatseal PSWE0001 (100 % knitted	P. aeruginosa S. aureus E. hirae C. albicans	Laboratory Surface test (with mechanical action) – EN16615. Temperature: 18-25 °C	The pass criteria for the test is ≥ 5 log reduction against bacteria and ≥ 4 log reduction against yeast. The product demonstrated the following	TRA-2018-334- 02 (to replace 2016 b

Interfering substances: 0.3 g/L BSA	log reductions.	and f)
(Clean conditions)	1 min:	
	P. aeruginosa – 5.09	
Contact times: 1 min for bacteria	S. aureus - >5.32	
and 3 mins for yeast.	E. hirae - >5.03	
	3 mins:	
	C. albicans - >4.06	
	·	
	·	
	, , , ,	
	3 minutes against yeast.	
	The centrals were sufficient to validate	
	the test.	
	(Clean conditions) Contact times: 1 min for bacteria	(Clean conditions) 1 min: P. aeruginosa – 5.09 S. aureus – >5.32 E. hirae – >5.03 3 mins:

Conclusion on the efficacy of the product

The Contec IPA Product Family claims the following:

- 'Suitable for use on clean hard surfaces'
- Specifically claimed are:
 - 1 minute contact time for bacteria (EN 13697).
 - 3 minute contact time for fungi (EN 13697).

The products of the family are proposed for use on hard surfaces in industrial and institutional areas.

Data for instrument disinfection, virus control and mycobacteria control have been submitted; however, the product family makes no claims for use as instrument disinfectants or as surface disinfectants in the medical area. Furthermore, no label claims have been made against viruses or mycobacteria (although both are included in the SPC). Nevertheless, the UK has evaluated all of these data and summarised the results in each section.

All of the products tested in the studies contain the same formulation and all of the products in the family contain the same formulation. Therefore, the products tested in the studies are acceptable in support of Contec IPA Product Family.

Efficacy against bacteria

For claims of bactericidal activity from disinfection in the food and feed area, it is a requirement to test the products in accordance with EN 1276 (phase 2, step 1) and EN 13697 (phase 2, step 2). For wipe products, testing with mechanical action according EN 16615 (phase 2, step 2), is typically required.

For bactericidal claims of instrument disinfection in the medical area, tests according to EN 13727 (phase 2, step 1) and EN 14561 (phase 2, step 2) are typically required. For bactericidal claims of surface disinfection in the medical area, tests following EN 13727 (phase 2, step 1) are required along with a test to EN 13697 (phase 2, step 2) for no mechanical action and a test to EN 16615 (phase 2, step 2) for mechanical action.

Against bacteria the product family has passed EN 1276 (2014a) under dirty conditions at a one minute contact time against the standard test organisms for use as a general disinfectant. In this test the liquid from extracted from a wipe product was used. The products also passed EN 1276 (2017a) when tested at a cold temperature (10°C) with a contact time of 1 minute in dirty conditions.

Efficacy has also been demonstrated in four separate tests following the EN 13697 standard. Two of these tests used dirty conditions and the standard test organisms; one of which used a 30 second contact time (2013a) and the other 1 minute (2016). The other two tests (2014a & 2014b) used clean conditions and a 1 minute contact time against the standard test organisms. One of these studies tested a cold temperature (4°C). The log reduction required to pass these tests was achieved in all cases.

The EN 1276 studies together with the EN 13697 studies sufficiently demonstrate that the product family is acceptable for use as a general surface disinfectant against bacteria in a general use area.

For use in the medical area studies according to EN 13727 and EN 14561 were submitted. Specifically for the wipe product numerous studies conducted to EN 16615 were submitted.

The study conducted to EN 13727 (2014b) demonstrates that the product family is significantly efficacious against the standard test bacteria when applied for one minute in dirty conditions.

The two studies conducted to EN 14561 both used a 30 second contact time and dirty conditions; but they differ in the bacteria used. 2013b included the standard test organisms for surface disinfection claims in the medical area whereas Watson, D.C; 2013c tested against methicillin resistant *S. aureus* and vancomycin resistant *E. faecium*. Both tests conducted according to EN 14561 demonstrate acceptable efficacy against the respective test organisms included.

Two separate studies were conducted according to the EN 16615 standard and demonstrated acceptable efficacy of the wipe products against *S. aureus* (the only tested organism) using clean conditions and a 1 minute contact time (2016a & 2016b). The five remaining EN 16615 studies all tested the efficacy of wipe products against *P. aeruginosa*, *S. aureus* and *E. hirae*. Of these five studies, two failed to demonstrate acceptable efficacy against *S. aureus* according to the standard under dirty conditions with a contact time of 1 minute (2016e & 2016f). The other three studies all support the product family against bacteria according to EN 16615. Two of these studies used a 5 minute contact time and clean conditions (2015a & 2015b) and 1 test (2016d) passed the requirements of EN 16615 under dirty conditions with a 1 minute contact time.

For use with mechanical action according to EN 16615, efficacy has been sufficiently demonstrated in clean conditions with a contact time of 1 minute. The products only claim to disinfect 'clean hard surfaces' so this is acceptable and efficacy in dirty conditions do not need to be considered. The UK CA considers that in areas of heavy soiling, the surface should be cleaned (until it is visibly clean) prior to disinfection. This could be achieved for example by using 1 wipe to clean the surface/equipment and another to disinfect it.

Efficacy against mycobacteria

At present there are no standard tests to demonstrate mycobactericidal activity in the food and feed area specified in the guidance. However, given the similarity of the conditions, the UK CA considers that the tests specified for PT 2 can also be used for PT 4. For claims of mycobactericidal activity instrument disinfection in the medical area EN 14348 (phase 2, step 1) and EN 14563 (phase 2, step 2) are typically required. For surface disinfection with or without mechanical action, EN 14348 is typically required.

A test according to EN 14348 (2014) which uses a 1 minute contact time under dirty conditions was submitted. Both *M. avium* and *M. terrae* show reductions that meet the test validation requirement.

A test according to EN 14563 (2017) which uses a 1 minute contact time under clean conditions was submitted. Both *M. avium* and *M. terrae* show reductions that meet the test validation requirement.

Overall, the UK CA considers that efficacy (with and without mechanical action) against mycobacteria has been sufficiently demonstrated in clean conditions with a contact time of 1 minute. This use is considered to be supported in both general use areas and in medical areas, including instrument disinfection.

Efficacy against fungi/yeasts

For claims of fungicidal/yeasticidal activity for disinfectants in the food and feed area, it is typically a requirement to test the products in accordance with EN 1650 (phase 2, step 1) and EN 13697 (phase 2, step 2) when mechanical action is not required. To assess mechanical action, a test to EN 16615 (phase 2, step 2) should normally be conducted. However, as no mechanical action test is available for fungi, testing can be done with liquid extracted from the wipes in an EN 13697 test.

For claims of fungicidal/yeasticidal activity for an instrument disinfectant in the medical area, it is typically a requirement to test the products in accordance with EN 13624 (phase 2, step 1) and EN 14562 (phase 2, step 2) when mechanical action is not required.

For claims of fungicidal/yeasticidal activity for a surface disinfectant in the medical area, it is typically a requirement to test the products in accordance with EN 13624 (phase 2, step 1), along with the same phase 2, step 2 testing required for the food and feed area.

In the first EN 1650 study (2014c), the products demonstrated sufficient efficacy against C. albicans according to the pass criterion, with a contact time of 1 minute in dirty conditions. However, the products did not demonstrate sufficient efficacy against A. niger to meet the pass criterion of the standard. In the second EN 1650 study (2017c) only C. albicans was tested. The products demonstrated sufficient efficacy according to the pass criterion, with a contact time of 15 minutes in dirty conditions and at a cold temperature (10°C). 2016c investigated the yeasticidal activity of the products in the medical area, according to EN 13624. Clean conditions were tested. This test demonstrated the efficacy of the products against C. albicans according to the pass criterion for the relevant standard protocol. The contact time used in the tests was 1 minute. 2006 investigated the yeasticidal and fungicidal activity of the products in the medical area, according to EN 13624. Clean conditions were tested. The products demonstrated sufficient efficacy against C. albicans according to the pass criterion, with a contact time of 1 minute. However, the products did not demonstrate sufficient efficacy

In one EN 13697 study (2016g), the products demonstrated sufficient efficacy against *C. albicans* according to the pass criterion, with a contact time of 1 minute in clean conditions.

against A. niger to meet the pass criterion of the standard.

In a second EN 13697 study 2017b), the products demonstrated sufficient efficacy against *C. albicans* according to the pass criterion, with a contact time of 15 minutes in clean conditions and at a cold temperature (10°C).

In the two other EN 13697 studies (2014a & 2014b), the products demonstrated sufficient efficacy against both *C. albicans* and *A. niger* according to the

pass criterion, with a contact time of 3 minutes in clean conditions.

2016 investigated the yeasticidal activity of the products in the medical area, according to EN 14562. Clean conditions were tested. This test demonstrated the efficacy of the products against *C. albicans* according to the pass criterion for the relevant standard protocol. The contact time used in the tests was 1 minute.

The yeasticidal activity of the product family was investigated in five studies conducted according to EN 16615 (2015a, 2015b, 2016d, 2016e & 2016f). Dirty conditions were used in 2 of the tests and clean conditions were used in the other three tests. In all of the studies, the products demonstrated sufficient efficacy against *C. albicans* according to the pass criterion, with a contact time of 3 to 5 minutes in dirty conditions was sufficient).

Overall, the UK CA considers that efficacy (with and without mechanical action) against yeast have been sufficiently demonstrated in clean conditions with contact time of 1 minute.

The use against yeast is supported in general use areas and in medical areas, including instrument disinfection. The use against fungi is supported in general use areas and on surfaces in the medical area; however, it is not supported for instrument disinfection, as efficacy against fungi has not been demonstrated in EN 13624 and EN 14562.

During the initial evaluation, the UK CA had concerns that a 3 minute contact time might not be appropriate for a wipe product.

The UK CA initiated an e-consultation with other MS on 4th January 2018 in order to discuss what contact times might be appropriate for wipe disinfectant products. Although there was no clear conclusion drawn in this e-consultation, it did appear that MS generally considered that contact times longer than 1 minute might be acceptable, provided a reasoned case is provided to justify why the times are appropriate for the formulation and use instructions.

The UK CA made the applicant aware that their claims against fungi are not considered to be supported and asked for them to provide a reasoned case to justify how a contact time of 3 minutes is appropriate for a wipe product. The applicant did not submit a reasoned case and chose to withdraw support for fungicidal claims. Therefore, the UK CA concludes that fungi claims are not currently supported.

If the applicant wishes to make fungicidal claims in the future, this may be possible if an acceptable reasoned case is provided to justify this use.

Efficacy against viruses

No claims have been made against viruses, nevertheless the UK CA has summarised the results of the submitted studies.

For claims of virucidal activity from a surface or instrument disinfectant in the medical area, it is typically a requirement to demonstrate \geq 4.0 log reduction of *Poliovirus*, *Adenovirus* and *Murine norovirus* in accordance with EN 14476 (phase 2, step 1).

In support of the efficacy against viruses one EN 14476 study (2016b) was submitted. The viruses tested in this study were *Adenovirus* and *Murine norovirus*. The

contact time used in the test was 1 minute and clean conditions were tested. According to the results of this study, the 70 % v/v propan-2-ol wipes possessed sufficient virucidal activity against *Murine norovirus*. However, the products did not possess sufficient virucidal activity against *Adenovirus* and failed the test for that virus.

Only efficacy against *Murine norovirus* has been demonstrated according to the pass criterion for the relevant standard protocol. Therefore, the UK considers that virucidal efficacy has not sufficiently been demonstrated in order to support a claim.

Decision

The UK CA concludes that the submitted data are sufficient to support all of the products in the Contec IPA Product Family.

All areas of use claimed for the product family are supported by the data provided with the exception of fungicidal activity with instrument disinfection.

Sufficient data were provided to demonstrate that the products are efficacious against bacteria, mycobacteria, and yeast in clean conditions with a contact time of 3 minutes. Efficacy against viruses is not supported.

New data submitted after the efficacy working group

After the efficacy working group, additional studies were provided to address the issues raised. These are summarised in the table above and are discussed below.

According to the conclusions of the Efficacy Working Group, December 2018, the open points related to two key issues:

- A. The efficacy of the spray product against yeast was proven only for the longer contact time of 15 minutes but not for the shorter contact times requested by the applicant of 1-3 minutes.
- B. Due to issues with the old EN 16615 studies, the efficacy of the wipe products was not demonstrated against bacteria and yeast. The standard wipe (from EN16615) or the exact wipes used for the product should be tested.

Five new studies were provided to address these issues.

- A. A new phase 2 step 2 study (TRA-2018-326-02_1) using the spray product against *C. albicans* was provided. This was conducted according to EN 13697. This new study demonstrates a sufficient reduction against yeast after contact times of 1, 3 and 5 minutes. The UK CA therefore considers that, alongside the existing acceptable 2,1 tests, sufficient evidence has now been provided to address this issue and the spray product can be authorised against yeast with a minimum contact time of 1 minute.
- B. A new EN16615 study (J000973-1) has been provided using the standard wipe 55% pulp, 45% PET Tork Premium Spezial Tucher. This study demonstrated sufficient log reductions against *P. aeruginosa, S. aureus* and *E. hirae* with a contact time of 1 minute and against *C. albicans* with a contact time of 3 minutes. These data, therefore, support the efficacy of the wipe products against bacteria and yeasts with contact times of 1 and 3 minutes respectively.

Additionally, as this wipe is the standard wipe described in EN 16615 as the

appropriate carrier to test where various different types of wipe may be used with the same product formulation, the UK CA considers that this study can also be used in support of the other wipe products as well. This seemed to be the view shared by the efficacy working group. We note that the applicant also provided a justification document to explain why they consider some types of wipe to be worst case. However, as the standard wipe has now been successfully tested, this justification may no longer be necessary.

In addition to this, the applicant has also provided new studies using other wipe materials to support the use of the product on other types of wipe. The studies TRA-2018-332-02, TRA-2018-333-02 and TRA-2018-334-02 tested 100% polypropylene wipes, 55% cellulose/45% polyester wipes, and 100 % knitted polyester respectively. In all of these studies the data demonstrate reductions sufficient to pass the test against bacteria with a 1 minute contact time and against yeast with a 3 minute contact time. Therefore, these data further support the efficacy of the wipe products against bacteria and yeasts with contact times of 1 and 3 minutes respectively.

Consequently, the UK CA considers that the new data provided addresses the concerns raised at the efficacy working group and supports the authorisation of all of the wipe products.

2.2.5.6 Occurrence of resistance and resistance management

The applicant has provided the following statement on the occurrence of resistance and resistance management:

'The Active Substance Doc IIA indicates the following:

Due to the unspecific mode of action of alcohols, i.e., denaturation and coagulation of proteins, cell lysis and disruption of the cellular metabolism, resistance against alcohols is not expected. In none of the presented data resistance has been reported. Also, according to (2004) no acquired resistance to alcohols has been reported. (1999) present an extensive overview over intrinsic and acquired resistance of antiseptics and disinfectants, but they did not find any evidence for acquired resistance of Propan-2-ol for bacteria, fungi or viruses.

Management

According to the Assessment Report of the active substance, Propan-2-ol, no known resistance has been reported against the target species.'

[UK CA note: this refers to the Doc IIA submitted by ASD Consortium in support of the alternative source of propan-2-ol]

The UK CA accepts that there is no significant risk of the development of resistance for this active substance and products, however, if the applicant becomes aware of any reports of resistance to the active substance propan-2-ol and/or the products these should be reported to appropriate bodies (such as the efficacy working group and/or concerned member states) so that it can be determined if further action is required.

2.2.5.7 Known limitations

There are no known limitations, but the products only claim to disinfect 'clean hard surfaces'. Therefore, the use in dirty conditions is not relevant.

2.2.5.8 Evaluation of the label claims

The UK CA considers that the following requested label claims are supported at room temperature:

- 'Suitable for use on clean hard surfaces'
- Specifically claimed are:
 - 1 minute contact time for bacteria (EN 13697).

Additionally the following usage instructions should be included on the label:

- Clean carefully the surfaces before application
- Apply the the wipe uniformly on the surface to be treated in sufficient quantity (One wipe per $1 1.5 \text{ m}^2$ of surface) so that the surface remains wet during the contact time.
- Wipes
 - "Pre-cleaning may be required prior to disinfection if surface is visually soiled"
 "Ensure the surface is uniformly covered with the product, <u>allow to act for a contact time of 1 minute for bactericidal and mycobactericidal action and 3 minutes for yeasticidal action"</u>
- Spray
 - "Ensure the surface is uniformly covered with the product, <u>allow to act for a contact</u> time of 1 minute"

Maximum of 2 wipes per m2

2.2.5.9 Relevant information if the product is intended to be authorised for use with other biocidal product(s)

The products of the family are not intended for use with other products.

2.2.6 Risk assessment for human health

2.2.6.1 Assessment of effects on Human Health

Skin corrosion and irritation

Conclusion used in	Conclusion used in Risk Assessment – Skin corrosion and irritation		
Value/conclusion	Contec IPA Product Family is not classified for skin corrosion or irritation.		
Justification for the value/conclusion	None of the components of the product family are classified for skin corrosion or irritation according to Regulation (EC) No 1272/2008. The product family does not therefore meet the criteria for classification for skin corrosion or irritation according to Regulation (EC) No 1272/2008.		
Classification of the product family according to CLP	Contec IPA Product Family is not classified for skin corrosion or irritation.		

Eye irritation

Conclusion used in	Risk Assessment - Eye irritation
Value/conclusion	Contec IPA Product Family is classified for eye irritation cat. 2 (H319).
Justification for the value/conclusion	Propan-2-ol is classified as an eye irritant (Cat. 2) and is present at a maximum of 70 % in the product family. No other components in the product family are classified as eye irritants. According to Regulation (EC) No 1272/2008, a mixture which contains a total of 10 % or more of a substance or substances classified in Category 2 for eye irritation shall be classified in Category 2 for eye irritation. Therefore, the product family meets the criteria to be classified for eye irritation category 2 (H319).
Classification of the product family according to CLP	Contec IPA Product Family is classified for eye irritation cat. 2 - H319: Causes serious eye irritation.

Respiratory tract irritation

Conclusion used in	Conclusion used in the Risk Assessment – Respiratory tract irritation		
Value/conclusion	Contec IPA Product Family is not classified for respiratory tract irritation.		
Justification for the conclusion	None of the components of the product family are classified for respiratory tract irritation according to Regulation (EC) No 1272/2008. The product family does not therefore meet the criteria for classification for respiratory tract irritation according to Regulation (EC) No 1272/2008.		
Classification of the product family according to CLP	Contec IPA Product Family is not classified for respiratory tract irritation.		

Skin sensitization

Conclusion used in Risk Assessment – Skin sensitisation		
Value/conclusion	Contec IPA Product Family is not classified for skin sensitisation.	
Justification for the value/conclusion	None of the components of the product family are classified for skin sensitisation according to Regulation (EC) No 1272/2008. The product family does not therefore meet the criteria for classification for skin sensitisation according to Regulation (EC) No 1272/2008.	
Classification of the product family according to CLP	Contec IPA Product Family is not classified for skin sensitisation.	

Respiratory sensitization (ADS)

Conclusion used in	Conclusion used in Risk Assessment – Respiratory sensitisation		
Value/conclusion	Contec IPA Product Family is not classified for respiratory sensitisation.		
Justification for the value/conclusion	None of the components of the product family are classified for respiratory sensitisation according to Regulation (EC) No 1272/2008. The product family does not therefore meet the criteria for classification for respiratory sensitisation according to Regulation (EC) No 1272/2008.		
Classification of the product family according to CLP	Contec IPA Product Family is not classified for respiratory sensitisation.		

Acute toxicity

Acute toxicity by oral route

Conclusion used in t	Conclusion used in the Risk Assessment – Acute oral toxicity		
Value/conclusion	Contec IPA Product Family is not classified for acute oral toxicity.		
Justification for the value/conclusion	None of the components of the product family are classified for acute oral toxicity according to Regulation (EC) No 1272/2008. The product family does not therefore meet the criteria for classification for acute oral toxicity according to Regulation (EC) No 1272/2008.		
Classification of the product family according to CLP	Contec IPA Product Family is not classified for acute oral toxicity.		

Acute toxicity by inhalation

Conclusion used in the Risk Assessment – Acute inhalation toxicity			
Value/conclusion	Contec IPA Product Family is not classified for acute inhalation toxicity.		
Justification for the value/conclusion	None of the components of the product family are classified for acute inhalation toxicity according to Regulation (EC) No 1272/2008. The product family does not therefore meet the criteria for classification for acute inhalation toxicity according to Regulation (EC) No 1272/2008.		
Classification of the product family according to CLP	Contec IPA Product Family is not classified for acute inhalation toxicity.		

Acute toxicity by dermal route

Conclusion used in the Risk Assessment – Acute dermal toxicity			
Value/conclusion	Contec IPA Product Family is not classified for acute dermal toxicity.		
Justification for the value/conclusion	None of the components of the product family are classified for acute dermal toxicity according to Regulation (EC) No 1272/2008. The product family does not therefore meet the criteria for classification for acute dermal toxicity according to Regulation (EC) No 1272/2008.		
Classification of the product family according to CLP	Contec IPA Product Family is not classified for acute dermal toxicity.		

Information on dermal absorption

Value(s) used in the Risk Assessment – Dermal absorption		
Substance	Propan-2-ol at 70 % (w/w)	
Value(s)	0.85 mg/cm ² /h (transdermal flux rate)	
Justification for the selected value(s)	Dermal absorption is read-across to the value presented in the agreed List of End Points (LoEP) for propan-2-ol which is based on <i>in vivo</i> data for male and female rats. The tested formulation was a dummy product, a model formulation consisting of the active substance and water; this is sufficiently similar to the product family to enable read-across of these data.	

Assessment of endocrine disruption (ED) properties of active substances and co-formulants in biocidal products

According to the CAR for propan-2-ol, there is no indication for endocrine disrupting properties of the active substance. Additionally, there is no indication for endocrine disrupting properties of the the co-formulants of the biocidal product.

In summary, there is no indication for endocrine disrupting properties of the biocidal product.

Available toxicological data relating to non active substance(s) (i.e. substance(s) of concern)

Contec IPA Product Family does not contain any substances of concern.

Available toxicological data relating to a mixture

None of the co-formulants of Contec IPA Product Family are mixtures.

2.2.6.2 Exposure assessment

Contec IPA Product Family contains three types of ready-to-use (RTU) disinfectant products: ready to use trigger spray liquid, liquid refill and impregnated wipe, containing 70% v/v (equivalent to 62.9% w/w) propan-2-ol as the active substance.

The products are for use by professionals for hard surface disinfection in industrial/manufacturing settings under two product types:

PT 2 – Cleanrooms for pharmaceutical, biopharmaceutical, medical device and diagnostic product manufacturing facilities and critical life science applications

PT 4 - Industrial food and feed preparation areas (excluding mopping of walls, floors and ceilings).

An overview of the products in the family is provided below.

Intended uses of the product family				
SPC	Product category	Use	Application rate	
	Ready-to-use liquid	PT2: Professional hard surface disinfectant for use in cleanrooms		
Meta SPC 1	trigger spray and liquid refill	PT4: Professional hard surface disinfectant for use in the preparation of food in industrial settings	50 ml/m²	
Meta	Ready-to-use	PT2: Professional hard surface disinfectant for use in cleanrooms	5-38 ml per wipe depending on specific product. (2 wipes/m²)	
SPC 2	impregnated disinfectant wipe	PT4: Professional hard surface disinfectant for use in the preparation of food in industrial settings	5-26 ml per wipe depending on specific product (2 wipes/m²)	

The RTU disinfectant liquid is supplied in 5 L containers for use to refill trigger sprayers and the RTU disinfectant trigger sprayer is supplied in 0.5 and 1 L trigger spray bottles.

The RTU impregnated disinfectant wipe product is supplied in a packet with varying numbers of wipes per pouch depending on the specific product. The in-use product density of 0.785 g/ml has been used in the exposure calculations.

It is assumed that trigger sprayers and impregnated wipes are intended for use to disinfection small surfaces.

Identification of main paths of human exposure towards active substance(s) and substances of concern from its use in biocidal product

Summary table: relevant paths of human exposure							
Primary (direct) exposure		Secondary (indirect) exposure					
Exposure path	Industrial use	Professional use	General public use	Industrial use	Professional use	Bystanders	Via food
Inhalation	N/A	Yes	N/A	N/A	Yes	N/A	N/A
Dermal*	N/A	Yes	N/A	N/A	Yes	N/A	N/A
Oral	N/A	No	N/A	N/A	No	N/A	N/A

^{*}According to cleanroom standards and protocols², cleanroom clothing must be worn to eliminate dispersion of contamination from skin and non-cleanroom clothing e.g. coveralls, gloves and face masks. Whilst cleanroom clothing covers any exposed skin and is worn over normal clothing, these may not protect workers from chemicals. Level of dermal exposure has been predicted but is recognised that the critical route of exposure is via inhalation given the high volatility of propan-2-ol.

List of scenarios for PT2

Critically for this risk assessment, the amount of product and frequency of use at any one time or during the day is undefined. Use instructions include 'As required and according to use instructions' and 'Use frequency could be several hours per day'. Although small area use/routine disinfection of 0.5 m² within laboratory environments is defined in the Assessment Report, there is no consideration of cleanroom environments or professional cleanroom users. As a result the UK CA has drawn values and approaches from various guidance (e.g. HEEG opinions, HEAdHoc recommendations, RIVM report 320104003/2006 Cleaning Products Fact Sheet) to construct a representative risk assessment.

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² Cleanrooms and associated controlled environments —BS EN ISO 14644-5

	Summary table: scenarios			
Scenario number	Scenario	Primary or secondary exposure Description of scenario	Exposed group	
1.	Pouring of RTU liquid to refill trigger sprayer	Primary exposure: professional user pouring RTU liquid from a 5 L container for refilling trigger sprayer	Professional	
2.	Routine disinfection of small surfaces in manufacturing/industrial settings e.g. cleanroom environments	routine disinfection of small surfaces e.g. equipment and work stations as part of	Professional	
3.	Inhalation of volatilised residues in industrial/manufacturing settings	, , ,		

Industrial exposure

Products of the Contec IPA Product Family are intended for use by professional users only.

Professional exposure

<u>Scenario 1: Primary exposure: professional user pouring RTU liquid from a 5 L container for refilling trigger sprayer</u>

According to the HEEG Opinion 1 (2008)³, a recommended alternative approach for repeated loading for small quantities is TNsG Mixing and loading model 4. It is noted that re-filling/pouring of the product will be covered within this mixing and loading step and does not need to be assessed separately. This is because the model covers all relevant mixing and loading tasks performed by a worker on an 8-hour working day.

Considering the application rate for the RTU liquid trigger sprayer is 50 ml/m^2 , a 5 L container contains sufficient product to treat an area of 100 m^2 . It is therefore not envisaged that a professional user would achieve a work rate of $>100 \text{ m}^2$ per day through a trigger sprayer. Moreover it is unlikely that 0.5 and 1 L trigger spray bottles will be used up by a professional user in one day as a trigger sprayer is unlikely to deliver large amounts of product per use/actuation e.g. 1 ml/actuation, thus the prolonged use of the trigger sprayer would lead to hand fatigue. As such, for the worst-case ten pouring/loading operations per day is assume for a professional user (5 L container would refill $10 \times 0.5 \text{ L}$ trigger spray bottles).

Inhalation exposure has been calculated using ConsExpo Web and the default parameters in the ConsExpo Cleaning Products Fact Sheet⁴ (p.58) for the exposure during mixing and loading of a liquid cleaner.

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³ HEEG Opinion on the use of available data and models for the assessment of the exposure of operators during the loading of products into vessels or systems in industrial scale

⁴ RIVM report 320104003/2006 Cleaning Products Fact Sheet

Description of Scenario 1

A professional user pouring RTU liquid disinfectants from a 5 L container into receiving vessel e.g. trigger sprayer bottle. TNsG Mixing and Loading Model 4 have been used to predict dermal exposure and ConsExpo exposure to vapour (evaporation from constant surface) has been used to predict inhalation exposure.

	Parameters	Value
	Adult body weight	60 kg
	Concentration of active in RTU spray	62.9% w/w
Dormal	Density of the product	0.786 g/ml (applicant information)
Dermal exposure	Contamination indicative value from a 5 L container	0.2 ml product/operation
	Dermal flux rate	0.85 mg/cm ² /h
	Number of operations	10 per day
	Exposure duration ⁴	0.75 min
	Product amount ¹	1965 g
	Room volume (user breathing zone) ⁴	1 m ³
Inhalation	Ventilation rate (default for unspecified room)	0.5/hr
	Release area ⁴ (circular opening of 5 cm diameter for a 5 L container)	20 cm ²
exposure	Application duration ⁴	0.25 min
	Temperature	20 °C
	Mass transfer rate	Thibodeaux model
	Molecular weight matrix ²	18 g/mol
	Vapour pressure (EU-agreed value (LoEP January 2015))	5780 Pa at 25°C

¹The product amount is half the amount of the bottle content 5 L (ConsExpo Product Fact Sheet)

Calculations for Scenario 1

Dermal exposure

The indicative dermal contamination value from TNsG Mixing and Loading Model 4 is 0.2 ml/operation for unspecified design (worst-case consideration). This would result in a total amount of 1.006 g propan-2-ol. Assuming that ten mixing/loading operation is performed in a day and the surface area of both hands is available for dermal exposure (820 cm²), following the approach in the propan-2-ol Assessment Report, the time of evaporation is calculated as follows:

 $T = mRTK/M\beta$ pA

²The molecular weight of the matrix is based on the formulation. Please see the confidential Annex Section 3.6.1 for details of the full formulation.

t = time [s]

m = mass of propan-2-ol on surface [mg] (1006 mg)

 $R = \text{gas constant [J K-1 mol-1] (8.314 J K}^{-1} \text{ mol}^{-1})$

 $T = \frac{\text{skin}}{\text{surface temperature [K] (303.15 K)}}$

K = conversion factor (36000)

 $M = \text{molar mass [g mol-1] (60.1 g mol^{-1})}$

 β = mass transfer coefficient [m h-1], for calculation see TGD (8.7 m h-1)

p = vapour pressure of the pure substance [Pa] (7600 Pa (30°C))

A = surface area [cm²] (820 cm²)

According to this equation the evaporation time is 28 secs. Therefore for a worst-case scenario, it is assumed that the total amount evaporates at once after this time interval. The absorption/dermal flux rate for propan-2-ol in a 70% w/w aqueous dilution is 0.85 mg/cm²/h (EU-agreed value (LoEP January 2015)). During a time interval of 28 secs on the skin surface of both hands 820 cm², this result in a total absorbed amount of 5.4 mg per day. This is equivalent to 0.09 mg/kg bw/d without PPE.

It should be noted that although the amount of product used per day varies depending on the situation of use, the predicted dermal exposure is considerably low such that an unrealistic number of mixing/loading operations would have to occur in order for a professional user to exceed the AEL.

Summary table: estimated exposure from professional uses			
Tier/PPE	Estimated inhalation uptake (mg/kg bw/d)	Estimated dermal uptake (mg/kg bw/d)	Estimated total uptake (mg/kg bw/d)
1 (no PPE)	0.0018	0.09	0.092
Please see Annex 3.2.1 for detailed calculations			

<u>Scenario 2: Primary exposure: a technician performs routine disinfection of small surfaces e.g. equipment and work stations as part of their working procedure in manufacturing/industrial cleanroom environments</u>

The application rate to surfaces is 50 ml/m^2 via the trigger spray and 2 wipes/m^2 for wet wipes containing 5-38 ml product per wipe. The worst-case scenario is therefore represented by the use of the products through wet wipes at an application rate of 76 ml/m^2 ($38 \text{ ml} \times 2 \text{ wipes}$). Thus the following risk assessment considers a professional user disinfecting surfaces using impregnated wet wipes and exposure from the use of trigger sprayer is within the risk envelope.

According to cleanroom protocols, professional workers working in cleanrooms are expected to disinfect personal work surfaces and equipment before and/or at the end of each work shift as good practice. Since disinfecting these surfaces is performed as part of their work day which is in addition to the performing assigned tasks in the cleanroom, disinfection will take place on a routine basis. In considering this pattern of use, the UK CA considers the scenario presented in the PT 2 Assessment Report for disinfection of small surfaces is representative for the risk assessment, where:

Work rate = Area to be disinfected per event $(0.5 \text{ m}^2 \text{ using } 1 \text{ wipe})$

Exposure duration = 45 mins per event

Application duration = 1 min per event

Frequency = 10 events per day (5 m² using 10 wipes)

Ventilation in cleanrooms varies based on cleanroom standards where lower ventilation rates are observed in higher classed cleanrooms with higher levels of particulates. A tier 1 worst-case assumption of 8/h ventilation in line with the ventilation rate for a laboratory in the propan-2-ol PT2 Assessment Report has been used. It should be recognised that a ventilation rate of 8/h is highly conservative for cleanrooms; according to the guidelines produced by the FDA⁵, minimum air exchange rate for IOS 8 cleanrooms is 20 per hour. The applicant has proposed a representative room volume of 55 m³ for a cleanroom from personal communication with Ctgb.

Description of Scenario 2

A professional worker in a cleanroom disinfects small surfaces e.g. equipment and work stations throughout the working day. Based on the approach outlined in the propan-2-ol PT 2 Assessment Report, it is assumed that a professional worker disinfects a work bench of $0.5 \, \text{m}^2$ every $45 \, \text{minutes}$.

Due to the high vapour pressure of propan-2-ol, the evaporation and airborne phase of the active from the surface is the critical route of exposure. In accordance with ConsExpo cleaning factsheet, inhalation exposure has been calculated using ConsExpo Web exposure to vapour: evaporation model (increasing area) for cleaning with wet tissue (surrogate for wet wipes).

	Parameters	Value	
	Adult body weight	60 kg	
	Concentration of active	62.9% w/w	
	Product density	0.786 g/ml (applicant information)	
	Inhalation rate	1.25 m ³ /hr	
	Molecular weight	60 g/mol	
	Vapour pressure (EU-agreed value (LoEP January 2015))	5780 Pa at 25°C	
Inhalation	Amount of product used to treat 0.5 m ²	30 g (1 wipe x 38 ml)	
exposure	Exposure duration (Assessment Report)	45 mins	
	Room volume	55 m³ (applicant information)	
	Ventilation rate	8/hr	
	Application duration (Assessment Report)	1 min	
	Release area (Assessment Report)	0.5 m ²	
	Molecular weight matrix ¹	18 g/mol	
	Mass transfer rate	Thibodeaux method	

⁵ Guidance for Industry: Sterile drug products produced by aseptic processing – Current Good Manufacturing Practice, 2004

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¹The molecular weight of the matrix is based on the formulation. Please see the confidential Annex Section 3.6.1 for details of the full formulation.

Calculations for Scenario 2

Tier 1 assessment

Dermal exposure

ConsExpo Cleaning Product Factsheet⁴ (p.64) estimated that during wiping with wet tissues, 1.4% of the total liquid of the wet tissue remains on the surface of the inner hand. Therefore the amount of product available for dermal exposure when using one wet wipe is 0.418 g (38 ml/wipe x 0.786 g/ml x 1.4%) or 268 mg propan-2-ol. Following the assumption in the propan-2-ol PT 2 Assessment Report and ECHA Methodology for disinfection by single use wiping tissue (p. 106), the time of evaporation is calculated according to TGD on risk assessment, App. I, App. IF:

$T = mRTK/M\beta$ pA

```
t = time [s]

m = mass of propan-2-ol on surface [mg] (268 mg)

R = gas constant [J K-1 mol-1] (8.314 J K<sup>-1</sup> mol<sup>-1</sup>)

T = skin/surface temperature [K] (303.15 K)

K = conversion factor (36000)

M = molar mass [g mol-1] (60.1 g mol<sup>-1</sup>)

R = mass transfer coefficient [m h-1], for calculation see TGD (8.7 m h-1)
```

p = vapour pressure of the pure substance [Pa] (7600 Pa (30°C))

According to this equation the evaporation time is 30 seconds. Therefore for a worst case scenario it is assumed that the total amount evaporates at once after this time interval. The absorption/dermal flux rate for propan-2-ol in a 70% w/w aqueous dilution is $0.85 \, \text{mg/cm}^2/\text{h}$ (EU-agreed value (LoEP January 2015)). During the time interval of 30 seconds on the skin surface of palm of one hand 205 cm², this result in a total amount absorbed per day is $1.45 \, \text{mg}$.

Therefore the total amount absorbed from the use of 10 wet wipes to disinfect 5 m^2 is 14.5 mg (0.24 mg/kg bw/d) without PPE.

Inhalation exposure

A = surface area [cm²] (205 cm²)

The estimated mean event concentration is 57 mg/m^3 calculated from ConsExpo Web. After 45 mins (one event), the residual air concentration has declined to 1.3 mg/m^3 from the extrapolated graph. Therefore the systemic inhalation exposure during an 8-hour working day is calculated as follows:

Systemic inhalation uptake (mg/kg bw/d) = ((mean event concentration + remaining air conc) x inhalation rate x total exposure duration)/body weight

Where:

```
Event concentration calculated from ConsExpo Web = 58.3 \text{ mg/m}^3
Inhalation rate = 1.25 \text{ m}^3/\text{h}
Total exposure duration = 8 \text{ h}
```

Body weight

S	Summary table: systemic	exposure from profess	sional uses	
Tier/PPE	Estimated inhalation uptake (mg/kg bw/d)	Estimated dermal uptake (mg/kg bw/d)	Estimated total uptake (mg/kg bw/d)	
1 (no PPE) 9.72		0.24	9.6	
Please see Annex 3.2.1 for detailed calculations				

= 60 kg

<u>Scenario 3: Secondary exposure: bystander inhalation of volatilised residues</u> <u>during disinfection in industrial/manufacturing cleanrooms</u>

Inhalation exposure may occur to re-entry of professional bystanders (e.g. cleanroom staff or manufacturing assistant) in manufacturing/industrial settings where surface disinfection is performed. The inhalation exposure will be in the same order of magnitude as for the person who disinfected the surfaces. In a worst case scenario it is assumed that the bystander stays for 8 hours in the room where surface disinfection is performed. Therefore the level of inhalation exposure of a bystander is estimated to be equivalent or lower compared to the professional user applying/using the products.

Consequently the dermal exposure of bystanders follows the assumption that there is a very low probability for direct contact to freshly disinfected surfaces, and a very short duration of dermal exposure if casual contact happens. Furthermore due to the high vapour pressure of the active substance, dermal exposure by contact (hands) with treated surfaces is likely to be very low due to rapid evaporation. Therefore the dermal exposure is considered to be negligible.

Combined exposure

Combined exposure has been considered for a professional using the RTU liquid to refill a trigger sprayer and subsequently disinfect hard surfaces via trigger sprayer and wiping. Therefore the combined exposures for professional primary exposure that could occur during a particular day are as follows:

- Scenario 1: Primary exposure: professional user pouring RTU liquid from a 5 L container for refilling trigger sprayer
- Scenario 2: Primary exposure: a technician performs routine disinfection of small surfaces e.g. equipment and work stations as part of their working procedure*

Summary table: combined systemic exposure from professional uses					
Scenarios combined	Tier	PPE	Estimated total uptake (mg/kg bw/day)		
Scenario 1: Dermal and inhalation exposure during pouring of RTU liquid to refill trigger sprayer	1	No PPE	0.092		
Scenario 2: Dermal and inhalation exposure during routine disinfection of small surfaces by a technician*	1	No PPE	<9.6		

Combined total exposure	29 7
Combined total exposure	\ \ J. /

*It should be noted that the worst-case representative product considered in this scenario is the wet wipes. As the exposure from the use of trigger sprayer is within the risk envelope, this scenario represents the worst-case combined exposure for a professional re-filling a trigger sprayer and subsequently disinfects surfaces with a trigger sprayer and wiping.

General public exposure

Products of the Contec IPA Product Family are intended for use by professional users only.

Exposure of bystanders

Contec IPA Product Family is intended for use in manufacturing/industrial cleanrooms where members of the general public e.g. children will be excluded from entry. As such no general public bystander exposure is foreseen.

List of scenarios for PT4

In the absence of specific guidance on surface disinfection in food preparation settings and product specific information regarding the scale of use/sizes of area being treated, the UK CA has followed the approach in the propan-2-ol PT 4 Assessment Report for surface disinfection by wiping in industrial food processing setting. It is noted that the Assessment Report considered surface disinfection of professional food preparation settings in canteen/kitchen however the applicant has clarified that Contec IPA Product Family is intended only for use in the professional preparation of food in industrial setting e.g. food processing industry.

Scenario number	Time, or secondar, experience				
4.	Disinfection of hard surfaces in food preparation settings	, , ,	Professional		
5.	Inhalation of volatilised residues in industrial/manufacturing settings	on of volatilised Secondary exposure: professional labeled bystander exposure to volatilised bystander exposure to volatilised bystander exposure to volatilised labeled bystander exposure to volatilised bystander exposure to volatilised labeled bystander exposure exposure to volatilised bystander exposure ex			

Industrial exposure

Products of the Contec IPA Product Family are intended for use by professional users only.

Professional exposure

<u>Scenario 4: Primary exposure: professional user disinfecting surfaces in professional food preparation settings e.g. food processing settings</u>

The application rate to surfaces is 50 ml/m^2 via the liquid trigger spray and 8 wipes/m^2 for impregnated wet wipes containing 5 - 26 ml product per wipe. The worst-case scenario is therefore represented by the use of the products through wet wipes at an application rate of 208 ml/m^2 ($26 \text{ ml } \times 8 \text{ wipes}$). Thus the following risk assessment considers a professional user disinfecting surfaces using impregnated wet wipes and exposure from the use of trigger sprayer is within the risk envelope.

The scenario outlined in the PT 4 Assessment Report considers a professional user disinfecting a cutting and packaging machine in a food processing setting.

Work rate = Area to be disinfected per event

 $(1 \text{ m}^2 - \text{cutting machine})$

(3.6 m² – packaging machine)

Application Duration = 5 min per event Exposure duration = 120 min per event Frequency = 4 events per day

The UK CA notes that the Assessment Report mentions the temperature in the working hall [food processing setting] is regulated to $0-5^{\circ}$ C and that the calculations were performed at 4° C. Given the high vapour pressure of propan-2-ol, volatility is likely to be significantly affected by the temperature. However it is unclear from the calculations whether they were performed at a lower temperature nor is the vapour pressure of propan-2-ol at this lower temperature stated. As such, the UK CA has performed calculations based on the vapour pressure of propan-2-ol of 5780 Pa at 25°C.

Description of Scenario 4

A professional user disinfects surfaces in professional food preparation setting e.g. food processing settings. The disinfection pattern considered in the Assessment Report has been followed to estimate the exposure from the use of this product family.

In accordance with ConsExpo cleaning factsheet, inhalation exposure has been calculated using ConsExpo Web exposure to vapour: evaporation model (increasing area) for cleaning with wet tissue (surrogate for wet wipes).

	Parameters	Value	
	Concentration of active in wet wipe	62.9% w/w	
	Inhalation rate	1.25 m³/hr	
	Molecular weight	60 g/mol	
	Vapour pressure (EU-agreed value (LoEP January 2015))	5780 Pa at 25°C	
	Product density	0.786 g/ml (applicant information)	
Inhalation	Molecular weight matrix ¹	18 g/mol	
exposure	Mass transfer rate	Thibodeaux method	
	Exposure duration (Assessment Report)	120 mins	
	Room volume (Assessment Report)	300 m ³	
	Ventilation (Assessment Report)	20/hr	
	Amount of product to treat 4.6 m ² (10 wipes) ²	204 g (10 x 26 ml = 260 ml)	
	Release area (Assessment Report)	4.6 m ²	
	Application duration (Assessment Report)	5 mins	

¹The molecular weight of the matrix is based on the formulation. Please see the confidential Annex Section 3.6.1 for details of the full formulation.

Calculations for Scenario 4

<u>Dermal exposure</u>

ConsExpo Cleaning Product Factsheet⁴ (p.64) estimated that during wiping with wet tissues, 1.4% of the total liquid of the wet tissue remains on the surface of the inner hand. Therefore the amount of product available for dermal exposure when using one wet wipe is 0.29 g (26 ml/wipe x 0.786 g/ml x 1.4%) or 183 mg propan-2-ol. Following the assumption in the propan-2-ol Assessment Report and ECHA Methodology for disinfection by single use wiping tissue (p. 106), one palm of hand (205 cm 2) is exposed during wiping; the time of evaporation is calculated according to TGD on risk assessment, App. I, App. If:

 $T = mRTK/M\beta$ pA

² The required number of wipes is calculated based on the nearest 0.5 m² area treated.

t = time [s]

m = mass of propan-2-ol on surface [mg] (183 mg)

 $R = \text{gas constant [J K-1 mol-1] (8.314 J K}^{-1} \text{ mol}^{-1})$

T = skin/surface temperature [K] (303.15 K)

K = conversion factor (36000)

 $M = \text{molar mass [g mol-1] (60.1 g mol^{-1})}$

 β = mass transfer coefficient [m h-1], for calculation see TGD (8.7 m h-1)

p = vapour pressure of the pure substance [Pa] (7600 Pa (30°C))

A = surface area [cm²] (205 cm²)

According to this equation the evaporation time is 20.4 seconds. Therefore for a worst case scenario it is assumed that the total amount evaporates at once after this time interval. The absorption/dermal flux rate for propan-2-ol in a 70% w/w aqueous dilution is 0.85 mg/cm²/h (EU-agreed value (LoEP January 2015)). During the time interval of 20.4 seconds on the skin surface of palm of one hand 205 cm², this result in a total amount absorbed per day is 0.99 mg.

Therefore the total amount absorbed from the use of 37 wet wipes to disinfect 18.4 m^2 per day $(4.6 \text{ m}^2 \text{ x 4})$ is 37 mg (0.61 mg/kg bw/d) without PPE.

Inhalation exposure

The estimated mean event concentration is 11 mg/m^3 calculated from ConsExpo Web. After 120 mins, the residual air concentration is $1.4 \times 10^{-7} \text{ mg/m}^3$. As the residual air concentration at the end of each event is essentially negligible (0.000001% of the mean air concentration), this has not been added to the mean event concentration to estimate worker inhalation exposure during a working day. Therefore the systemic inhalation exposure during an 8-hour working day is calculated as follows:

Systemic inhalation uptake (mg/kg bw/d) = (mean event concentration x inhalation rate x total exposure duration)/body weight

Where:

Event concentration calculated from ConsExpo Web $= 11 \text{ mg/m}^3$ Inhalation rate $= 1.25 \text{ m}^3/\text{h}$ Total exposure duration = 8 hBody weight = 60 kg

Summary table: estimated exposure from professional uses					
Scenario/PPE	Estimated inhalation uptake (mg/kg bw/d)	Estimated dermal uptake (mg/kg bw/d)	Estimated total uptake (mg/kg bw/d)		
4 (no PPE)	1.83	0.61	2.44		
Please see Annex 3.2.1 for detailed calculations					

Combined scenarios

Combined exposure has been considered for a professional using the RTU liquid to refill a trigger sprayer and subsequently disinfect hard surfaces via trigger sprayer and wiping.

Therefore the combined exposures for professional primary exposure that could occur during a particular day are as follows:

- Scenario 1: Primary exposure: professional user pouring RTU liquid from a 5 L container for refilling trigger sprayer
- Scenario 4: Primary exposure: professional user disinfecting surfaces in professional food preparation settings e.g. food processing settings *

Summary table: combined systemic exposure from professional uses						
Scenarios combined	Tier PPE/clothing		Estimated total uptake (mg/kg bw/day)			
Scenario 1: Dermal and inhalation exposure during pouring of RTU liquid to refill trigger sprayer	1	No PPE	0.092			
Scenario 4: Dermal and inhalation exposure during disinfection of hard surfaces*	1	No PPE	<2.44			
Combined total exposure		·	<2.53			

^{*}It should be noted that the worst-case representative product considered in this scenario is the wet wipes. As the exposure from the use of trigger sprayer is within the risk envelope, this scenario represents the worst-case combined exposure for a professional re-filling a trigger sprayer and subsequently disinfects surfaces with a trigger sprayer and wiping.

<u>Scenario 5: Secondary exposure: professional bystander exposure to volatilised</u> residues during disinfection in professional food preparation setting

Inhalation exposure may occur to re-entry by professional bystanders (e.g. manufacturing assistant) in manufacturing/industrial food preparation settings where surface disinfection is performed. The inhalation exposure will be in the same order of magnitude as for the person who disinfected the surfaces. In a worst case scenario it is assumed that the bystander stays for 8 hours in the room where surface disinfection is performed. Therefore the level of inhalation exposure of a bystander is estimated to be equivalent or lower compared to the professional user applying the products.

Consequently the dermal exposure of bystanders follows the assumption that there is a very low probability for direct contact to freshly disinfected surfaces, and a very short duration of dermal exposure if casual contact happens. Furthermore due to the high vapour pressure of the active substance, dermal exposure by contact (hands) with treated surfaces is likely to be very low due to rapid evaporation. Therefore the dermal exposure is considered to be negligible.

General public exposure

Products of the Contec IPA Product Family are intended for use by professional users only.

Bystander exposure

Contec IPA Product Family is intended for use in controlled professional environments e.g. manufacturing/industrial and food preparation settings where members of the general public e.g. children will be excluded from entry. As such no general public bystander exposure scenarios are foreseen.

Dietary exposure

The formulation of the products of the family is similar to the representative formulation considered at active substance approval and therefore the same conclusion is applicable. As such no residues in food or feed are expected to arise from the use of the products of the family (PT 4) due to the high vapour pressure of the active substance (5780 Pa at 25°C).

Summary of exposure assessment

Scenarios and values to be used in risk assessment					
Scenario number	Exposed group	Tier/PPE	Estimated total uptake (mg/kg bw/d)		
Scenario 1: Primary exposure: professional user pouring RTU liquid from a 5 L container for refilling trigger sprayer	Professionals	No PPE	0.092		
Scenario 2: Primary exposure: a technician performs routine disinfection of small surfaces e.g. equipment and work stations as part of their working procedure	Professionals	No PPE	9.6		
Scenario 3: Secondary exposure: bystander inhalation of volatilised residues during disinfection in industrial/manufacturing cleanroom	Professional bystanders	N/A	<9.6		
Scenario 4: Primary exposure: professional user disinfecting surfaces in professional food preparation settings e.g. food processing settings	Professionals	No PPE	2.44		

Scenario 5: Secondary exposure: professional bystander exposure to volatilised residues during disinfection in professional food preparation setting	Professional bystanders	N/A	<2.44
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2.2.6.3 Risk characterisation for human health

Reference values to be used in Risk Characterisation

Reference	Study	NOAEL (LOAEL)	AF¹	Correction for oral absorption	Value
Professional workers AELacute/ Medium-term/ long-term	EU-agreed value (LoEP January 2015)	NOAEC – 200 ppm LOAEL 400 ppm for acute systemic (neurological) effects (over 8 hours), based on the deterioration of postural balance	3.8	-	17.9 mg/kg bw/d (52.6 ppm for 8 hours/d)
Professional workers Reference value for inhalation (proposed OEL)	EU-agreed value (LoEP January 2015)	-	-	-	200 ppm
General population AEL acute/medium/ long-term		NOAEC – 200 ppm LOAEL 400 ppm	6.4	-	10.7 mg/kg bw/d (31.25 ppm for 8 hours/d)
ARfD	Not required; no residues in food expected				
Not required; no residues in food expected					

¹ Default assessment factors of 6.4 for the general population and 3.8 for professional users were applied to account for intraspecies variability.

Note on local effects: The AEC $_{acute/medium/long-term}$ is assumed to also sufficiently cover local irritant effects in the eyes/airways.

Risk for industrial users

Products of the Contec IPA Product Family are intended for use by professional users only.

Risk for professional users PT 2

Systemic effects

Task/ Scenario	Tier	AEL mg/kg bw/d	Estimated uptake mg/kg bw/d	Estimated uptake/ AEL (%)	Acceptable (yes/no)
Scenario 1: Primary exposure: professional user pouring RTU liquid from a 5 L container for refilling trigger sprayer	1 (no PPE)	17.9	0.092	<1	Yes
Scenario 2: Primary exposure: a technician performs routine disinfection of small surfaces e.g. equipment and work stations as part of their working procedure	1 (no PPE)	17.9	9.6	54	Yes
Scenario 3: Secondary exposure: bystander inhalation of volatilised residues during disinfection in industrial/manufacturing setting e.g. cleanroom	-	-	<9.6	<54	Yes

Combined scenarios

Combined exposure has been considered for a professional using the RTU liquid to refill a trigger sprayer and subsequently disinfect hard surfaces via trigger sprayer and wiping in manufacturing/industrial cleanrooms.

Scenarios combined	Tier	MEL mg/kg	uptake ma/ka	IIIDTAKE/	Acceptable (yes/no)
1 and 2	1 (no PPE)	17.9	9.7	54	Yes

Local effects

Contec IPA Product Family is classified for Eye Irritant 2 (H319: Causes serious eye irritation) and STOT SE 3 (H336: May cause drowsiness or dizziness). The propan-2-ol Assessment Report informs that the AEC for professional users of 52.6 ppm for 8 hours/day (converted to a systemic AEL of 17.9 mg/kg bw/d) also sufficiently covers local irritant effects in the eyes/airways. As all professional exposure scenarios were below the AEL, no further consideration for local effects is necessary.

Conclusion

On the basis of the risk assessment and considering local effects have been taken into account in the setting of the AEL, exposure is within acceptable limits for professional users without PPE.

Risk for bystanders PT 2

Contec IPA Product Family is intended for use in controlled professional environments of manufacturing/industrial cleanrooms where members of the general public e.g. children will be excluded from entry. As such no general public bystander exposure scenarios are foreseen.

Risk for professional users PT 4

Systemic effects

Task/ Scenario	Tier	AEL mg/kg bw/d	Estimated uptake mg/kg bw/d	Estimated uptake/ AEL (%)	Acceptable (yes/no)
Scenario 4: Primary exposure: professional user disinfecting surfaces in professional food preparation settings e.g. food processing settings	1 (no PPE)	17.9	2.44	14	Yes
Scenario 5: Secondary exposure: professional bystander exposure to volatilised residues during disinfection in professional food processing setting	-	-	<2.44	<14	Yes

Combined scenarios

Combined exposure has been considered for a professional using the RTU liquid to refill a trigger sprayer and subsequently disinfect hard surfaces via trigger sprayer and wiping in food preparation settings.

Scenarios combined	Tier	AEL ma/ka	uptake ma/ka	IIINTAKE/	Acceptable (yes/no)
1 and 4	1 (no PPE)	17.9	2.53	14	Yes

Local effects

Contec IPA Product Family is classified for Eye Irritant 2 (H319: Causes serious eye irritation) and STOT SE 3 (H336: May cause drowsiness or dizziness). The propan-2-ol Assessment Report informs that the AEC for professional users of 52.6 ppm for 8 hours/day (converted to a systemic AEL of 17.9 mg/kg bw/d) also sufficiently covers local irritant effects in the eyes/airways. As all professional exposure scenarios were below the AEL, no further consideration for local effects is necessary.

Conclusion

On the basis of the risk assessment and considering local effects have been taken into account in the setting of the AEL, exposure is within acceptable limits for professional users without PPE.

Risk for bystanders PT 4

Contec IPA Product Family is intended for use in controlled professional food preparation setting (industrial/institutional) where members of the public will be excluded. As such no general public exposure scenarios are foreseen.

Risk for consumers via residues in food

The formulation of the products of the family is similar to the representative formulation considered at active substance approval and therefore the same conclusion is applicable. No residues in food or feed are expected to arise from the use of the products of the family (PT 4) due to the high vapour pressure of the active substance (5780 Pa at 25°C). Furthermore, reference values for dietary intake (ADI or ARfD) have not been derived for propan-2-ol (2015 Assessment Report).

Risk characterisation from combined exposure to several active substances or substances of concern within a biocidal product

Contec IPA Product Family contains only one active substance and no substances of concern relevant for human health. Therefore the consideration of combined exposure is not required.

2.2.7 Risk assessment for animal health

Primary and secondary exposure of animals to the biocidal product family is not expected via the intended uses.

2.2.8 Risk assessment for the environment

The Contec IPA Product Family products are a pour-on liquid (intended only for refilling a trigger spray), a liquid trigger spray, and pre-saturated wipes. The trigger spray and wipe products are available ready-to-use (RTU) and each contain 70 % (v/v) propan-2-ol as the active substance. The products are intended for disinfectant use in cleanrooms for biotechnology, pharmaceutical, manufacture of medical devices, healthcare industries and other critical life science applications; industrial food and feed preparation areas. The application rate to surfaces is 50 ml/ $\rm m^2$ via the liquid trigger spray products and, based on a maximum application rate of 2 wipes per square metre and a maximum of 38 ml per wipe, a worst case has been assumed of 76 ml/ $\rm m^2$ for the wipe product. The applicant states that the products are formulated by mixing 700 ml of propan-2-ol (density 0.785 g/ml) with 300 ml of water, giving 0.5502 kg/l of active substance in the products.

The UK CA notes that $0.785 \text{ g/ml} \times 700 \text{ ml} = 549.5 \text{ g}$ - then correcting for the percentage technical material of 99 % w/w gives a value of $(549.5 \times 100/99) = 555 \text{ g/l}$ (0.555 kg/l) which has been used in the risk assessment.

Environmental and ecotoxicological data specific to the Contec IPA Product Family are not available. Instead, information required for the environmental risk assessment is based on the active substance, propan-2-ol, which is available in the EU Assessment Report (January 2015). This approach is justified because the type of formulation and inert substances used in the products are not expected to affect the environmental properties or ecotoxicological profile of propan-2-ol. Data generated with unformulated propan-2-ol can be extrapolated to the formulated product and environmental properties of the products do not need to be specifically tested.

2.2.8.1 Effects assessment on the environment

The product family contains only one active substance and no substances of concern. Therefore all toxicity data can be obtained from the Assessment Report. The PNECs are summarised below:

Aquatic = 2.82 mg a.s./L
Sediment = 2.41 mg/kg ww sediment
Sewage = 10 mg/L
Soil = 0.496 mg/kg ww soil
Log Pow = 0.05 and no surface tension properties

Not B or T

Information relating to the ecotoxicity of the biocidal product which is sufficient to enable a decision to be made concerning the classification of the product is required

No additional data submitted. The agreed LoEP for propan-2-ol has been used in the risk assessment.

Further Ecotoxicological studies

No additional data submitted. The agreed LoEP for propan-2-ol has been used in the risk assessment.

Effects on any other specific, non-target organisms (flora and fauna) believed to be at risk (ADS)

No additional data submitted. The agreed LoEP for propan-2-ol has been used in the risk assessment.

Supervised trials to assess risks to non-target organisms under field conditions

No additional data submitted. The agreed LoEP for propan-2-ol has been used in the risk assessment.

Studies on acceptance by ingestion of the biocidal product by any non-target organisms thought to be at risk

No additional data submitted. The agreed LoEP for propan-2-ol has been used in the risk assessment.

Secondary ecological effect e.g. when a large proportion of a specific habitat type is treated (ADS)

No additional data submitted. The agreed LoEP for propan-2-ol has been used in the risk assessment.

Foreseeable routes of entry into the environment on the basis of the use envisaged

The products in the family are applied indoors to hard surfaces in industrial and institutional areas; no direct exposure of soil or surface waters is expected.

The emission pathways considered in the risk assessment include indirect emissions of the formulation to surface water (and sediment), the STP itself, groundwater and air. The main emission pathway during use will be to air since propan-2-ol is known to evaporate completely within a short time due to its relatively high vapour pressure.

The deposition of propan-2-ol to soil and subsequent movement to groundwater has also been considered.

Further studies on fate and behaviour in the environment (ADS)

The fate and behaviour of the active substance propan-2-ol is not expected to be altered by the co-formulants in the products. As such, the data submitted for the active

substance is considered sufficient to cover all endpoints for environmental fate and behaviour. No further product specific studies are required.

Leaching behaviour (ADS)

The products of the family are formulated products and not treated articles. The products are not intended for addition to a matrix or impregnation into another material. As such, testing for leaching behaviour is neither relevant nor required according to the Guidance on the Biocidal Product Regulation, Volume IV, Part A.

Testing for distribution and dissipation in soil (ADS)

The fate and behaviour of the active substance propan-2-ol is not expected to be altered by the co-formulants in the products. As such, the data submitted for the active substance is considered sufficient to cover all endpoints for environmental fate and behaviour. No further product specific studies are required.

Testing for distribution and dissipation in water and sediment (ADS)

The fate and behaviour of the active substance propan-2-ol is not expected to be altered by the co-formulants in the products. As such, the data submitted for the active substance is considered sufficient to cover all endpoints for environmental fate and behaviour. No further product specific studies are required.

Testing for distribution and dissipation in air (ADS)

The fate and behaviour of the active substance propan-2-ol is not expected to be altered by the co-formulants in the products. As such, the data submitted for the active substance is considered sufficient to cover all endpoints for environmental fate and behaviour. No further product specific studies are required.

If the biocidal product is to be sprayed near to surface waters then an overspray study may be required to assess risks to aquatic organisms or plants under field conditions (ADS)

No additional data submitted. The agreed LoEP for propan-2-ol has been used in the risk assessment.

If the biocidal product is to be sprayed outside or if potential for large scale formation of dust is given then data on overspray behaviour may be required to assess risks to bees and non-target arthropods under field conditions (ADS)

No additional data submitted. The agreed LoEP for propan-2-ol has been used in the risk assessment.

2.2.8.2 Exposure assessment

The Contec IPA Product Family biocidal products are used under PT 2 and PT 4.

Exposure assessments are conducted separately for each product type. It is stated in the propan-2-ol BPC opinion that the distribution of releases between air and waste water at a ratio of 90 % and 10 % should be re-evaluated at product authorisation stage. In the absence of any further information and in view of the high similarity between this product and the representative product we assume that the 90 : 10 ratio is still appropriate for this propan-2-ol containing product.

Product type 2

The product type is PT 2 (disinfectants not intended for direct application to humans or animals, mainly healthcare applications).

Products in the Contec IPA Product Family are ready-for-use, alcoholic disinfectant products which are intended to be used as private and public health area disinfectants. Their use is restricted to routine hard surface disinfection in industrial/manufacturing settings (e.g. cleanrooms, for biotechnology, pharmaceutical, manufacture of medical devices, healthcare industries and other critical life science applications; industrial food and feed preparation areas). They may be used in hospital pharmacy clean-room facilities where drugs are compounded.

These products are used in strictly controlled, occupational settings where all members of the public will be excluded. The products are applied to surfaces as such (i.e. undiluted) and no post-application rinsing of surfaces is required.

There is no general public (consumer use) of these products.

The products are produced in different formulation types: wipes, trigger sprays and ready-to-use liquids for pouring into trigger spray bottles. The products can therefore be applied by wiping or spraying, and will be used according to different disinfection regimes.

The products in this family are intended for professional use only and are provided ready-to-use. It was considered that any losses from the pouring of liquid into trigger spray bottles would be negligible and could be considered as being included in a number of the worst case scenarios assessed.

General information for PT 2

Assessed PT	PT 2
Assessed scenarios	Based on the environmental emission scenarios for Product Type 2: Private and public health area disinfectants and other biocidal products, the following scenarios were identified as relevant to generate local emission values (i.e. emission rate to wastewater (standard STP)).
	Scenario 1 : "PT 2-med sector-rooms furniture/objects" (emission scenario for calculating the release of disinfectants used for sanitary purposes in hospitals based on the amount

of solution of disinfection used on a day).
Scenario 2: "PT 2-industrial areas" (emission scenario for
calculating the releases of disinfectants used in industrial
areas).
Environmental Emission Scenarios for Product Type 2:
Private and public health area disinfectants and other
biocidal products
Emission estimations were calculated for the three relevant
scenarios identified listed above, using ECHA's environmental
emission scenarios for PT 2 Excel spread sheet.
Calculated based on the ECHA Guidance on Environmental
Risk Assessment, Volume IV, Part B + C.
Tier 1 assessment plus argumentation
No
Industrial use
Exposure based on worst case assumptions and relevant
guidance documents
Consumption based modelling carried out (as found to be
worst case in the propan-2-ol AR).

Emission estimation for PT 2

Scenario 1: "PT 2-med sector-rooms furniture/objects" (emission scenario for calculating the release of disinfectants used for sanitary purposes in hospitals based on the amount of solution of disinfection used on a day)

Although sanitary uses have not been requested by the applicant, the default value for brushes has been considered in this scenario to reflect the use of this product family in a hospital on a daily basis.

The emission rate to wastewater (standard STP) was calculated based on the amount of active substance in the products (0.555 kg/I) used in a hospital to disinfect brushes. The propan-2-ol AR states that the distribution of releases between air and wastewater occurs at a ratio of 90 % and 10 % respectively, this has been applied in the emissions assessment.

For all other parameters the default values were used (*i.e.* amount of water with active substance).

This scenario is most applicable to the trigger spray products, and may also be considered protective for the use of wipes.

Input parameters for calculating the local emission					
Input	Value	Unit	Remarks		
Scenario 1: "PT2-med sector-rooms furniture/objects" (emission scenario for calculating the release of disinfectants used for sanitary purposes in hospitals based on the amount of solution of disinfection used on a day).					
Active substance in product	0.555	kg/l	Calculated based on products composed of 70% propan-2-ol		
Fraction released to wastewater	0. 10	-	ESD default		
Fraction released to air	0. 90	-	ESD default		
Emission to wastewater (standard STP)	1.39	kg/d	Output		
Emission to air	12.5	kg/d	Output		

Scenario 2: "PT 2-industrial areas" (emission scenario for calculating the releases of disinfectants used in industrial areas)

The local release was calculated based on an extreme worst case application rate of the biocidal wipes of 0.076 l/m^2 (38 ml per wipe and 2 wipes per square metre - Scenario $\frac{3}{2}$ 2a) and the more realistic application rate of 50 ml/ m² for the trigger spray products (Scenario 2b). A concentration of active substance in the products of 0.555 g/l and "Small scale application (RTU)" were also assumed.

The propan-2-ol AR states that the distribution of releases between air and wastewater occurs at a ratio of 90 % and 10 % respectively. The fraction released to wastewater was therefore set to 0.1 to derive the release to wastewater and the release to air calculated from this value (i.e. (Elocal_{water} \times 10) \times 0.9).

It was agreed in the TAB version 1.3 for PT 2 that the treated area for a RTU product could be reduced to 25 m^2 (when product is applied by means of trigger spray or wipes), this has been used in the exposure calculation and can be assumed to be protective for a number of smaller repeat applications made to a localised area during the day.

All other parameters used were defaults (number of applications per day and fraction of substance disintegrated during or after application (before release to the sewage system)).

Input parameters for calculating the local emission- Small scale use (2a)- Wipes					
Input	Value	Unit	Remarks		
Scenario 3a: "PT 2-industrial areas" (emission scenario for calculating the releases of disinfectants used in industrial areas).					
Application rate of biocidal product	0.076	I/m²	Based on 76 ml/m² worst case application rate (wipes)		
Concentration of active substance in the product	555	g/l	Based 70 % v/v propan-2-ol		
Surface area to be disinfected	25	m ²	TAB 1.3 RTU small scale		

Input parameters for calculating the local emission- Small scale use (2a)- Wipes					
Input	Value	Unit	Remarks		
			applications		
Fraction released to wastewater	0.1	-	Assessment report 2015		
Fraction released to air	0.9	-	Assessment report 2015		
Local release to wastewater	0.105	kg/d	Output		
Local release to air	0.949	kg/d	Output		

Input parameters for calculating the local emission- Small scale use (3 2b)- Trigger spray					
Input Value Unit Remarks					
Scenario 3b: "PT 2-industrial areas"	-		nario for calculating the		
releases of disinfectants used in indu	ıstrial aı	reas).			
Application rate of biocidal product	0.050	I/m²	Based on 50 ml/m² worst case application rate (wipes)		
Concentration of active substance in the product	555	g/l	Based 70 % v/v propan-2-ol		
Surface area to be disinfected	25	m²	TAB 1.3 RTU small scale applications		
Fraction released to wastewater	0.1	-	Assessment report 2015		
Fraction released to air	0.9	-	Assessment report 2015		
Local release to wastewater	0.069	kg/d	Output		
Local release to air	0.624	kg/d	Output		

Product type 4

The product type is PT 4 (disinfectants used in food and feed areas).

Products in the Contec IPA Product Family are also used to disinfect hard surfaces and equipment which are used in the preparation of food in industrial settings (e.g. in food, beverage and dairy industries). Products in the Contec IPA Product Family intended for food and feed area disinfection are all ready-for-use, alcoholic disinfectants which are used indoors and are effective against bacteria and yeast. These products are used in controlled, occupational settings in areas where members of the public will be excluded. The products are applied to surfaces as such and no post-application rinsing of surfaces is required.

There is no general public (consumer use) of these products.

The products are produced in different formulation types: wipes, trigger sprays and ready-to-use liquids for pouring into trigger spray bottles. The products can therefore be applied by wiping or spraying, and will be used according to different disinfection regimes.

The products in this family are intended for professional use only and are provided ready-to-use. It was considered that any losses from the pouring of liquid into trigger spray bottles would be negligible and could be considered as being included in a number of the worst case scenarios assessed.

General information for PT 4

Assessed PT	PT 4
Assessed scenarios	Based on ECHAs environmental emission scenarios for Product Type 4: Disinfectants used in food and feed areas, the following two scenarios were identified as relevant to generate local emission values (i.e. effluent concentration of active substance in the effluent of the on-site STP, influent concentration of active substance in the off-site STP and local emission to waste water). Scenario 1: "PT 4-FDM industries" (assessment of entire plants e.g. breweries, dairies, beverage processing plants). Scenario 2: "PT 4-large scale kitchens, etc" (emission scenario for calculating the release of disinfectants used in large scale catering kitchens, canteens, slaughterhouses and butcheries).
ESD(s) used	Environmental Emission Scenarios for Product Type 4: Disinfectants used in food and feed areas
Approach	Emission estimations were calculated for the four relevant scenarios identified listed above, using ECHA's environmental emission scenarios for PT 4 Excel spread sheet.
Distribution in the environment	Calculated based on the ECHA Guidance on Environmental Risk Assessment, Volume IV, Part B + C.
Groundwater simulation	Tier 1 assessment only.
Confidential Annexes	No
Life cycle steps assessed	Industrial use
Remarks	Exposure based on worst case assumptions and relevant guidance documents

Emission estimation for PT 4

Scenario 1: "PT 4-FDM industries" (assessment of entire plants e.g. breweries, dairies, beverage processing plants).

An influent concentration of active substance to the off-site STP of 0.003 mg/l was calculated using the "PT 4-FDM industries" model (assessment of entire plants e.g. breweries, dairies, beverage processing plants) from ECHAs "Environmental Emission Scenarios for Product Type 4: Disinfectants used in food and feed areas". This scenario is based on a number of default parameters and the assumption that 143 kg/yr of propan-2-ol is used in a brewery (using the agreed ECHA excel sheet).

All other parameters were default parameters (i.e. amount of biocidal active substance used per year in the local plant, number of emission days per year, fraction released to waste water, capacity of STP and dilution factor in surface water). For the fraction released to waste water, the default value of 1000 % loss was assumed as a worst case, as assumed in the AR (relating to a surface treatment).

Input parameters for calculating the local emission						
Input	Value	Unit	Remarks			
Scenario 1: "PT 4-FDM industries" (assessment of entire plants e.g. breweries, dairies, beverage processing plants).						
Active ingredient applied in breweries	Propan-2-ol	N/A	Pick-list			
Amount of active substance used per year in local plant (Qai)	143	kg/yr	Default from ESD table 6			
Number of emission days per year (Temission)	231	d/yr	Default			
Fraction released to wastewater (Fwater)	0.1	-	AR default			
Effluent concentration from the onsite STP Ceffluent = Clocal water	3.43E-03	mg/l	Output			
Influent concentration of active substance in the off-site STP Cinfluent	3.10E-02	mg/l	Output			
Emission to Air (Elocalair)	5.57E-01	kg/d	Output			

Scenario 2: "PT 4-large scale kitchens, etc" (emission scenario for calculating the release of disinfectants used in large scale catering kitchens, canteens, slaughterhouses and butcheries).

The local release to wastewater was calculated using the "PT 4-large scale kitchens, etc" model (emission scenario for calculating the release of disinfectants used in catering kitchens, canteens, slaughterhouses and butcheries) from the ESD "Environmental Emission Scenarios for Product Type 4: Disinfectants used in food and feed areas".

Following item 54 in the TAB version 1.3, a default surface area of 50 m^2 can be applied for a small scale RTU product such as a trigger spray or wipes. This value is also protective for the use in slaughterhouses where a default area of 10 m^2 is assumed (for small scale use).

The propan-2-ol AR states that the distribution of releases between air and waste water occurs at a ratio of 90 % and 10 % respectively. The fraction released to waste water was therefore set to 0.1 to derive the release to waste water and the release to air calculated from this value (i.e. (Elocal $_{water}$ x 10) x 0.9). All other parameters were default parameters (volume to be disinfected and number of applications per day).

The application rate was calculated based on the worst case of $0.076 \text{ l/m}^2 \times 555 \text{ g/l} = 42.2 \text{ g/m}^2$ for wipes (Scenario 2a) and more typical usage of $0.05 \text{ l/m}^2 \times 555 \text{ g/l} = 28 \text{ g/m}^2$ for the spray products (Scenario 2b).

Input parameters for calculating the local emission - Small scale use (2a)- Wipes					
Input	Value	Unit	Remarks		
Scenario 2a: "PT 4- Emission scenario for calculating the release of disinfectants used in large scale catering kitchens, canteens, slaughterhouses and butcheries.					
Type of application	Wiping	-	Pick-list		
Size of the area treated	50	m ²	TAB 1.3 RTU small scale applications		
Fraction released to waste water (Fwater)	0.1	-	Assessment Report 2015		
Fraction released to air (Fair)	0.9	-	Assessment Report 2015		
Application rate of the active substance	42.2	g/m²	Based on a worst-case application rate of 304 ml/m ²		
Local release to waste water	0.211	kg/d	Output		
Local release to air	1.90	kg/d	Output		

Input parameters for calculating the local emission - Small scale use (2b)- Trigger spray					
Input	Value	Unit	Remarks		
Scenario 2b: "PT 4- Emission scenario for calculating the release of disinfectants used in large scale catering kitchens, canteens, slaughterhouses and butcheries.					
Type of application	Spraying	-	Pick-list		
Size of the area treated	50	m ²	TAB 1.3 RTU small scale applications		
Fraction released to waste water (Fwater)	0.1	-	Assessment Report 2015		
Fraction released to air (Fair)	0.9	-	Assessment Report 2015		
Application rate of the active substance	27.8	g/m²	Based on a worst-case application rate of 50 ml/m ²		
Local release to waste water	0.139	kg/d	Output		
Local release to air	1.25	kg/d	Output		

It has then been assumed from the release to air that deposition to soil can take place.

Fate and distribution in exposed environmental compartments

Identif	Identification of relevant receiving compartments based on the exposure pathway									
	Fresh- water	Freshwater sediment	Sea- water	Seawater sediment	STP	Air	Soil Ground- water		Other	
PT 2	PT 2									
Scenario 1	Yes	Yes	No	No	Yes	Yes	Yes	Yes	No	
Scenario 3	Yes	Yes	No	No	Yes	Yes	Yes	Yes	No	
PT 4	PT 4									
Scenario 1	Yes	Yes	No	No	Yes	Yes	Yes	Yes	No	
Scenario 2	Yes	Yes	No	No	Yes	Yes	Yes	Yes	No	

Input parameters (only set values) for calculating the fate and distribution in the environment (PT 2 & PT 4)							
Input	Value	Unit	Remarks				
Molecular weight	60.09	g/mol	EU-agreed value (LoEP January 2015)				
Melting point	-89.5	°C	EU-agreed value (LoEP January 2015)				
Boiling point	82.5	°C	EU-agreed value (LoEP January 2015)				
Vapour pressure at 25°C (12°C)	5780 (2302)	Pa	EU-agreed value (LoEP January 2015)				
Water solubility at 25°C (12°C)	1000000 (831846)	mg/l	EU-agreed value (LoEP January 2015)				
Log Octanol/water partition coefficient	0.05	Log 10	EU-agreed value (LoEP January 2015)				
Organic carbon/water partition coefficient (K _{OC})	3.3	l/kg	EU-agreed value (LoEP January 2015)				
Henry's Law Constant at 25°C (12°C)	0.80 (0.383)	Pa/m³/mol	EU-agreed value (LoEP January 2015)				
DT ₅₀ for degradation in air	3.1	days	EU-agreed value (LoEP January 2015)				
Biodegradability	Readily biodegradable	N/A	EU-agreed value (LoEP January 2015)				

Fate and distribution in the STP PT 2 and PT 4 (using SimpleTreat 3.0)					
Commontment	Percentage [%]	Domonyka			
Compartment	All scenarios	Remarks			
Air	0.3	EU-agreed value (LoEP January 2015)			
Water	12.5	EU-agreed value (LoEP January 2015)			
Sludge	0	EU-agreed value (LoEP January 2015)			
Degraded in STP	87.1	EU-agreed value (LoEP January 2015)			

Resulting local emission to relevant environmental compartments						
Compartment	Local emission (Elocal _{water}) [kg/d]	Local emission (Elocal _{air}) [kg/d]				
PT 2						
Scenario 1	1.39	12.5				
Scenario 2a	0.105	0.949				
Scenario 2b	0.069	0.624				
PT 4						
Scenario 1*	6.19E-02*	0.557				
Scenario 2a	0.211	1.90				
Scenario 2b	0.139	1.25				

^{*}Not relevant as C_{eff} and C_{inf} calculated in scenario equations Back calculated from C_{inf} value of 0.031 mg/ l

Calculated PEC values

Please refer to Annex 3.2.2 for the calculation of PECs following Volume IV Part B + C, guidance on environmental risk assessment.

Summary table on calculated PEC values								
	PEC _{STP} [mg/l]	PEC _{water}	PEC _{sed} [mg/kg _{wwt}]	PEC _{soil} [mg/kg _{wwt}]	Tier 1 PEC _{GW} [µg/l]	PEC _{air} [mg/m³]		
PT 2								
Scenario 1	8.67E-02	8.67E-03	NC	3.42E-04	1.94	3.47E-03		
Scenario 2a	6.59E-03	6.59E-04	NC	2.60E-05	0.148	2.64E-04		
Scenario 2b	4.34E-03	4.34E-04	NC	1.71E-05	0.097	1.74E-04		
PT 4								
Scenario 1	3.87E-03	3.43E-03* 3.87E-04#	NC	1.52E-05	0.086	1.55E-04		
Scenario 2a	1.32E-02	1.32E-03	NC	5.19E-05	0.295	5.28E-04		
Scenario 2b	8.67E-03	8.67E-04	NC	3.42E-05	0.194	3.47E-04		

^{*}Value from on-site STP; #Value from off-site STP; NC Not calculated; Where a and b refer to emissions from wiping and spraying respectively

PEC_{sediment} values were not calculated as there are no sediment effects data available for propan-2-ol. It is assumed that the levels of risk to the sediment will be the same as the values calculated for surface water.

Primary and secondary poisoning

Primary poisoning

The proposed uses of the products preclude any risk of primary poisoning.

Secondary poisoning

There is a low risk of secondary poisoning from the use of propan-2-ol products as the active substance has a very low potential for secondary poisoning. Propan-2-ol has a low log Kow of 0.05 and the calculated bioaccumulation factors (BCFs) for fish and earthworms are 0.22 and 0.85 l/kg respectively. Consequently, the approach taken in the Assessment Report was followed and no PNEC or PEC values have been calculated.

2.2.8.3 Risk characterisation

Atmosphere

<u>Conclusion</u>: As stated in the propan-2-ol AR there is a potential for long range environmental transport, however effects on stratospheric ozone and acidification are not expected as propan-2-ol does not contain halogens, nitrogen or sulphur and is not listed as an ozone depleting substance. Inhalation studies with mammals also indicate that adverse effects are not expected to occur to terrestrial mammals.

As there are no effects data available it was accepted in the AR that the risk to air can be considered to be acceptable.

Sewage treatment plant (STP)

Summary table on calculated PEC/PNEC values				
	PEC/PNEC _{STP}			
PT 2				
Scenario 1	0.082			
Scenario 1	0.009			
Scenario 2a	<0.001			
Scenario 2b	<0.001			
PT 4				
Scenario 1	<0.001*			
Scenario 2a	0.001			
Scenario 2b	<0.001			

^{*}Value from on-site STP; Where a and b refer to emissions from wiping and spraying respectively

<u>Conclusion</u>: PEC/ PNEC ratios for the STP are below the value of 1 for all scenarios indicating an acceptable level of risk for all proposed uses.

Aquatic compartment

Summary table on calculated PEC/PNEC values						
	PEC/PNECwater	PEC/PNEC _{sed}				
PT 2						
Scenario 1	0.003	NC				
Scenario 2a	< 0.001	NC				
Scenario 2b	< 0.001	NC				
PT 4						
Scenario 1	0.001*	NC				
Scenario 1	< 0.001#	IVC				
Scenario 2a	< 0.001	NC				
Scenario 2b	< 0.001	NC				

^{*}Value from on-site STP; #Value from off-site STP; NC Not calculated; Where a and b refer to emissions from wiping and spraying respectively

<u>Conclusion</u>: As the PEC/ PNEC ratios for the aquatic compartment are below the trigger value of 1 for all scenarios, acceptable risks to the aquatic compartment are indicated. As effects data have not been provided for the sediment compartment it is accepted that the level of risk to this compartment can be considered to be the same as that for the aquatic compartment.

Terrestrial compartment

Summary table on Calculated PEC/PNEC val		
	PEC/PNEC _{soil}	
PT 2		
Scenario 1	< 0.001	
Scenario 2a	< 0.001	
Scenario 2b	< 0.001	
PT 4		
Scenario 1	< 0.001	
Scenario 2a	< 0.001	
Scenario 2b	< 0.001	

Where a and b refer to emissions from wiping and spraying respectively

<u>Conclusion</u>: As the PEC/ PNEC ratios for the terrestrial compartment are below the trigger value of 1 for all scenarios, acceptable risks to the terrestrial compartment are indicated.

Groundwater

When a tier 1 approach to the calculation of groundwater levels is applied following guidance in Vol IV Part B, acceptable levels are predicted for some scenarios. This porewater calculation is generally assumed to be a conservative approach as it does not

take into account any lateral movement processes, degradation within the soil or removal by volatilisation from the soil.

Summary table on Tier 1 Porewater concentration					
PECIocal _{soil, porewater} [µg/I]					
PT 2					
Scenario 1	1.94				
Scenario 2a	0.148				
Scenario 2b	0.097				
PT 4					
Scenario 1	0.086				
Scenario 2a	0.295				
Scenario 2b	0.194				

Where a and b refer to emissions from wiping and spraying respectively

Following discussions at WG-VII-2018 it was agreed that the following argument forms an acceptable weight of evidence approach to support FOCUS PEARL not offering an appropriate tier 2 refinement for these proposed uses of propan-2-ol.

For the environmental risk assessment, dry and wet deposition, expressed as DEPTtotal_{ann}, is assumed to be the main emission pathway to the soil and subsequently to the groundwater compartment due to the high volatility of propan-2-ol. The DEPTtotal_{ann} is calculated by use of the OPS model (as described in the Guidance on the BPR IV ENV B, 2015), which assumes that the major fraction (90%) of the applied propan-2-ol is released to the ambient air and subsequently deposited in close vicinity (within a radius of 1000 m) to the source of emission. In the case of propan-2-ol this assumption represents an unrealistic worst-case and it can be considered highly unlikely that the assumed magnitude of exposure truly occurs under relevant field conditions.

- The FOCUS PEARL model was developed for the determination of groundwater concentrations related to the application of plant protection products (PPP) on agricultural land. Accordingly, the model assumptions for the nine locations rely on e.g. soil properties that are representative for agriculturally used areas in Europe. The unlimited applicability of the model for the very diverse field of biocidal applications is thus questionable. For biocidal applications where the release of active substances to the environment is related to the application of sewage sludge or manure/slurry to agricultural land, the applicability of FOCUS PEARL might be given. In the present case, where the products of the BPF are used in urban areas where a direct exposure to the urban environment is assumed, the model assumptions of FOCUS PEARL may not be accurate and the results of such a refinement should be evaluated with caution. The same applies, when FOCUS PEARL is used for the groundwater assessment of volatile compounds, for which the model is might not be suitable, since it might overestimate the leaching rate to the groundwater for such compounds. Consequently, the results of the

refined groundwater assessment with FOCUS PEARL must also be considered as an unrealistic worst-case.

This discussion is supported by the conclusion at the 21st BPC meeting that if not all nine scenarios show a safe use and the applicability of the models for the substance evaluated can be questioned, a qualitative approach could be applied using expert judgement in a weight of evidence approach.

An acceptable risk of propan-2-ol to groundwater is therefore expected.

Primary and secondary poisoning

Primary poisoning

The proposed uses of the products preclude any risk of primary poisoning.

Secondary poisoning

There is an acceptable level of risk of secondary poisoning from the use of propan-2-ol products as the active substance has a very low potential for secondary poisoning. Propan-2-ol has a low log Kow of 0.05 and the calculated bioaccumulation factors (BCFs) for fish and earthworms are 0.22 and 0.85 l/kg respectively. Consequently, no risk of secondary poisoning is foreseen and no PNEC or PEC values have been calculated.

Mixture toxicity

As the biocidal products contain only one active substance, an assessment of mixture toxicity is not required.

Aggregated exposure (combined for relevant emmission sources)

Propan-2-ol is used in a number of biocidal PTs (1, 2 and 4) and has a number of other non-biocidal uses. An aggregated exposure assessment was performed in the propan-2-ol Assessment Report (2015) and is reproduced below.

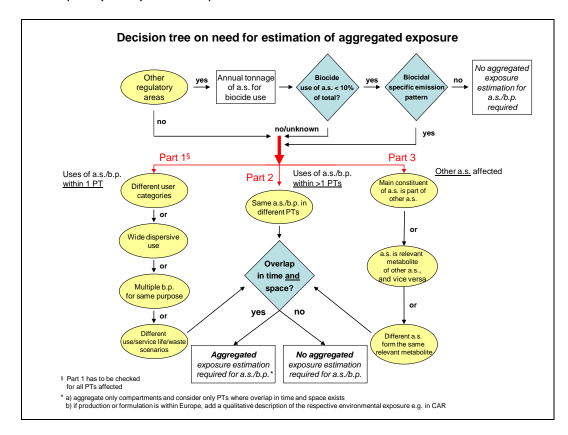


Figure 1: Decision tree on the need for estimation of aggregated exposure

"Propan-2-ol is notified for Annex I inclusion in PT 1, 2, and 4. For all mentioned PTs, DE is RMS. The respective CA reports consider the following uses: PT 1 - skin and hand disinfectant in hospitals; PT 2 - disinfection of rooms, furniture and objects in the sanitary sector; PT 4 - assessment of small-scale applications (spraying of surfaces) / industrial kitchens / meat processing industry. As b.p. containing propan-2-ol are used in a wide dispersive way, an aggregated environmental exposure assessment may be reasonable. According to the "Decision tree on the need for estimation of aggregated exposure" (BIP6.7 Decision Tree Agg Expo), the requirement for aggregated exposure estimations was checked for propan-2-ol. In summary, it has been concluded that no aggregated exposure assessment for propan-2-ol has to be performed as the biocidal uses of propan-2-ol is less than 10 % of the total tonnage produced and no specific biocidal emission patterns are identified."

On this basis further consideration of aggregated exposure is not necessary.

Overall conclusion on the risk assessment for the environment of the product

A number of scenarios have been used to assess this biocidal product family to cover the liquid trigger spray (applied at 50 ml/m^2) and RTU wipes (applied at 76 ml/m^2).

When RTU wipes and trigger sprays have been considered, the assessment has followed the agreements made within the propan-2-ol AR and assumed a 90 % loss to air and 10 % loss to drain following application.

Application Rate

Two application rates have been considered, a worst case value of 76 ml/m² for the wipes 50 ml/m² for the trigger spray.

The UK CA notes that the value used in risk assessment of 76 ml/m² is based on a number of worst case assumptions, notably the total amount of liquid (38 ml) contained in the largest wipe in the product family is assumed to be lost on wiping. This is unrealistic as there will always be an amount of liquid retained in a wipe following use.

RTU Trigger spray (Meta SPC 1)

Acceptable levels of risk to all environmental compartments have been demonstrated for the proposed uses of the RTU trigger spray product.

RTU wipes (Meta SPC 2)

Acceptable levels of risk to all environmental compartments have been demonstrated for the proposed uses of the RTU wipes.

2.2.9 Measures to protect man, animals and the environment

Please see section 2.1.4.

2.2.10 Assessment of a combination of biocidal products

The products of the Contec IPA Product Family are not intended to be used in combination with other biocidal products.

2.2.11 Comparative assessment

A comparative assessment is not required as the active substance is not a candidate for substitution.

3 Annexes

3.1 List of studies for the biocidal product family

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
	2016	Determination of Storage Stability and Shelf Life Specification Data for a Formulation containing 70% Isopropanol stored at various temperatures for Two Years, in Compliance with Good Laboratory Practice.	David Norris Analytic al Laborat ories Ltd.	DNA346 5	Yes	No	Yes		3.1-
	2016	Cosmetic Regulation Stability Test Report	Microbio logical Solution s Limited	15/0155 1/2	No	No	Yes	.III	3.1-
	2016	Expert Statement on the EU CLP Charachteristic s of a Propan- 2-ol/water Solution (70:30): Explosivity, Oxidising Properties, Self-Reacting, Self-Heating and Corrosive to Metals	TSGE Consulti ng Limited	TSGE_1 6-070- 01_IPA_ CLP_2.0	No	No	Yes	Contec Cleanro om (UK) Limited,	4.1
	2016	Validation of the Methods of	David Norris	DNA346 6	Yes	No	Yes		5.1

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
		Determination of a Formulation containing Isopropanol, in Compliance with Good Laboratory Practice.	Analytic al Laborat ories Ltd.						
	2014	Microbiological Analysis Based on EN 1276 (2009) Quantitative suspension test for the evaluation of bactericidal activity of chemical disinfectants and antiseptics (Phase 2 / Step 1)	MGS Laborat ories Limited	TRA- 2014- 091-01	No	No	Yes		6.7- 01
	2014	Microbiological Analysis Based on EN 1650 (2008) + A1:2013 Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of fungicidal or yeasticidal activity of chemical disinfectants and antiseptics used in food, industrial, domestic and	MGS Laborat ories Limited	TRA- 2014- 089-01	No	No	Yes		6.7- 02

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
		institutional areas – Test method and requirements (Phase 2, Step 1)							
	2013	EN 13697, Chemical disinfectants and antiseptics - Quantitative non-porous surface test for the evaluation of bactericidal and/or fungicidal activity of chemical disinfectants used in food, industrial, domestic and institutional areas – Test method and requirements without mechanical action (Phase 2, Step 2)	Abbott Analytic al	13C.079 SB.PAL	No	No	Yes		6.7-03
	2014	Microbiological Analysis Based on EN 13727 (2012) Chemical Disinfectant and Antiseptics - Quantitative Suspension Test for the Evaluation of Bactericidal Activity in the Medical Area –	MGS Laborat ories Limited	TRA- 2014- 090-01	No	No	Yes		6.7- 04

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
		Test Method and Requirements (phase 2 / step 1)							
	2006	Test Report BS EN 13624:2003 Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of fungicidal activity of chemical disinfectants for instruments used in the medical area – Test method and requirements (phase 2, step 1)	BluScien tific Test Data	N/A	Yes	No	Yes		6.7- 05
	2015	Test for the evaluation of bactericidal activity on non-porous surfaces with mechanical action in the medical area BSEN 16615:2015	Microbio logical Solution s Limited	15/0460 6-1	No	No	Yes		6.7- 06
	2015	Test for the evaluation of bactericidal activity on non-porous surfaces with mechanical action in the	Microbio logical Solution s Limited	15/0460 6-3	No	No	Yes		6.7- 07

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
		medical area BSEN 16615:2015							
	2014	EN 14348 (2005) Quantitative suspension test for the evaluation of tuberculocidal activity of chemical disinfectants in the medical area including instrument disinfectants. (Phase 2 / Step 1)	MGS Laborat ories Limited	TRA- 2014- 110-01	No	No	Yes		6.7- 08
	2013	EN 14561, Chemical disinfectants and antiseptics - Quantitative carrier test for the evaluation of bactericidal activity of chemical disinfectants used in the medical area – Test method and requirements (phase 2, step 2)	Abbott Analytic al	13C.079 CB.PAL	No	No	Yes		6.7- 09
	2013	EN 14561, Chemical disinfectants and antiseptics - Quantitative carrier test for the evaluation of bactericidal	Abbott Analytic al	13C.079 CVrMr.P AL	No	No	Yes		6.7-10

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
		activity of chemical disinfectants for instruments used in the medical area – Test method and requirements (phase 2, step 2)							
	2016	Quantitative carrier test for evaluation of fungicidal or yeasticidal activity for instruments used in the medical area BSEN 14562:2006	Microbio logical Solution s Limited	J000181	No	No	Yes		6.7-11
	2017	Chemical disinfectants and antiseptics — Quantitative carrier test for the evaluation of mycobactericid al or tuberculocidal activity of chemical disinfectants used for instruments in the medical area BS EN 14563:2008	Microbio logical Solution s Limited	J000181	No	No	Yes		6.7-12
	2016	BS EN 13624:2013 Quantitative	Microbio logical Solution	J000181	No	No	Yes		6.7- 13

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
		suspension test for the evaluation of bactericidal / fungal activity in the medical area	s Limited						
	2016	Quantitative non-pourous surface test for the evaluation of bactericidal / fungal activity of chemical disinfectants	Microbio logical Solution s Limited	J000188	No	No	Yes		6.7- 14
	2016	Test Report: EN 14476 2013 + A1 2015 Chemical disinfectants and antiseptics - Quantitative suspension test for the evaluation of virucidal activity in the medical area- Test method and requirements (Phase 2/Step 1)	BluTest Laborat ories Ltd	N/A	Yes	No	Yes		6.7- 15
	2016	Test for the evaluation of bactericidal activity on non-porous surfaces with mechanical action in the medical area BSEN 16615:2015	Microbio logical Solution s Limited	16/0087 2-2	No	No	Yes	Contec Cleanro om (UK) Limited	6.7-16

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
	2016	Test for the evaluation of bactericidal activity on non-porous surfaces with mechanical action in the medical area BSEN 16615:2015	Microbio logical Solution s Limited	16/0087 2-1	No	No	Yes	Contec Cleanro om (UK) Limited	6.7- 17
	2016	Test for the evaluation of bactericidal activity on non-porous surfaces with mechanical action in the medical area BSEN 16615:2015	Microbio logical Solution s Limited	16/0087 2-1 RETEST	No	No	Yes	Contec Cleanro om (UK) Limited	6.7-18
	2016	Test for the evaluation of bactericidal activity on non-porous surfaces with mechanical action in the medical area BSEN 16615:2015	Microbio logical Solution s Limited	16/0087 2-3	No	No	Yes	Contec Cleanro om (UK) Limited	6.7- 19
	2016	Test for the evaluation of bactericidal activity on non-porous surfaces with mechanical action in the medical area BSEN 16615:2015	Microbio logical Solution s Limited	16/0087 2-3 RETEST	No	No	Yes	Contec Cleanro om (UK) Limited	6.7- 20

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
	2017	Investigation into the Effectiveness of FH55 When Tested in Accordance With: UKAS Accredited Method for BS EN 1276:2009 Chemical disinfectants and antiseptics – Quantitative suspension test for the evaluation of bactericidal activity of chemical disinfectants and antiseptics used in food, industrial, domestic and institutional areas	Microbio logy Dept Selden Researc h Limited	43	No	No	Yes		6.7-21
	2016	Investigation into the Effectiveness of FH55 When Tested in Accordance With: Method for BS EN 13697:2015 Chemical disinfectants and antiseptics – Quantitative non-porous surface test for the evaluation of bactericidal	Microbio logy Dept Selden Researc h Limited	3	No	No	Yes		6.7- 22

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
		and/or fungicidal activity of chemical disinfectants used in food, industrial, domestic and institutional areas – Test method and requirements without mechanical action							
	2017	Investigation into the Effectiveness of FH55 When Tested in Accordance With: Non-UKAS Accredited Method for BS EN 1650:2008 + A1:2013 Chemical disinfectants and antiseptics – Quantitative suspension test for the evaluation of fungicidal or yeasticidal activity of chemical disinfectants and antiseptics used in food, industrial, domestic and institutional areas	Microbio logy Dept Selden Researc h Limited	4	No	No	Yes		6.7-23

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
	2017	Investigation into the Effectiveness of FH55 When Tested in Accordance With: Non-UKAS Accredited Method for BS EN 13697:2015 Chemical disinfectants and antiseptics – Quantitative non-porous surface test for the evaluation of bactericidal and/or fungicidal activity of chemical disinfectants used in food, industrial, domestic and institutional areas – Test method and requirements without mechanical action	Microbio logy Dept Selden Researc h Limited	4	No	No	Yes		6.7-24
	2014	Certificate of Analysis	Food & Drug Analytic al Services Limited	8992 SBT170I W	No	No	Yes	Contec Cleanro om (UK) Limited	6.7- 25
	2014	Certificate of Analysis	Food & Drug Analytic al Services	8992	No	No	Yes	Contec Cleanro om (UK) Limited	6.7- 26

Autho r(s)	Year	Title	Testin g Compa ny	Report No.	GLP Stu dy (Ye s/N o)	Publis hed (Yes/ No)	Data Prote ction Claim ed (Yes/ No)	Data Owner	IUC LID Sec tion No.
			Limited						
	2021	Shelf-life stability study at 25°C for 24 months on the test item "cellulose/polye ster nonwoven wipes (9"x11") 70/30 IPA/DI water" (Product Code PSC20006)	Eurofins Biolab S.r.l.	2019/12 1AM	Yes			Contec Cleanro om (UK) Limited	
	2021	Shelf-life stability study at 25°C for 24 months on the test item "cellulose/polye ster nonwoven wipes (6"x9") 70/30 IPA/DI water" (Product Code SAT-C1- 7030/18)	Eurofins Biolab S.r.l.	2019/12 2AM	Yes			Contec Cleanro om (UK) Limited	

3.2 Output tables from exposure assessment tools

3.2.1 Human Health Exposure Output Tables

Scenario 1: Primary exposure: professional user pouring RTU liquid from a 5 L container for refilling trigger sprayer

ConsExpo Web Substance

Name Propan-2-ol Molecular weight 60 g/mol

Kow -

Product Name

Weight fraction substance 62.9 %

Population

Name

Body weight 60 kg

Frequency 1 per day

Description

Inhalation

Exposure model Exposure to vapour - Evaporation

Exposure duration 0.75 minute
Product amount 1970 g
Weight fraction substance 62.9 %

Room volume 1 m³
Ventilation rate 0.5 per hour

Inhalation rate 1.25 m³/hr

Application temperature 20 °C

Vapour pressure 5.78E+03 Pa
Molecular weight 60 g/mol
Mass transfer coefficient 20.1 m/hr
Release area mode Constant
Release area 20 cm²

Emission duration 0.25 minute

Product in pure form No

Molecular weight matrix 18 g/mol
Absorption model Fixed fraction

Absorption fraction 100 %

Dermal

Exposure model n.a. Absorption model n.a.

Oral

Exposure model n.a. Absorption model n.a.

Results for scenario pouring

Inhalation

Mean event concentration 6.9 mg/m³ Mean concentration on day of exposure $3.6 \times 10^{-3} \text{ mg/m}^3$ $3.6 \times 10^{-3} \text{ mg/m}^3$ Year average concentration External event dose 1.8×10^{-3} mg/kg bw 1.8×10^{-3} mg/kg bw External dose on day of exposure Internal event dose 1.8×10^{-3} mg/kg bw Internal dose on day of exposure 1.8×10^{-3} mg/kg bw/day Internal year average dose 1.8×10^{-3} mg/kg bw/day

Integrated

Internal event dose 1.8×10^{-3} mg/kg bw Internal dose on day of exposure 1.8×10^{-3} mg/kg bw/day Internal year average dose 1.8×10^{-3} mg/kg bw/day

Scenario 2: Primary exposure: a technician performs routine disinfection of small surfaces e.g. equipment and work stations as part of their working procedure

ConsExpo Web

Substance

Name Propan-2-ol Molecular weight 60 g/mol

Kow -

Product Name

Weight fraction substance 62.9 %

Population Name

Body weight 60 kg

Scenario Technician in cleanroom

Frequency 1 per day

Description

Inhalation

Exposure model Exposure to vapour - Evaporation

Exposure duration 45 minute

Product amount 30 g
Weight fraction substance 62.9 %
Room volume 55 m³
Ventilation rate 8 per hour
Inhalation rate 1.25 m³/hr

Application temperature 25 °C

Vapour pressure 5.78E+03 Pa
Molecular weight 60 g/mol
Mass transfer coefficient 20.1 m/hr
Release area mode Increasing
Release area 0.5 m²
Application duration 1 minute

Product in pure form No

Molecular weight matrix 18 g/mol Absorption model Fixed fraction

Absorption fraction 100 %

Dermal

Exposure model n.a. Absorption model n.a.

Oral

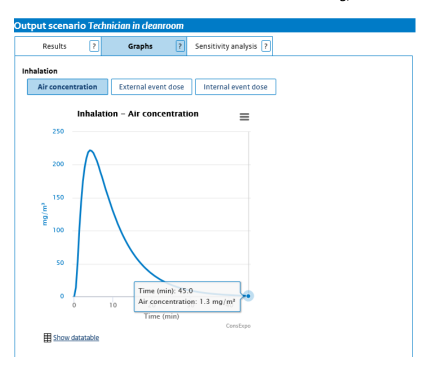
Exposure model n.a. Absorption model n.a.

Results for scenario Technician in cleanroom

Inhalation

Mean event concentration

5.7 mg/m³



Scenario 4: Primary exposure: professional user disinfecting surfaces in professional food preparation settings e.g. food processing settings

ConsExpo Web -

Substance

Name Propan-2-ol Molecular weight 60 g/mol

Kow -

Product Name

Weight fraction substance 62.9 %

Population Name

Body weight 60 kg

Scenario professional user in food processing

Frequency 1 per day

Description

Inhalation

Exposure model Exposure to vapour - Evaporation

Exposure duration 120 minute

Product amount 204 g
Weight fraction substance 62.9 %
Room volume 300 m³
Ventilation rate 20 per hour
Inhalation rate 1.25 m³/hr

Application temperature 25 °C

Vapour pressure 5.78E+03 Pa
Molecular weight 60 g/mol
Mass transfer coefficient 20.1 m/hr
Release area mode Increasing
Release area 4.6 m²
Application duration 5 minute

Product in pure form No

Molecular weight matrix 18 g/mol
Absorption model Fixed fraction

Absorption fraction 100 %

Dermal

Exposure model n.a. Absorption model n.a.

Oral

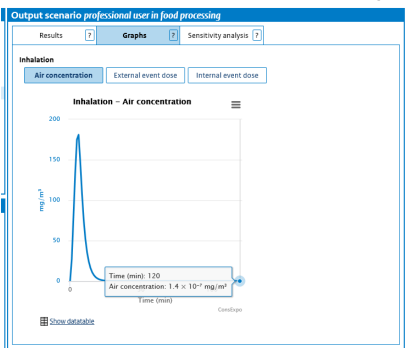
Exposure model n.a. Absorption model n.a.

Results for scenario professional user in food processing

Inhalation

Mean event concentration

1.1 x 10¹ mg/m³



3.2.2 Environmental Exposure Output Tables

PECair

Calculation of the emission to air has been made following the calculations laid out in the ECHA Guidance on ERA as follows.

$$Estp_{air} = Fstp_{air} \cdot Elocal_{water}$$
 (35)

$$Clocal_{air} = \max (Elocal_{air}, Estp_{air}) \cdot Cstd_{air}$$
 (42)

Where:

 $Fstp_{air} = 0.003$

Cstd_{air} = 2.78E-04 mg/m³ (default value)

Following the ECHA guidance on ERA the local concentration in the air has been calculated based on the maximum value of Estpair or Elocalair.

The fraction of active associated with aerosol particles (Fassaer) was then calculated following the equations in the ECHA guidance on ERA.

Using as a default $CON_{junge} \times SURF_{aer} = 10^{-4} Pa$ and a vapour pressure for propan-2-ol of 2302 Pa (corrected to 12°C).

$$Fass_{aer} = \frac{CONjunge \cdot SURF_{aer}}{VP + CONjunge \cdot SURF_{aer}}$$
(19)

Hence $Fass_{aer} = 1.73E-08$

The deposition flux can then be calculated summing the emission to air from the STP and direct emission from the indoor application using equations 43 and 44.

$$DEP total = \left(Elocal_{air} + Estp_{air}\right) \cdot \left(Fass_{aer} \cdot DEP std_{aer} + (1 - Fass_{aer}) \cdot DEP std_{gas}\right)$$
(43)

$$DEPtotal_{ann} = DEPtotal \cdot \frac{Temission}{365}$$
 (44)

If it is assumed that emissions take place throughout the year then T_{emission} is 365 and the annual average total deposition flux is the same as the total deposition per emission episode.

Where:

 $\begin{array}{ll} \text{DEPstd}_{\text{air}} & = 1.00\text{E-}002\\ \text{DEPstd}_{\text{gas}} & = 4.00\text{E-}04 \end{array}$

Scenario	[Elocal _{water} al		Local emission to air [Elocal _{air} kg/d]	Local concentration in air during emission episode [Clocalair kg/d]	DEPtotal (DEPtotal _{ann}) [mg m²/day]		
PT 2							
Scenario 1	1.39	4.16E-03	12.5	3.47E-03	5.00E-03		
Scenario 2a	0.105	3.16E-04	0.949	2.64E-04	3.80E-04		
Scenario 2b	0.069	2.08E-04	0.624	1.74E-04	2.50E-04		
PT 4							
Scenario 1	6.19E-02*	1.86E-04	0.557	1.55E-04	2.23E-04		
Scenario 2a	0.211	6.33E-04	1.90	5.28E-04	7.59E-04		
Scenario 2b	0.139	4.16E-04	1.25	3.47E-04	5.00E-04		

Where a and b refer to emissions from wiping and spraying respectively; *From back calculation from $Clocal_{inf}$ of 0.031~mg/ I

The aerial deposition flux per kg of soil, D_{air} is then derived from the total deposition flux (DEPtotal_{ann}) and is used in the calculation of PEC_{soil}.

$$D_{air} = \frac{DEPtotal_{ann}}{DEPTH_{soil} \cdot RHO_{soil}}$$
(52)

Scenario	Ecosystem and arable crops Aerial deposition flux Dair	Grassland Aerial deposition flux D _{air}
PT 2		
Scenario 1	1.47E-05	2.94E-05
Scenario 2a	1.12E-06	2.23E-06
Scenario 2b	7.35E-07	1.47E-06
PT 4		
Scenario 1	6.56E-07	1.31E-06
Scenario 2a	2.23E-06	4.47E-06
Scenario 2b	1.47E-06	2.94E-06

Where a and b refer to emissions from wiping and spraying respectively

Calculation of PEC_{STP} and PEC_{surface_water} (and PEC_{sediment})

Indoor scenarios

Taking the Elocal_{water} values previously calculated, the aquatic PEC values can be calculated using the following equations and default values taken from the ECHA guidance on ERA.

$$Clocal_{inf} = \frac{Elocal_{water} * 10^{6}}{EFFLUENT_{stp}}$$
122

(32)

$$Clocal_{eff} = Clocal_{inf} \cdot Fstp_{water}$$
(33)

$$Clocal_{water} = \frac{Clocal_{eff}}{(1 + Kp_{susp} * SUSP_{water} * 10^{-6}) * DILUTION}$$
(45)

As this product is intended for daily use, the $\mathsf{Clocal}_{\mathsf{eff}}$ will be used to assess the risk to microorganisms at STP.

Calculation of PEC_{stp} for Indoor application

Parameters	Nomenclature	Value	Unit	Origin
Effluent discharge rate of STP	EFFLUENT _{stp}	2000000	I	Default
Conc in untreated wastewater PEC _{STP}	Clocaleff		mg/l	Output
PT 2				
Scenario 1		8.67E-02	mg/l	Output
Scenario 2a		6.59E-03	mg/l	Output
Scenario 2b		4.34E-03	mg/l	Output
PT 4				
Scenario 1		3.87E-03	mg/l	Output
Scenario 2a		1.32E-02	mg/l	Output
Scenario 2b		8.67E-03	mg/l	Output

Where a and b refer to emissions from wiping and spraying respectively

Calculation of PEC_{surface water} for Indoor application

Parameters	Nomenclature	Value	Unit	Origin
Fraction directed to water by STP (Simpletreat)	Fstp _{water}	0.125	[-]	Input
Weight fraction organic carbon in suspended solids	FoC _{susp}	0.1		Default
Partition coefficient organic carbon - water	Koc	3.30	l/kg	Input
Partition coefficient solid – water in suspended matter	Kp _{susp}	0.330		Default
Concentration of suspended matter	$SUSP_{water}$	15	mg/l	Default
	DILUTION	10		Default
PEC _{surface} water	Clocalwater		mg/l	Output
PT 2				
Scenario 1		8.67E-03	mg/l	Output
Scenario 2a		6.59E-04	mg/l	Output
Scenario 2b		4.34E-04	mg/l	Output
PT 4				
Scenario 1		3.87E-04	mg/l	Output
Scenario 1 (on site STP to water)		3.43E-03	mg/l	Output
Scenario 2a		1.32E-03	mg/l	Output
Scenario 2b		8.67E-04	mg/l	Output

Where a and b refer to emissions from wiping and spraying respectively

Calculation of PECsoil and PECgroundwater

Calculation of Soil removal rate constants

Given that propan-2-ol is a volatile substance, volatilisation as an additional route of removal from soil was considered appropriate when calculating the PEC $_{\text{soil}}$. Following the ECHA guidance on risk assessment, Volume IV Part B- the total rate constant for removal is made up of several parts:

- Biodegradation rate constant (30 days based on propan-2-ol ready biodegradability) kbio_{soil}
- Volatalisation of substance from soil k_{volat}
- Leaching to deeper soil layer k_{leach}

As the soil concentration will be used to calculate pore water concentrations the third of the above rate constants (leaching) will not be considered in the following calculations. The overall rate constant is given by

$$K = K_{volat} + K_{leach} + Kbio_{soil}$$
(56)

The diffusive transfer from soil to air is estimated using the classical two film resistance model.

$$\frac{1}{K_{volat~i}} = \left(\frac{1}{Kasl_{air} * K_{air-watsr}/K_{soii-watsr}} + \frac{1}{Kasl_{soilair}}\right)$$

Where:

Kasl_{air} = partial mass transfer coeff. at air-side of the air soil-interface $\lceil m/d^{-1} \rceil$

(90.72 based on 1.05E-03 m s⁻¹ x 60 x 60 x 24 a correction of Vol IV Part

B+C of 2017 from the TGD of 2003)

Kasl_{soilair} = partial mass transfer coeff. at soil air-side of the air soil-interface $[m/d^{-1}]$

(calculated value see footnote at end of emissions)

 $K_{air-water}$ = air-water partitioning coefficient [m³/m⁻³] (1.62E-04) (see calculation below) $K_{soil-water}$ = soil-water partitioning coefficient [m³/m⁻³] (0.299) (see calculation below)

Depth_i = mixing depth of soil [m] (0.1 (grassland); 0.2 (agricultural soil))

 $K_{\text{volat i}}$ = rate constant for volatilisation from soil $I [d^{-1}]$

$$K_{cir-water} = \frac{HENRY}{R*TEMP}$$

Where:

HENRY = Henry's law constant [Pa/m³/mol⁻¹] (0.8 at 25°C corrected to 0.383 at 12°C

using equation 25 Volume IV Parts B + C 2017 as HENRY was derived

experimentally)

R = Gas constant $[Pa/m^3/mol^{-1}k^{-1}]$ (8.314)

TEMP = temperature at the air-water interface [k] (285) $K_{air-water}$ = air-water partitioning coefficient [m³/m⁻³] (7.02E-05)

 $\mathbf{K}_{soli-water} = Fair_{comp} * K_{air-water} + Fwater_{comp} + Fsolid_{comp} * rac{Kp_{comp}}{1000} * RHOsolid$

Where:

Fair_{comp} = fraction air in soil compartment $[m^3/m^{-3}]$ (0.2) Fwater_{comp} = fraction water in soil compartment $[m^3/m^{-3}]$ (0.2) Fsolid_{comp} = fraction solids in soil compartment $[m^3/m^{-3}]$ (0.6)

 Kp_{comp} = solids-water part. coeff. in soil compartment [I/kg] (0.066) (Koc x Foc_{soil};

 3.3×0.02)

RHOsolid = density of the solid phase $[kg/m^{-3}]$ (2500)

 $K_{\text{soil-water}}$ = soil-water partitioning coefficient [m³/m⁻³] (0.299)

As the mixing depth for soil varies between the different soil types, two values for kvolat can be calculated.

Ecosystem and arable soil kvolat = $1.19E-02 d^{-1}$ Grassland kvolat = $2.38E-02 d^{-1}$

When combined with the soil rate constant derived from the default value of 30 days the following values for k are given.

Ecosystem and arable soil $k = 4.30E-02 d^{-1}$

Grassland $k = 6.29E-02 d^{-1}$

These values are then used in equation 59 to calculate the deposition to soil following 10 years of use.

$$Cdep_{soil10}(0) = \frac{D_{air}}{k} - \frac{D_{air}}{k} \cdot e^{-365 \cdot 10 \cdot k}$$
(59)

Parameters	Ecosystem and arable crop Cdep _{soil10} (0)	Grassland Cdep _{soil10} (0)	Unit	Origin
PT 2				
Scenario 1	3.42E-04	4.67E-04	mg/kg	Output
Scenario 2a	2.60E-05	3.55E-05	mg/kg	Output
Scenario 2b	1.71E-05	2.33E-05	mg/kg	Output
PT 4				
Scenario 1	1.52E-05	2.08E-05	mg/kg	Output
Scenario 2a	5.19E-05	7.10E-05	mg/kg	Output
Scenario 2b	3.42E-05	4.67E-05	mg/kg	Output

Where a and b refer to emissions from wiping and spraying respectively

As only a negligible amount of a.s. reaches the sludge via STP the contribution from Csludge $_{\text{soil}}$ 10 (0) can be ignored and the concentration of propan-2-ol in soil can be assumed to come only via deposition.

$$C_{soil\ 10}\ (0) = Cdep_{soil\ 10}\ (0) + Csludge_{soil\ 10}\ (0)$$
 (63)

This initial soil concentration can then be used in equation 54 and 55 to calculate the average concentration in soil over 180 or 30 days.

$$Clocal_{soil} = \frac{D_{air}}{k} + \frac{1}{kT} \left[C_{soil}(0) - \frac{D_{air}}{k} \right] \cdot \left[1 - e^{-kT} \right]$$
(55)

Parameters	Ecosystem (30 days)- PEC _{soil}	Arable (180 days)	Grassland (180 days)	Unit	Origin
PT 2					
Scenario 1	3.42E-04	3.42E-04	4.67E-04	mg/kg	Output
Scenario 2a	2.60E-05	2.60E-05	3.55E-05	mg/kg	Output
Scenario 2b	1.71E-05	1.71E-05	2.33E-05	mg/kg	Output
PT 4					

Scenario 1	1.52E-05	1.52E-05	2.08E-05	mg/kg	Output
Scenario 2a	5.19E-05	5.19E-05	7.10E-05	mg/kg	Output
Scenario 2b	3.42E-05	3.42E-05	4.67E-05	mg/kg	Output

Where a and b refer to emissions from wiping and spraying respectively

The PEC_{soil} value taken from the arable PEC after 180 days has then been used to calculate the porewater concentration using the following equations from the ECHA guidance on ERA.

Where:

$$K_{soil-water} = Fair_{soil} \times K_{air-water} + Fwater_{soil} + Fsolid_{soil} \times (Kp_{soil}/1000) \times RHO_{solid}$$
 (24)

and

$$PEClocal_{soil,porewater} = (PEClocal_{soil} x RHO_{soil}) / (K_{soil-water} x 1000) (67)$$

Tier 1 Calculation of concentration in Porewater

Parameters	Nomenclature	Value	Unit	Origin				
Fraction of air in soil	Fair _{soil}	0.2	m _{air} ³/m _{soil} ³	Default				
Fraction of water in soil	Fwater _{soil}	0.2	m _{water} ³ /m _{soil} ³	Default				
Fraction of solids in soil	Fsolid _{soil}	0.6	$m_{\text{solid}}^3/m_{\text{soil}}^3$	Default				
Solids - water partitioning coefficient in soil	Kp _{soil}	0.066	l/kg	Calculated Foc _{soil} x Koc				
Density of the solid phase	RHO _{solid}	2500	kg _{solid} /m _{solid} ³	Default				
	Foc _{soil}	0.02	kgoc/kgsolid	Default				
Soil- water partitioning coefficient	K _{soil-water}	0.3		Output				
PT 2								
Scenario 1	PEClocal _{soil} , porewater	1.94E-03	mg/l	Output				
Scenario 2a	PEClocal _{soil} , porewater	1.48E-04	mg/l	Output				
Scenario 2b	PEClocal _{soil} , porewater	9.71E-05	mg/l	Output				
PT 4								
Scenario 1	PEClocal _{soil} , porewater	8.66E-05	mg/l	Output				
Scenario 2a	PEClocal _{soil} , porewater	2.95E-04	mg/l	Output				
Scenario 2b	PEClocal _{soil} , porewater	1.94E-04	mg/l	Output				

Where a and b refer to emissions from wiping and spraying respectively

Calculation of kasl _{soil}					
Parameter	Symbol	Unit	Value	Comment	Equation
Molecular weight	М	kg _c mol ⁻¹	6.01E-02	Input value	
Molecular diffusivity of the substance in the gas phase	DIFFgas	$m^2 d^{-1}$	1.22E+00	OUTPUT	equation 79
Molecular diffusivity of the substance in the water phase	DIFFwater	$m^2 d^{-1}$	1.26E-04	OUTPUT	equation 80
Volume fraction of water in the soil compartment	Fwater _{soil}	m _{water} 3.m _{soil} 3	0.20	Table 3	
Volume fraction of air in the soil compartment	Fair _{soil}	m _{air} 3 m _{soil} 3	0.20	Table 3	
Air-water partitioning coefficient	K _{air-water}	(-)	1.62E-04	OUTPUT	Equation 24
Volume fraction of solids in the soil compartment	Fsolid _{soil}	m _{solid} 2.m _{soil} -3	0.60	Table 3	
Partition coefficient solid-water in soil	Kp soil	L kg ⁻¹	6.60E-02	OUTPUT	Equation 26
Density of the solid phase	RHOsolid	kg m ⁻³	2500	Table 3	
Mass fraction of the substance in the water phase of the soil	FRw.soil	(-)	0.669	OUTPUT	Equation 76
Mass fraction of the substance in the solid phase of the soil	FRs.soil	(-)	0.331	OUTPUT	Equation 77
Mass fraction of the substance in the air phase of soil	FRa.soil	(-)	1.08E-04	OUTPUT	Equation 78
Average daily rate of wet precipitation	RAINRATE	m d ⁻¹	1.92E-03	BPR guidance value	
Fraction of precipitationthat penetrates into the soil	Finf _{soil}	(-)	2.50E-01	BPR guidance value	
Rate of advective downward transport of soil particles	SOLIDadv.soil	m d ⁻¹	5.48E-07	BPR guidance value	
Solid phase diffusion coefficient in the soil compartment	SOLIDdiff.soil	$m^2 d^{-1}$	5.50E-07	BPR guidance value	
Effective advection (with penetrating porewater)	Veff _{soil}	m d ⁻¹	1.61E-03	OUTPUT	Equation 74
Effective diffusion coefficient	Deff _{soil}	m^2d^{-1}	9.68E-05	OUTPUT	Equation 75
Rate constant for degradation in bulk soil	kdeg _{soil}	d ⁻¹	0.0231	INPUT	
Substance-dependent penetration depth	dp	m	1.08E-01	OUTPUT	Equation 73
Partial mass-transfer coefficient at soil side at the air-soil interface	kas I _{soil}	m d ⁻¹	2.50E-03	ОИТРИТ	Equation 72
Partial mass transfer coefficient at air side of the air-soil interface	kasl _{air}	m s ⁻¹	1.05E-03	BPR guidance value	
Partial mass transfer coefficient at air side of the air-soil interface	kasl _{air}	m d ⁻¹	90.72	BPR guidance value	
Mixing depth of soil i soil	DEPTH _{soil}	m	0.20	Table 9	
agric. soil	DEPTH _{soil}	m	0.20	Table 9	
grassland	DEPTH _{soil}	m	0.10	Table 9	

3.3 New information on the active substance

No new information on the active substance has been provided in support of this biocidal product family.

3.4 Residue behaviour

No residues in food or feed are expected to arise from the use of the products of the family (PT 4) due to the high vapour pressure of the active substance (5780 Pa at 25°C).

3.5 Summaries of the efficacy studies

Please see section 3.1 above and the efficacy section 2.2.5 of this PAR which summarises these data.

3.6 Confidential annex

Please refer to the separate document.

3.7 Other