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

PRODUCT ASSESSMENT REPORT OF A BIOCIDAL PRODUCT FOR NATIONAL AUTHORISATION APPLICATIONS



SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

SANITERPEN INSECTICIDE DK PAE

SANITERPEN INSECTICIDE EFFET CHOC

Product type 18

Deltamethrin

BC-TF001462-52

France

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1 GENERAL INFORMATION ABOUT THE PRODUCT APPLICATION

1.1 Applicant

Company Name:	ACTION PIN
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Country:	France
Telephone:	+33.5.5855.0700
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E-mail address:	Antoine.Andrieux@action-pin.fr

1.1.1 Person authorised for communication on behalf of the applicant

Name:	Antoine ANDRIEUX, Nicolas HUGUET
Function:	Regulation
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City:	Castets
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1.2 Proposed authorisation holder

Company Name:	ACTION PIN
Address:	Z.I. de Cazalieu 1078, route André Dupuy BP 30
City:	Castets
Postal Code:	40 260
Country:	France
Telephone:	+33.5.5855.0700
Fax:	+33.5.5855.0707
E-mail address:	Antoine.Andrieux@action-pin.fr
Letter of appointment for the applicant to represent the authorisation holder provided (yes/no):	no

1.3 Information about the product application

Application received:	20/12/2013
Application reported complete:	23/01/2014
Type of application:	Application for national authorisation (NA-APP)
Further information:	-

1.4 Information about the biocidal product

1.4.1 General information

Trade name:	SANITERPEN INSECTICIDE DK PRET A L'EMPLOI
Manufacturer's development code number(s), if appropriate:	Development code: 27214-B1 Development name: SANITERPEN INSECTICIDE effet choc
Product type:	18
Composition of the product (identity and content of active substance(s) and substances of concern; full composition see confidential annex):	0.02 % deltamethrin w/w
Formulation type:	Ready-for-use liquid (AL).
Ready to use product (yes/no):	No
Is the product the very same (identity and content) to another product already authorised under the regime of directive 98/8/EC (yes/no); If yes: authorisation/registration no. and product name: or Has the product the same identity and composition like the product evaluated in connection with the approval for listing of active substance(s) on to Annex I to directive 98/8/EC (yes/no):	No

1.4.2 Information on the intended use(s)

Overall use pattern (manner and area of use):	Insecticide and acaricide for animal's house and shelters IV.1 Indoor use IV.1.3.4 Animal houses/shelters The product is used indoors, in empty animals' houses and shelters (animals not intended for human consumption).
Target organisms:	Scientific name: <i>Dermanyssidae</i> : <i>Dermanyssus gallinae</i> , common name: poultry red mite, development stage: all. Scientific name: <i>Culicidae</i> : <i>Culex pipiens</i> , common name: mosquito, development stage: adults. Scientific name: <i>Muscidae: Musca domestica</i> ,

Category of users:	common name: house fly, development stage: all. Scientific name: <i>Ceratopogonidae</i> : <i>Ctenocephalides felis</i> , common name: cat flea, development stage: larvae and adults. Professional users
Directions for use including minimum and maximum application rates, application rates per time unit (e.g. number of treatments per day), typical size of application area:	 Method of application: spraying Detailed description of the method: The product is sprayed on the ground, walls, ceiling, and around windows and doors to control flies and mosquitoes, or on walls and floor to control fleas or red mites. The product is ready to use and then it is sprayed on relevant surfaces using the handheld trigger of the sprayer on the surfaces. After application the surfaces shall not be cleaned as the product is able to control the insect infestation for a 3- months period. Cleaning should only be done before the next application. Dry cleaning shall be considered and dusts shall be collected and treated as a waste. Relevant dose is 50 mL of un-diluted product for 1 m². According to the volume delivered by the spray, this corresponds to 37 sprayings / m².
	sufficient to control the populations for a 3-month period. The product can be applied maximum 3 times a year. Maximum 350 m2 of surface can be treated per application. If the infestation requires a treatment for a surface above this limit, another option shall be considered.
Potential for release into the environment (yes/no):	Yes
Potential for contamination of food/feedingstuff (yes/no)	No,
Proposed Label:	See docIIIB-12
Use Restrictions:	

For full details of the intended uses claimed by the applicant, please see annex 0a.

1.4.3 Information on active substance(s)

Active substance chemical name:	Deltamethrin (S)-α-cyano-3-phenoxybenzyl (1R, 3R)-3-(2,2- dibromovinyl)- 2,2-dimethylcyclopropanecarboxylate
CAS No:	52918-63-5
EC No:	258-256-6
Purity (minimum, g/kg or g/l):	Deltamethrin technical: 0.20% Deltamethrin pure: 0.02%
Inclusion directive:	Commission Directive 2011/81/EU
Date of inclusion:	20/12/2011
Is the active substance equivalent to the active substance listed in Annex I to 98/8/EC (yes/no):	Yes
Manufacturer of active substance(s) used in the biocidal product:	BAYER SAS, Environmental Science

Company Name:	SBM Formulation - Béziers
Address:	CS 621 ZI avenue Jean Foucault
City:	Béziers
Postal Code:	34535
Country:	France
Telephone:	
Fax:	
E-mail address:	

1.4.4 Information on the substance(s) of concern

The product contains no substance of concern.

The product contains preservatives currently in the review program of active substances for PT6 or already approved for this type of product. The data related to this preservative shall be taken into account in the evaluation after its approval at European level, at product's renewal stage.

1.5 Documentation

1.5.1 Data submitted in relation to product application

Identity, physicochemical and analytical method data

Physico-chemical properties studies and analytical methods on the biocidal product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI were provided by Action Pin.

Action Pin has access to physico-chemical properties studies and analytical methods on the active substance with a letter of Access of Bayer.

Efficacy data

- Laboratory study according to CEB N°135¹ method with the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI, on *Musca domestica* (house fly), *Culex pipiens* (mosquito), *Ctenocephalides felis* (cat flea) and *Demanyssus gallinae* (poultry red mite).
- Laboratory study according to CEB N°135 method with the product SANITERPEN INSECTICIDE EFFET CHOC PRET A L' EMPLOI, on *Musca domestica* (house fly).
- Laboratory study according to CEB n° 135 method with the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI on *Musca domestica* (house fly) and *Aedes aegypti* (mosquito).
- Semi-field study according to BSI 4172² method with the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI on *Aedes aegypti* (mosquito), *Culex pipiens* (mosquito) and *Ctenocephalides felis* (cat flea).
- Field test conducted in breeding premises according to CEB N°107³ method with the product SANITERPEN INSECTICIDE DK EXTRA, (0.2 % w/w deltamethrin) on *Musca domestica* (house fly).

Toxicology data

Studies on the product were submitted: acute oral and dermal studies, dermal and ocular irritation studies and skin sensitisation assays.

¹ CEB n°135 method: « Efficacy trial method for acaricide / insecticide products intended for surface treatment of storage facilities, processing and marketing if industrial products of animal or vegetal». ² BS 4172-21993 Hand held processing and marketing if industrial products of animal or vegetal.

² BS 4172-2:1993 Hand-held pressurized aerosol dispensers against houseflies. Method for determination of insecticidal efficiency

³ CEB n°107 method: « Efficacy trial method for insecticide products intended to control stable flies in breeding buildings »

Residue data

No studies were submitted.

Ecotoxicology data

No studies were submitted.

1.5.2 Access to documentation

Action Pin has access to physico-chemical properties studies and analytical methods on the active substance with a letter of Access of Bayer.

2 SUMMARY OF THE PRODUCT ASSESSMENT

2.1 Identity related issues

The source of the active substance used in the biocidal product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is the source used for annex I inclusion.

It has to be noted that the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI contains a substance (terpineol, CAS: 8000-41-7), even if presented as a solvent in the formulation, that is listed on the annex I of the regulation (EU) 1451/2007⁴ as an existing active substance. No active substance dossier has been submitted to support the approbation of this substance on the list of the active substances of the Union.

The laboratory study n°1948c/0615, 2015/11 was conducted with the products SANITERPEN INSECTICIDE DK PAE, (0.02 % w/w deltamethrin) and SANITERPEN INSECTICIDE DK PAE without deltamethrine. Based on the results, It can be concluded that the product without active substance presents no efficacy on its own at the recommended application rate and that terpineol is a co-formulant that has no insecticide properties in this product. More details are presented at the section 2.5).

2.2 Classification, labelling and packaging

2.2.1 Harmonised classification of the active substance deltamethrin

Classification - Regulation (EC) 1272/2008	
Class of danger	Acute tox. 3* Aquatic acute 1 Aquatic chronic 1
Hazard statements	H331: Toxic if inhaled H301: Toxic if swallowed H400: Very toxic to aquatic life (M=1 000 000) H410: Very toxic to aquatic life with long lasting effects (M=1 000 000)

⁴ Régulation of the European Comission of 4 December 2007 in the second phase of the 10-year work programme referred to in Article 16(2) of Directive 98//8 EC of the European Parliament of the Council concerning the placing of biocidal products on the market

2.2.2 Classification of the biocidal product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

Classification - Regulation (EC) 1272/2008			
Class of danger	Eye Irrit.2		
	Aquatic acute 1		
	Aquatic chronic 1		
Hazard statements	H319: Causes serious eye irritation		
	H400 : Very toxic to aquatic life		
	H410: Very toxic to aquatic life with long lasting effects		
Precautionary statements	P264: Wash thoroughly after handling.		
	P280: Wear protective gloves/protective clothing/eye protection/face protection.		
	P305 + P351 + P338: IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses if present and easy to do – continue rinsing.		
	P337 + P313: If eye irritation persists get medical advice/attention.		
	P273: Avoid release to the environment.		
	P391: Collect spillage		
	P501: Dispose of this material and its container to hazardous or special waste collection point, in accordance with local, regional, national and/or international regulation.		

2.2.3 Labelling of the biocidal product

The proposed labelling according to the CLP regulation is:

Pictograms:	
Signal words:	Warning
Class of danger	Eye Irrit. 2 Aquatic chronic 1
Hazard statements	H319: Causes serious eye irritation H410: Very toxic to aquatic life with long lasting effects Contains C(M)IT/MIT and OIT. May produce an allergic reaction.
Precautionary statements	 P264: Wash thoroughly after handling. P280: Wear protective gloves/protective clothing/eye protection/face protection. P305 + P351 + P338: IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses if present and easy to do – continue rinsing. P337 + P313: If eye irritation persists get medical advice/attention. P273: Avoid release to the environment. P391: Collect spillage.

P501: Dispose of this material and its container to hazardous
or special waste collection point, in accordance with local, regional, national and/or international regulation.

There is a sensitizing substance in the composition of the formulation for which a specific labelling is needed: Contains a mixture of 5-chloro-2-methyl-4-isothiazolin-3-one/2-methyl-2H-isothiazol-3-one (C(M)IT/MIT) and 2octyl-2H-isothiazol-3-one (OIT). May produce an allergic reaction

Substance chemical name	C(M)IT/MIT (mixture of 5-chloro-2-methyl-4-isothiazolin-3- one/2-methyl-2H-isothiazol-3-one)	
CAS No:	55965-84-9	
Relevant toxicological/ecotoxicological information:	May cause sensitisation by skin contact	

Substance chemical name OIT (2-octyl-2H-isothiazol-3-one)	
CAS No:	26530-20-1
Relevant toxicological/ecotoxicological information:	May cause sensitisation by skin contact

2.2.4 Packaging of the biocidal product

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is supplied in 750 mL Individual PET hand-operated trigger sprayer.

2.3 Physico/chemical properties and analytical methods

2.3.1 Active ingredient

2.3.1.1 Identity, origin of active ingredient

The source of the active substance used in the biocidal product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is the source used for Annex I inclusion to Directive 98/8/EC.

Manufacturer of the active substance:

Name:	Bayer CropScience AG
Address:	Alfred-Nobel Strasse 50
	40789 Monheim am Rhein
	Germany
Contact person:	Laurent Patty
Telephone:	+33 (0)472 85 46 85
Fax number:	+33 (0)472 85 47 75
E-mail address:	laurent.patty@bayercropscience.com

Manufacturer of the active substance: confidential data please refer to confidential PAR.

2.3.1.2 Physico-chemical properties

Physico-chemical properties of the active substance Deltamethrin have already been evaluated at EU level and are presented in the Competent Authority Report (CAR, 2011). The applicant Action Pin has a letter of access to these data.

2.3.1.3 Analytical method for determination of active ingredient and impurities in the technical active ingredient

Analytical method(s) for the determination of pure active substance Deltamethrin in the technical active substance as manufactured has already been performed and validated at EU level in the CAR (2011). The applicant Action Pin has a letter of access to these data.

Summary:

	Principle of method
Technical active substance as manufactured:	HPLC-UV and chiral HPLC-UV
Impurities in technical active substance:	-

2.3.1.4 Analytical method for determining relevant components and/or residues in different matrices

Analytical methods for the determination of residues of the active substance Deltamethrin in the different matrices have already been performed and validated at EU level in the CAR (2011). The applicant Action Pin has a letter of access to these data. Please refer to Annex 3 of this document.

Summary:

Soil (principle of method and LOQ)	LC-MS/MS using 1 transition LOQ 0.1 µg/kg			
Air (principle of method and LOQ)	GC-ECD for quantification and GC-MS for confirmation LOQ 0.27 μg/m ³			
Water (principle of method and LOQ)	Drinking water GC-ECD for quantification and confirmation LOQ 0.05 μg/L LC-MS/MS using 1 transition LOQ 5.9 ng/L GC-ECD for quantification and GC-MS/MS for confirmation LOQ 3 ng/L Surface water GC-ECD for quantification and GC-MS/MS for confirmation LOQ 3 ng/L			
Body fluids and tissues (principle of method and LOQ)	TissuesGC-ECD for quantification and confirmationLOQ 0.02 mg/kg for milk, eggs, meat, fat, liver and kidneyFluidsGC-MS for quantification and confirmationLOQ 200 µg/l for whole bloodGC-MSGC-MSmulti-methodforpyrethroidsfor			
	Paga			

	quantification
	LOQ 20 ng/L for whole blood
Food/feed of plant origin (principle of method	Not required as the intended uses will not result in
and LOQ for methods for monitoring purposes)	significant residues when the label instruction is followed.
	However two methods are provided which can be
	used in case of suspected contamination:
	GC-ECD for quantification
	LOQ 0.02 mg/kg for rice, flour, bread, meat, candy,
	butter, banana cream pie and lettuce
	LC-MS/MS
	LOQ 0.01 mg/kg for edible materials
	LOQ 0.05 mg/kg for non-edible materials for
	barley, broccoli, corn, melon, lettuce, olive, pepper,
	sugar beet, tobacco, tomato, wheat and zucchini
Food/feed of animal origin (principle of method	GC-ECD for quantification and confirmation
and LOQ for methods for monitoring purposes)	LOQ 0.02 mg/kg for milk, eggs, meat, fat, liver and
	kidney

2.3.2 Biocidal product

2.3.2.1 Identity, composition of the biocidal product, packaging

The biocidal product is not the same as the one assessed for the inclusion of the active substance in Annex 1 of Directive 98/8/EC.

The identification of the biocidal product is given in the table below:

Trade Name	SANITERPEN INSECTICIDE DK Prêt à l'emploi		
Manufacturer's development code number	Development code: 27214-B1 Development name: SANITERPEN INSECTICIDE effet choc		
Ingredient of preparation	Function Content (% w/w)		
Deltamethrin (CAS No.52918-63-5)	Active substance 0.02		
Formulants	Details on the composition of the product are included in the Confidential part (Doc. C1 and C2)		
Physical state of preparation	Liquid		
Nature of the preparation	AL (Any other Liquid)		

Manufacturer of the biocidal product:

Name:	Action PIN
Address:	Z.I. de Cazalieu
	1078, route André Dupuy
	40260 Castets
	France
Contact person:	Nicolas Huguet
Telephone:	+33 (0)558 55 07 00
E-mail address:	Nicolas.huguet@action-pin.fr
Plant location:	
Address:	448 route de l'océan
	40560 Vielle Saint Girons
	France
Contact person:	Antoine Andrieux
Telephone:	+33 (0)558 55 07 00

E-mail address:

antoine.andrieux@action-pin.fr

2.3.2.2 Physico-chemical properties

The tested product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is a ready-to-use any other liquid formulation (AL).

Bridging data for SANITERPEN INSECTICIDE DK PRET A L'EMPLOI from SANITERPEN INSECTICIDE DK: Explosive properties, oxidising properties, auto-flammability

An explosive properties study according to EC A.14 method, an oxidising properties study according to EC A.21 method and an auto-flammability study according to EC A.15 method were performed with the product SANITERPEN INSECTICIDE DK, a product similar to SANITERPEN INSECTICIDE DK PRET A L'EMPLOI but containing 0.20% w/w Deltamethrin (concentrate).

The comparison between SANITERPEN INSECTICIDE DK and SANITERPEN INSECTICIDE DK PRET A L'EMPLOI will demonstrate that the explosive properties, oxidising properties and autoflammability of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI can be extrapolated from the data obtained with SANITERPEN INSECTICIDE DK.

The detailed compositions of both formulations are presented in the confidential part.

Deltamethrin's content in tested product is:

- 0.173 % w/w in the Batch 27152-B1/95002 of SANITERPEN INSECTICIDE DK
- 0.020 % w/w in the Batch 27214-B1/95001 of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI (Effet choc)

The biocidal product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is a ready to use formulation contain in an individual PET hand-operated trigger sprayer and with a 0.20 % w/w of deltamethrin.

Properties	Method	Purity/ Specification	Results	Reference	Acceptable Yes/no
B3 – Physical, chem	nical and technica	l properties			
B3.1 Appearance					
B3.1.1 - Physical state and nature B3.1.2 - Colour B3.1.3 - OdourVisual examination Organoleptic determination	examination Organoleptic	Batch 27214- B1/95001	Homogeneous white and opaque liquid	B3.1 – C. Da Costa, B. Demangel 2010 Report No.09- 901011-006 GLP	Acceptable
		Batch 27214- B1/95001	Homogeneous white and opaque liquid	B3.2 – Demangel B. 2013a Report No.09- 901011-007 GLP	Acceptable
		Batch 12157001200301	white opaque liquid with a strong terpenic odour	B3.3 – Demangel B. 2013b Study No.13-901011- 015 No GLP	Acceptable
B3.2 Acidity/alkalini					
рН	CIPAC MT 75.3	Batch 27214- B1/95001	Pure test item: After 1 min pH value 4.07 at 19.6°C After 10 min pH value 4.08 at 19.4°C	B3.1 – C. Da Costa, B. Demangel 2010 Report No.09- 901011-006 GLP	Acceptable
Acidity	CIPAC MT 191	Batch 27214- B1/95001	<u>After 2 years at $20 \pm 2^{\circ}C$</u> : The mean value of a 10% w/v aqueous dilution of the test item acidity was 0.018 ± 0.001% w/w.	B3.2 – Demangel B. 2013a Report No.09- 901011-007 GLP	Acceptable
B3.3 Relative densit	ty and bulk, tap de	ensity		·	
Relative density	EC Method A3 OECD No. 109 method	Batch 27214- B1/95001	Pycnometric method: $D^{21.9^{\circ}C}_{4.0^{\circ}C} = 0.994 \pm 0.001$	B3.4 – C. Da Costa, B. 2009a Report No.09- 901011-005 GLP	Acceptable
B3.4 Storage stabili		nelf-life			
B3.4.1 Storage stab	ility tests				

Table 1: Physico-chemical properties of the biocidal product

Properties	Method	Purity/ Specification	Results				Reference	Acceptable Yes/no
B3 – Physical, cher	nical and technica	I properties						
B3.1 Appearance			1			.		
B3.4.1.1 – Accelerated storage study (2 weeks at 54°C)	CIPAC MT 41	Batch 27214- B1/95001		Initial	After storage 14 days at 54 ±2 °C in		B3.1 – C. Da Costa, B. Demangel 2010 Report No.09- 901011-006	The product was considered stable
	HPLC/UV method for deltamethrin		Annorrange	Homogonoo	PET spray and in glass flask		GLP	after 14 days at 54°C in glass flask and in the spray commercial
	opaque The asp to be s no sign	opaque liquid The aspect w to be stable	as considered after storage, at change of			The HPLC-UV method, used for the determination		
			A.s. content deltamethrin	0.020% w/v	0.020 % w/v			of deltamethrin content was
			pH (pure test item) after 1 min after 10 min	4.07 at 19.6°C 4.08 at 19.4°C	19.8°C			validated in this report (part 2.3.2.3).
			Dilution stability	at 20°C	IPAC water D			
			At T ₀	Homogene ous with a white foam (2 mL)	Homogeneo us			
			At T _{18h}	Homogene ous	Homogeneo us			

Properties	Method	Purity/ Specification	Results				Reference	Acceptable Yes/no
B3 – Physical, che	nical and technica							100/110
B3.1 Appearance								
B3.4.1.2 – Ambient shelf life study	GIFAP No. 17 Visual examination CIPAC MT 41 HPLC/UV	Batch 27214- B1/95001		Initial	After storage 2 years at 20 ±2 °C in PET spray		B3.2 – Demangel B. 2013a Report No.09- 901011-007 GLP	Acceptable The product was considered stable after 2 years at
	method for deltamethrin CIPAC MT 75.3 CIPAC MT 191		Appearance	opaque liquid The aspect w to be stable	s white and as considered after storage, at change of			20°C in the spray commercial packaging in PET. The deltamethrin content was
			A.s. content deltamethrin	0.020% w/v	0.017 % w/v (-15%)			decreased more than 10% (15%), but at this
			pH (pure test item) after 1 min after 10 min	4.07 at 19.6°C 4.08 at 19.4°C	3.89 at 20.2°C 3.86 at 19.8°C			substance active content the limit of 10% degradation is not relevant.
			Acidity (10 % aqueous dilution) Dilution	- 5 % v/v in C	0.018 ± 0.001% w/w			The HPLC-UV method, used for the determination of deltamethrin
			stability	at 20°C				content was
			At T ₀	Homogene ous with a white foam (2 mL)	Homogeneo us			validated in this report (part 2.3.2.3).
			At T _{18h}	Homogene	Homogeneo			
B3.4.1.3 – Low	CIPAC MT 39.3	Batch 27214-		ous <u>e test:</u> the tes	us t item was a h	nomogeneous white and		Acceptable
temperatures stability test (liquids)		B1/95001	and turbid liquid w	ith a white crea	amy phase (abo	was an offwhite opaque but 2 mL). vhite opaque and turbid	901011-005	The product was considered stable after 7 days at 0°C:

Properties	Method	Purity/ Specification	Results	Reference	Acceptable Yes/no
B3 – Physical, cher	nical and technica	I properties			
B3.1 Appearance	1	1			
			liquid with a white creamy phase (about 3 mL) (with the same black line). <u>After inverting the cones</u> : homogeneous offwhite and turbid liquid.		Nevertheless, the measure is mentioned on the label: "Before use, well stir the product."
	ontent of the active	e substance and teci	hnical characteristics of the biocidal product	[
B3.4.2.1 – Light	-	-	No required as the biocidal product is packaged in white PET bottle. The test item is considered to be stable after 14 days at $54 \pm 2^{\circ}C$	-	Acceptable
B3.4.2.2 –	-	-	(please refer to 3.4.1.1) and after 2 years at 20 \pm 2°C (please refer to	-	Acceptable
Temperature and			3.4.1.2).		
humidity			With its hand-operated trigger sprayer, the bottle is leak-tight.		
B3.4.2.3 –	-	-	Experience on the product has proved that no reactivity is expected	-	Acceptable
Reactivity towards			towards container material. The packaging material was considered to		•
container material			be stable after 2 years of the storage procedure at $20 \pm 2^{\circ}C$ (please refer to 3.4.1.2).		
B3.5 Technical cha	racteristics of the	biocidal product			
B3.5.1 – Wettability	-	-	Not required as the product is a ready-to-use liquid.	-	Acceptable
B3.5.2 – Suspensibility, spontaneity and dispersion stability	-	-	Not required as the product is a ready-to-use liquid.	-	Acceptable
B3.5.3 – Wet sieve analysis and dry sieve test	-	-	Not required as the product is a ready-to-use liquid.	-	Acceptable
B3.5.4 – Emulsifiability, re- emulsifiability and emulsion stability	-	-	Not required as the product is a ready-to-use liquid.	-	Acceptable
B3.5.5 – Disintegration tima	-	-	Not required as the product is a ready-to-use liquid.	-	Acceptable
B3.5.6 – Particle size distribution,	-	-	Not required as the product is a ready-to-use liquid.	-	Acceptable

Properties	Method	Purity/ Specification	Results	Reference	Acceptable Yes/no
B3 – Physical, cher	nical and technica	l properties			
B3.1 Appearance	•				
content of dust/ fines attrition, friability					
B3.5.7 – Persistent foaming	CIPAC MT 47.2	Batch 27214- B1/95001	In pure test item at 20°C: <u>After 10 s</u> : 7 mL <u>After 1 min</u> : 6 mL <u>After 3 min</u> : 5 mL After 12 min: 5 mL	B3.4 – C. Da Costa, B. 2009a Report No.09- 901011-005 GLP	Acceptable
B3.5.8 – Flowability/ Pourability/ Dustability	-	-	Not required as the product is a ready-to-use liquid.	-	Acceptable
B3.5.9 – Burning rate – smoke generators	-	-	Not applicable. The product is a ready-to-use liquid and is not intended to be applied as a smoke.	-	Acceptable
B3.5.10 – Burning completeness – smoke generators	-	-	Not applicable. The product is a ready-to-use liquid and is not intended to be applied as a smoke.	-	Acceptable
B3.5.11 – Composition of smoke – smoke generator		-	Not applicable. The product is a ready-to-use liquid and is not intended to be applied as a smoke.	-	Acceptable
B3.5.12 –Spraying pattern - aerosols	-	-	Not applicable. The product is a ready-to-use liquid and is not intended to be applied by spray with propellant gas under pressure.	-	Acceptable
B3.5.13 – Other technical characteristics		Batch 27214- B1/95001	trigger sprayer of the 750 mL PET bottle Dv (0%) = 5 μm Dv (0.4%) = 10 μm Dv (10%) = 52.42 μm Dv (90%) = 221.8 μm.	B3.5 – Buton N. 2010 Report 12/07/2010 No GLP B3.6 – Demangel B. 2013c No GLP	Acceptable
		ility with other produ	ucts including other biocidal products with which its use is to be author	orised	
B3.6.1 – Physical compatibility	-	-	Not applicable. The product is a ready-for-use liquid and is not intended to be added to any other product.		Acceptable

Properties	Method	Purity/ Specification	Results	Reference	Acceptable Yes/no
B3 – Physical, chen	nical and technica	I properties			
B3.1 Appearance					
B3.6.1 –Chemical compatibility	-	-	Not applicable. The product is a ready-for-use liquid and is not intended to be added to any other product.		Acceptable
B3.7 Degree of diss	olution and dilution	on stability			
Dilution stability	CIPAC MT 41	Batch 27214- B1/95001	In 5 % v/v in CIPAC water D at 20°C: <u>At T0</u> : Homogeneous white and opaque liquid with a white foam (2 mL) <u>After 18h</u> : Homogeneous white and opaque liquid	B3.1 – C. Da Costa, B. Demangel 2010 Report No.09- 901011-006 GLP	Acceptable
B3.8 Surface tensio					
Surface tension	EC Method A5 OECD No. 115	Batch 27214- B1/95001	The mean surface tension at 20.1 °C of the test item at 10% v/v aqueous dilution was: 32.5 ± 0.1 mN/m (corrected value). The test item was considered as surface-active in our experimental conditions.	B. 2009a	Acceptable The product is considered as surface-active.
B3.9 Viscosity	1	-	1	r	
Viscosity B4 – Physical hazar	OECD No. 114	Batch 27214- B1/95001	At 20°C (shear rate 48.5 s ⁻¹): 1.85 mPa.s (increasing rotation speed) At 20°C (shear rate 183.5 s ⁻¹): 1.92 mPa.s (increasing rotation speed) 1.92 mPa.s (decreasing rotation speed) At 40°C (shear rate 68 s ⁻¹): 1.15 mPa.s (increasing rotation speed) 1.16 mPa.s (decreasing rotation speed) At 40°C (shear rate 111 s ⁻¹): 1.17 mPa.s (increasing rotation speed) 1.18 mPa.s (decreasing rotation speed) 1.18 mPa.s (decreasing rotation speed) 1.18 mPa.s (decreasing rotation speed) Taking into account the results obtained at 20.0 and 40.0 °C, the test item was considered to have newtonian properties in our experimental conditions.	B3.4 – C. Da Costa, B. 2009a Report No.09- 901011-005 GLP	Acceptable The product is considered as a newtonian liquid.
					A
B4.1 – Explosives	Bridging DETERMINATIO N	Batch 27152- B1/95002	The comparison between SANITERPEN INSECTICIDE DK and SANITERPEN INSECTICIDE DK PRET A L'EMPLOI demonstrated that the explosive properties of SANITERPEN INSECTICIDE DK PRET A	2009a	Acceptable The product is not expected to have

Properties	Method	Purity/ Specification	Results	Reference	Acceptable Yes/no
B3 – Physical, chen	nical and technica	l properties			
B3.1 Appearance	[1			
			L'EMPLOI can be extrapolated from the data obtained with SANITERPEN INSECTICIDE DK. According to EC A.14 method, SANITERPEN INSECTICIDE DK is not explosive.	901011-001 GLP	explosive properties.
B4.2 – Flammable gases	-	-	Not required as the product is a liquid.	-	Acceptable
B4.3 – Flammable aerosols	-	-	Not required as the product is a liquid.	-	Acceptable
B4.4 – Oxidising gases	-	-	Not required as the product is a liquid.	-	Acceptable
B4.5 – Gases under pressure	-	-	Not required as the product is a liquid.	-	Acceptable
B4.6 – Flammable liquids	EC Method A9	Batch 27214- B1/95001	Automatic Setaflash closed tester: No flash point was observed up to 110.0 °C (corrected value).	B3.4 – C. Da Costa, B. 2009a Report No.09- 901011-005 GLP	Acceptable The product is not highly flammable.
B4.7 – Flammable solids	-	-	Not required as the product is a liquid.	-	Acceptable
B4.8 – Self- reactive substances and mixtures	-	-	Not required as the product is not explosive and not flammable according to EC method A.9.	-	Acceptable
B4.9 – Pyrophoric liquids	-	-	Not required as experience in manufacture and handling shows that the product does not ignite spontaneously on coming into contact with air at normal temperature.	-	Acceptable
B4.10 – Pyrophoric solids	-	-	Not required as the product is a liquid.	-	Acceptable
B4.11 – Self heating substances and mixtures	-	-	Not required as the product is a liquid.	-	Acceptable
B4.12 – Substances and	-	-	Not required as experience in handling and use shows that the product does not react with water.	-	Acceptable

Properties	Method	Purity/ Specification	Results	Reference	Acceptable Yes/no
B3 – Physical, chen	nical and technica	I properties			
B3.1 Appearance					
mixtures which in					
contact with water emit flammable					
B4.13 – Oxidising			The comparison between SANITERPEN INSECTICIDE DK and		
liquids	DETERMINATIO N	B1/95002	SANITERPEN INSECTICIDE DK PRET A L'EMPLOI demonstrated that the oxidising properties of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI can be extrapolated from the data obtained with SANITERPEN INSECTICIDE DK. According to EC A.21 method, SANITERPEN INSECTICIDE DK is not oxidising.	Study No. 09-	The formulation is not expected to have oxidising properties.
B4.14 – Oxidising solids		-	Not required as the product is a liquid.	-	Acceptable
B4.15 – Organic peroxides	-	-	Not required as the product not contains organic peroxide	-	Acceptable
B4.16 – Corrosive to metals	-	-	Not required as the product has not a low or high pH value ($pH = 4.1$) and does not contain halogen.	-	Acceptable
B4.17 Additionnal p	hysical indication	s of hazard			
B4.17.1 – Auto- ignition			The comparison between SANITERPEN INSECTICIDE DK and SANITERPEN INSECTICIDE DK PRET A L'EMPLOI demonstrated that		Acceptable The product is not
temperatures of	-	D1/90002	auto-flammability of SANITERPEN INSECTICIDE DK PRET A		expected to have
products (liquids					self-ignition
and gases)			SANITERPEN INSECTICIDE DK.	GLP	properties at
June guess)			According to EC A.15 method, SANITERPEN INSECTICIDE DK is not		ambient
			auto-flammable (mean self-ignition temperature: $265 \pm 4^{\circ}$ C).		temperature.
B4.17.2 – Relative self-ignition temperature for solids	-	-	Not required as the product is a liquid.	-	Acceptable
B4.17.3 – Dust explosion hzard	-	-	Not required as the product is a liquid.	-	Acceptable

Conclusion:

The biocidal product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is a homogeneous white and opaque liquid with a strong terpenic odour. The product has not explosive properties, nor oxidising properties. It is not highly flammable (flash point is > 110°C) and not auto-flammable at ambient temperature. The pH of the product in aqueous dilution (pure test item) is about 4.1 at 20°C and the density of the product is 0.994. The product is considered as surface-active (mean surface tension at 20.1°C of the pure test item is 32.5 ± 0.1 mN/m) and has non-Newtonian properties at 20.0°C and 40.0°C.

After the accelerated storage procedure (14 days at 54 \pm 2°C), no significant change of the product was observed, regarding the deltamethrin content, the aspect of the product and the pH. SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is considered stable after the accelerated storage during 14 days at 54 \pm 2°C in the individual PET hand-operated trigger sprayer.

After the long term storage procedure (2 years at ambient temperature), no significant change of the product was observed, regarding the deltamethrin content, the appearance of the product and its commercial packaging material (opaque white plastic flask of 750 mL with a green spray gun) and the pH. SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is considered as stable after the long term storage during 2 years at 20 \pm 2°C. Accordingly, defined shelf-life is 2 years.

Significant change was observed in the deltamethrin content (-15% deviation from t = 0 value) after 2 years of the storage procedure at $20 \pm 2^{\circ}$ C, nevertheless at this substance active content (0.02 % w/w) the limit of 10% degradation is not relevant.

After storage of the product for 7 days at $0 \pm 2^{\circ}$ C, the test item was an off-white opaque and turbid liquid with a white creamy phase (about 2 mL). Due to this result, an additional statement is mentioned on the label: "Before use, well shake the product".

The mean volume of foam produced after several inversions of the pure test item at $20 \pm 2^{\circ}$ C was 6 mL after 1 min of standing.

The particle size distribution has been determined with the hand-operated trigger sprayer of the 750 mL PET bottle (Dv (0%) = 5 µm, Dv (0.4%) = 10 µm, Dv (10%) = 52.42 µm and Dv (90%) = 221.8 µm).

2.3.2.3 Analytical method for determining the active substance and relevant component in the biocidal product

An analytical method for the determination of the active substance deltamethrine in the formulation SANITERPEN INSECTICIDE DK PRET A L'EMPLOI has been developed. The following analytical method for the determination of the active substance in the formulation performed on SANITERPEN INSECTICIDE DK PRET A L'EMPLOIhas not previously been reviewed and is provided in support of this assessment.

Report:	B5.1 – RICAU H., 2010a
Title:	Validation of the analytical method for the determination of deltamethrin in SANITERPEN INSECTICIDE EFFET CHOC, in compliance with SANCO/3030/99 rev.4 EU
Document No:	Defitraces, Report No. 09-901011-008
GLP	Yes
Report:	B5.2 – RICAU H., 2010b
Title:	Validation of the analytical method for the determination of deltamethrin in SANITERPEN INSECTICIDE DK
Document No:	Defitraces, Report No. 09-901011-004
GLP	

Principle of the method

A quantity of about 1.0 g (to the nearest 0.1 mg) of the test item was weighed into a 100-mL volumetric flask, and the volume was made with acetonitrile. Deltamethrin was analyzed by liquid chromatography using an UV detector (225nm).

Specificity for SANITERPEN INSECTICIDE DK (RICAU H., 2010b)

To define the specificity of the analytical method, the following solutions were analyzed: solvent, blank formulation, reference item and test item SANITERPEN INSECTICIDE DK.

In the reference item and in the test item, the peak at the retention time around 21 min represents deltamethrin. Three impurities are present in the blank formulation but their areas represent less than 3% of active substance area peak, then the specificity was therefore defined for deltamethrin.

Representative chromatograms of standard solution, sample solution of formulation and formulation blank were provided.

Specificity for SANITERPEN INSECTICIDE DK PRET A L'EMPLOI (RICAU H., 2010a)

To define the specificity of the analytical method, the following solutions were analyzed: solvent, blank formulation, reference item and test item SANITERPEN INSECTICIDE DK PRET A L'EMPLOI.

In the blank formulation and in the solvent, no peak appears at the retention time near the analyzed substance. In the reference item and in the test item, the peak at the retention time around 19 min represents deltamethrin.

Representative chromatograms of standard solution, sample solution of formulation and formulation blank were provided.

Linearity (RICAU H., 2010b)

For the calibration of deltamethrin, a calibration curve based on one injection per dilution was generated. The results were used to calculate the calibration curves to verify the linearity of detector response.

The analytical system gave a linear response (n=5) between 0.01 g/L and 0.030 g/L of Deltamethrin. The linear correlation coefficient for the calibration range was found to be > 0.99.

equation of the calibration line :	$Y = 4.90^{*}10^{8} X + 2.78^{*}10^{5}$
correlation coefficient :	$r^2 = 0.9984$

Accuracy for SANITERPEN INSECTICIDE DK (RICAU H., 2010b)

For accuracy 100%

A quantity of about 99.8 mg (to the nearest 0.1 mg) of blank formulation was weighed into a 10-mL volumetric flask a volume of 2-mL of the REF02 Delta was added and the volume was made up with acetonitrile (EXACT 1 100%). An identical solution was prepared with REF03 Delta (EXACT 2 100%).

For accuracy 50%

A quantity of about 49.9 mg (to the nearest 0.1 mg) of blank formulation was weighed into a 10-mL volumetric flask a volume of 1-mL of the REF02 Delta was added and the volume was made up with acetonitrile (EXACT 1 50%). An identical solution was prepared with REF03 Delta (EXACT 2 50%).

	Fortification Level [mg/L]	Number of Analyses	Mean Recovery [%]
Accuracy	20.6	2	96.9
100%	20.8	2	96.2
	10.3	2	101.8
Accuracy 50%	10.4	2	95.1

Accuracy for SANITERPEN INSECTICIDE DK EXTRA (RICAU H., 2010a)

For accuracy 100%

The recovery rate of deltamethrin in two reconstituted samples of the test item at 20.6 and 21.2 mg/L are given in the following table.

For accuracy 50%

The recovery rate of deltamethrin in two reconstituted samples of the test item at 10.3 and 10.6 mg/L are given in the following table.

	Fortification Level [mg/L]	Number of Analyses	Mean Recovery [%]
Accuracy	20.6	2	102.0
100%	21.2 2	2	99.0
Accuracy 50%	10.3	2	99.9
	10.6	2	99.2

Repeatability (RICAU H., 2010b)

The precision was determined by analyzing twice five specimen samplings. The content of deltamethrin for each analysis was calculated with the average value of the response factor of the two calibration solutions. Then, the average value of the content, the standard deviation and the Relative Standard Deviation (R.S.D.) were calculated.

The acceptability of the precision was based on the following modified Horwitz equation.

Precision results for Pyrimethanil

Mean Fortification Level	Number of Analyses	RSD	RSD Horwitz
[% w/w]		[%]	[%]
0.193	10	2.89	3.44

Conclusion

Specificity, linearity, precision and recovery were provided and found acceptable for Deltamethrin.

Analytical methods using HPLC/UV (**RICAU H., 2010a**, report No. 09-901011-008 and **RICAU H., 2010b**, report No. 09-901011-004) for the determination of deltamethrin in the formulation (SANITERPEN INSECTICIDE DK PRET A L'EMPLOI) as manufactured has been performed and validated in accordance to guidance of Regulation (EU) No 528/2012.

2.3.2.4 Analytical methods for determining relevant components and/or residues in different matrices

Analytical methods for deltamethrin residues in soil, air, water (including drinking water) and sediment, animal and human body fluids and tissues and deltamethrin residues in food/feed of plant and animal origin are available in Assessment Report Deltamethrin Product-type 18 (insecticides), May 2011.

A Letter of Access from Bayer has been provided.

2.4 Risk assessment for Physico-chemical properties

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is a ready-to-use product containing deltamethrin (0.02 % (w/w)) spray application. It is not highly flammable, not auto-flammable, not explosive and does not have oxidizing properties.

The product is stable for 14 days at 54°C and 2 years at ambient temperature.

The product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is compatible with PET packagings

Risk mitigation measures linked to risk assessment for physico-chemical

- Shake before use.

Disposal considerations

None.

Required information linked to risk assessment for physico-chemical properties

None.

2.5 Effectiveness against target organisms

2.5.1 Function

MG 03: Pest Control

Product Type 18: Insecticides, acaricides and products to control other arthropods.

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is presented as a liquid ready to use supplied in a 750 ml PET bottle with a hand sprayer.

The formulation contains 0.02 % w/w of the insecticidal active substance, deltamethrin.

The biocidal product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is used by professional, for the control of poultry red mites, mosquitoes, flies and fleas in small and pet breeding, in animal boarding and pet shop.

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

According to the uses claimed by the applicant, the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is intended to be used to control poultry red mites (eggs, larvae and adults), mosquitoes (adults), flies (adults) and fleas (adults).

The specific target organisms to be controlled are:

- Poultry red mites: Dermanyssus gallinae
- Mosquitoes: Culex spp. and Aedes spp.
- House fly: *Musca domestica,*
- And Fleas: Pulicidae spp

The solution must be sprayed on the floor, walls, ceiling, and around windows and doors to control flies and mosquitoes, or applied on the floor and walls to control fleas and red mites.

The application rates recommended by the applicant are the following:

The recommended application rate (for porous and non-porous surfaces) is 100 % v/v, i.e 50 ml of the product to treat 1 m².

The product is claimed as efficient during 3 months.

Animals, textiles, animal food and sources of water shall be cleared off the treated areas before using the product. They can be reintroduced only when the treated surfaces are dried. A default time of 24 hours shall be considered.

2.5.3 Effects on target organisms and efficacy

The studies submitted to demonstrate the efficacy of the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI according to the uses and doses claimed, are described below. These studies were carried out with the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI (0.02 % w/w deltamethrin), with a similar product SANITERPEN INSECTICIDE EFFET CHOC PAE (0.02 % w/w deltamethrin) and the product SANITERPEN INSECTICIDE DK EXTRA (0.2 % w/w deltamethrin), diluted at 10 % v/v

The product SANITERPEN INSECTICIDE DK PAE contains terpineol as solvent. The component terpineol is identified in annex 1 of Commission Regulation 1451/2007/CE of 4th December 2007, and is also known in the literature as having potential insecticide effects. In order to demonstrate that Terpineol doesn't have any insecticidal properties in this formulation, a laboratory study has been carried out with SANITERPEN INSECTICIDE DK PAE.

The laboratory study n°1948c/0615, 2015/11 was conducted with the products SANITERPEN INSECTICIDE DK PAE (0.02 % w/w deltamethrin) and SANITERPEN INSECTICIDE DK PAE without deltamethrin, on *Musca domestica* (house fly, adults), *Stomoxys calcitrans* (Stable fly, adults), *Phlebotomus duboscqi* (sandfly, adults), *Culex pipiens* and *Aedes aegypti* (mosquitoes, adults), *Ctenocephalides felis* (cat flea, adults and larvae) and *Dermanyssus gallinae* (poultry red mite, adults and larvae) according to a methodology adapted from the standard ASTM E 654-96 (Reapproved 2009): "Standard test method for effectiveness of aerosol and pressurized spray Insecticides against cockroaches".

The products were sprayed directly at the claimed application rate of 50 mL of product per m² on the arthropods. After the spraying, different parameters were measured, including the KD1 hour, KT100 and mortality after 1 day and 3 days. Four replicates were made for each species.

The untreated controls demonstrated the validity of the test, with less than 5 % of mortality. The product SANITERPEN INSECTICIDE DK PAE with active substance showed total efficacy against all tested target organisms. The product without active substance gave no significant efficacy (<10% mortality within 72 hours).

It can be concluded that the product without active substance presents no efficacy on its own at the recommended application rates and that terpineol is a co-formulant that has no insecticide properties in this product.

1. Laboratory study n°1558c/1112R, 2013/04 conducted with the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI, (0.02 % w/w deltamethrin) on *Musca domestica* (house fly, eggs and larvae), *Culex pipiens* (mosquito, adults), *Ctenocephalides felis* (cat flea, adults and

larvae) and *Dermanyssus gallinae* (poultry red mite, adults and larvae) according to CEB N°135 method.

The product was sprayed at the dose of 100 % v/v, at the rate of 50 mL per m² on 4 representative surfaces (concrete, wood, plaster and ceramic tiles) and the arthropods were placed in contact with these surfaces for an exposure time of 4 hours. The same test was done with treated surfaces after 1, 2 and 3 months of storage. The mortality of the arthropods was recorded during the exposure period and during post monitoring. Four replicates were made for each test condition (surface*treatment*storage*arthropod).

The untreated controls demonstrated the validity of the test, with less than 5 % of mortality. The product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI showed total efficacy against flies (eggs and larvae), fleas (adults and larvae), mosquitoes (adults) and poultry red mites (adults and larvae), on both porous and non-porous surfaces. This efficacy lasted for at least 3 months after application.

2. Laboratory study n°1296b/1008R, 2011/02 conducted with the product SANITERPEN INSECTICIDE EFFET CHOC PAE, (0.02 % w/w deltamethrin) on *Musca domestica* (house fly, adults) following a method similar to CEB N°135 method.

The product was sprayed at the dose of 100 % v/v, at the rate of 50 ml/m² on the surface (ceramic tiles) and the arthropods were placed in contact with these surfaces during 2 hours. The mortality is recorded every 10 minutes. The mortality is controlled again 24 and 48 hours after the application. The same procedure is applied with aged product of 3, 6, 12, 18, 24, 30 and 36 months.

Four replicates were made for each test condition (surface*treatment*storage*arthropod).

The untreated controls demonstrated the validity of the test, with an absence of mortality. The product SANITERPEN INSECTICIDE EFFET CHOC PAE (considered as similar to the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI) showed a knock-down effect (100 %) from 35 minutes to 3 hours of contact depending on the age of the product and, 100 % of mortality in 24 hours against flies on a non-porous surface even after storage of three years in his packaging.

3. Laboratory study n°1826a/0914R, 2014/12 conducted with the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI, (0.02 % w/w deltamethrin) on *Musca domestica* (house fly) and Aedes aegypti (mosquito) according to CEB N°135 method.

The product was sprayed at the dose of 100 % v/v, at the rate of 50 mL per m² on 2 representative surfaces (concrete and ceramic tiles) and the arthropods were placed in contact with these surfaces for an exposure time of 1 hour. The same test was done with treated surfaces after 1, 2 and 3 months of storage. Four replicates were made for each test condition (surface*treatment*storage*arthropod).

The untreated controls demonstrated the validity of the test, with less than 4 % of mortality. The product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI showed knock-down effect (100 %) within 5 minutes and a total efficacy against flies (adults), and mosquitoes (adults), on both porous and non-porous surfaces. This efficacy lasted for at least 3 months after application.

4. Semi-field tests n°1826b/0914, 2014/11 conducted with the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI, (0.02 % w/w deltamethrin) on *Aedes aegypti*, *Culex pipiens* (mosquitoes) and *Ctenocephalides felis* (fleas), according to BSI 4172 method.

The product was sprayed using the provided spray trigger container at the rate of 50 ml/m² in a test chamber (30 m³ with 12 m² floor) and the treated surface (ceramic tiles) is half of the test chamber (6 m²). To simulate what happens in premises, some panels of polystyrene blocks and cardboards are set into the test chamber to

be harbourages and a water+food sources (these ones are not treated). The insects are able to reach water and food sources without being in contact with the insecticide and they have the choice not to be in contact with the product.

Three replicates were made for each species.

The untreated controls demonstrated the validity of the test, with less than 10 % of mortality for each species. The product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI showed 100 % of efficacy 7 days after the application.

Nevertheless it shall be noted than experimental conditions described in this simulated test are different from the real conditions observed in breeding premises (behaviour of insect's populations more complex, nature of building materials, variability of parameters such as temperature, humidity and light are not taken into account). Moreover, residual activity of the product until 3 months was not assessed.

It shall be also noted that these semi-field tests were performed only with adult's insects whereas larvae stage were also claimed for fleas.

5. Field test n° 1527/07/12, 2012/11 conducted in breeding premises with the product SANITERPEN INSECTICIDE DK EXTRA, on *Musca domestica* (house fly), according to CEB n°107 method

The product used in this study is SANITERPEN INSECTICIDE DK. The product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is a dilution at 10 % v/v of the product SANITERPEN INSECTICIDE DK and then is considered as equivalent.

After confirmation and evaluation of the initial infestation level with sticky traps, the product was diluted in water (10 % v/v) and applied by spraying to vertical surfaces, at the application rate of 50 ml of diluted product per m². The reduction of the insect's population was assessed after 7, 14, 30, 60 and 90 days, by counting flies trapped on sticky traps let for 24 hours in the premises.

A reference product is included and three premises are treated with each product.

The population decreased in the treated breeding premises until 90 % from the first week and 95 % from two weeks after treatment and still of 97.9 % after 3 months. The test product gave results similar to the reference product.

In conclusion, in accordance with the tests submitted and the requirements of the TNsG on PT18, the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is efficient against domestic flies (*Musca domestica*, adults) but the efficacy against poultry red mite (*Dermanyssus gallinae*) is not proved as no field test has been provided for this target organism.

Semi-field tests are submitted for mosquitoes and fleas but experimental conditions are a bit far from the field and therefore not sufficiently reliable. So, in the absence of sufficient supporting data on mosquitoes (genus *Culex* and *Aedes*) and fleas (*Pulicidae*), suitable information (as semi-field or field tests) demonstrating the efficacy of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI against these target organisms, will need to be provided in support of the authorisation, within one year

Conclusion

All efficacy studies are presented in Annex 9.

Target Organismes	Rates and uses acceptable	Method of application	Time delay of the biocidal product
House flies, adults (<i>Musca domestica</i>)	100 % v/v		
Mosquitoes, adults genus Culex, genus Aedes	The product is ready to use and supplied in a 750 ml PET bottle with a hand sprayer Application rate : 50 ml/m ²	Porous and non porous surfaces	after a few hours
Fleas (<i>Pulicidae</i>), adults			

Based on these efficacy data, the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI (0.02 % w/w deltamethrin), formulated as ready to use liquid, at a rate of 100 % v/v, showed an efficacy against house flies (*Musca domestica*, adults), mosquitoes (*genus Culex and Aedes*, adults) and fleas (*Pulicidae*, adults) over a 3 months period.

In laboratory tests, the effect began a few hours after application.

2.5.4 Mode of action including time delay

Deltamethrin is a pyrethroid insecticide which acts on harmful organisms by contact and ingestion. It expresses a strong knock-down effect.

Pyrethroids impair ion transport through the membrane of nerve axons, causing muscular paralysis in the insect; death seems to follow a nervous system impairment that occurs a few minutes to several hours after pesticide absorption. The primary site of activity of deltamethrin is the voltage sensitive sodium channel in nerve membrane. Deltamethrin prolongs the opening of the sodium channels (i.e. the channels directly responsible for generating nerve action potentials) leading to neuronal hyper excitability.

The effect begins around some hours after contact of the product in the laboratory trials submitted by the applicant.

2.5.5 Occurrence of resistance – resistance management / unacceptable effect

Deltamethrin is a pyrethroid insecticide. Deltamethrin products are widely used for various applications: veterinary medicine, crop protection, indoors and outdoors biocide, and against numerous arthropods target organisms. Resistance to deltamethrin has already been reported in several insects, including important nuisance insects of breeding premises.

Resistant populations of house flies (*Musca domestica*) have been identified in the whole world (Asia, Europe, and America). Several mechanisms are involved in resistance to pyrethroïds in house fly. This includes detoxification of the active substances, knockdown resistance (also called kdr), correlated with decrease nerve sensitivity, and decrease in the rate of penetration of the products. These mechanisms can co-exist in a resistant strain⁵. Cross-resistance also exists in resistant strains, among pyrethroïds, but also other insecticide types (chlorpyrifos and imadacloprid).

A recent French study with two populations of stable fly (*Stomoxys calcitrans*) concluded that the population from the "conventional" farm, with cattle and stable walls frequently treated with deltamethrin, was resistant to the five tested pyrethroïds, including deltamethrin⁶.

Concerning *Culex pipiens* (or *C.quinquefasciatus*, very close species), deltamethrin resistant populations have been identified in many part of the world, including North and West Africa and Asia⁷. Resistant populations have not been identified in Europe yet.

⁵ Nannan L, Xin Y. Insecticide resistance and cross-resistance in the house fly (Diptera:Muscidae).J. Econ. Entomol. 93(4):1269-1275 (2000).

⁶ Salem A, Bouhsira E, Liénard E, Bousquet Melou A, Jacquiet P, Franc M. Susceptibility of two European strains of *Stomoxys calcitrans* (L.) to Cypermethrin, Deltamethrin, Fenvalerate, λ-cyalothrin, Permethrin and Phoxim. Intern J Appl Res Vet Med. Vol. 10, N°.3, 2012.

Populations of fleas (*Ctenocephalides felis*) resistant to deltamethrin have not been identified in Europe yet. Resistance to insecticide is difficult to identify in fleas, because of important intra-population variability⁸.

Concerning poultry red mites (*Dermanyssus gallinae*), populations resistant to permethrin, another pyrethroïds active substance, have been identified in France and other European countries. However, resistance to deltamethrin is not reported in the scientific literature⁹.

To ensure a satisfactory level of efficacy and avoid the development of resistance in susceptible insect populations, the following recommendations have to be implemented:

- Always read the label or leaflet before use and respect follow all the instructions provided.
- Take into account the life cycle and characteristics of target insects to adapt treatments. In particular, target the most susceptible stage of the pest, timing of applications and areas to be treated.
- Adopt integrated pest management methods such as the combination of chemical, physical control methods and other public health measures, taking into account local specificities (climatic conditions, target species, conditions of use, etc).
- Alternate products containing active substances with different mode of action, (to remove resistant individuals from the population).
- Establish a baseline and monitor levels of effectiveness on populations in key areas (at least one survey per year) in order to detect any significant changes in susceptibility to active substance. Information from resistance monitoring programs allows early detection of problems and gives information for correct decision making.
- The users should inform if the treatment is ineffective and report straightforward to the registration holder
- The authorization holder should report any observed resistance incidents to the Competent Authorities (CA) or other appointed bodies involved in resistance management.

2.5.6 Evaluation of the Label Claims

French competent authorities (FR CA) assessed that the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI, as a ready to use 750 ml PET bottle with a hand sprayer, applied at the application rate 50 ml /m² has shown a sufficient efficacy for the control of domestic flies (*Musca domestica*), mosquitoes (genus *Culex spp.* and *Aedes spp.*) and fleas (*Puclicidae*), up to 3 months.

The product is applied indoor on the porous and non-porous surfaces in empty small and pet breeding, animal boarding and pet shop.

As only efficacy has been demonstrated against adult's insects, treatment with the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI shall be completed with a product efficient on larvae stages.

2.5.7 Summary of efficacy assessment

The efficacy level of the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI (0.02 % w/w deltamethrin) is satisfactory for the uses proposed in Table 2 below.

Conditions of use linked to efficacy assessment

To ensure a satisfactory level of efficacy and avoid the development of resistance in susceptible insect populations, the following recommendations have to be implemented:

- Always read the label or leaflet before use and respect follow all the instructions provided.

⁷ Tahir HM, Butt A, Khan SY. Response of Culex quinquefasciatus to deltamethrin in Lahore district. Journal of Parasitology and Vector Biology Vol. 1 (3) pp. 019-024, October, 2009

⁸ Bossard RL, Hinkle NC, Rust MK. Review of insecticide resisitance in cat fleas (Siphonatera : Pulicidae). J. Med. Entomol. 35(4):415-422 (1998)

⁹ Marangi M, Cafiero MA, Capelli G, Camarda A, Sparagano OAE, Giangaspero A. Evaluation of the poultry red mite, *Dermanyssus gallinae* 'Acari:Dermanyssidae) susceptibility to some acaricides in field populations from Italy. Exp Appl Acarol (2009) 48:11-18.

- Take into account the life cycle and characteristics of target insects to adapt treatments. In particular, target the most susceptible stage of the pest, timing of applications and areas to be treated.
- Adopt integrated pest management methods such as the combination of chemical, physical control methods and other public health measures, taking into account local specificities (climatic conditions, target species, conditions of use, etc).
- Alternate products containing active substances with different mode of action, (to remove resistant individuals from the population).
- The users should inform if the treatment is ineffective and report straightforward to the registration holder.

Recommendations to be taken into account by the authorisation holder

- The authorization holder should report any observed resistance incidents to the Competent Authorities (CA) or other appointed bodies involved in resistance management.

Required information linked to efficacy assessment

- Suitable information (as semi-field or field tests) demonstrating the efficacy of SANITERPEN INSECTICIDE DK EXTRA against target organisms mosquitoes (Culex pipiens and Aedes aegypti) and fleas (Pulicidae), will need to be provided in support of the authorisation, within one year.
- Establish a baseline and monitor levels of effectiveness on populations in key areas (at least one survey per year) in order to detect any significant changes in susceptibility to active substance. Information from resistance monitoring programs allows early detection of problems and gives information for correct decision making.

2.6 Description of the intended use(s)

MG/PT	Field of uses envisaged	Likely concentrations at which product will be used
Main Group 03; Pest Control	Professional uses	
PT18: insecticides, acaricides and products to control other arthropods	Insecticide for use against house flies, mosquitoes and fleas infestation (adult stage) indoor in small and pet breeding, in animal boarding and pet shop.	100 % v/v The product is ready to use and supplied in a 750 ml PET bottle with a hand sprayer Application rate : 50 ml/m ²

Table 2: Summary of intended uses

2.7 Risk assessment for human health

2.7.1 Hazard potential

2.7.1.1 Toxicology of the active substance

The toxicology of the active substance was examined extensively according to standard requirements. The results of this toxicological assessment can be found in the CAR. The threshold limits and labelling regarding human health risks listed in Annex 4 "Toxicology and metabolism" must be taken into consideration.

The following corresponds to the summary of the effect assessment available in the assessment report of Deltamethrine.

Absorption, distribution, metabolism and excretion

The rate of oral absorption of deltamethrin was approximately 75%, this based on urinary and biliary excretion data in rats. Deltamethrin was rapidly absorbed when orally administered to rats (the majority of the radioactivity was eliminated within 24 hrs after dosing, 19-47% with the urine; 32-55% in faeces) and distributed to most tissues. Residues in tissues and carcass were low. The highest residues were found in fat. There was no indication of accumulation, although the residue of deltamethrin in adipose tissue eliminated with a half-life of >24 hrs.

Deltamethrin was rapidly excreted in both urine and faeces. 7 days postdose, 31% to 56% of the oral dose was excreted with the urine and 36% to 59% in faeces. No 14CO2 was formed according to data from the open literature. Deltamethrin was rapidly and extensively metabolised in rats. The main route of metabolism was via cleavage of the ester bond with or without hydroxylation at the 4' position of the alcohol moiety. The acid moiety and alcohol moiety were further transformed and excreted in urine in free forms and as conjugated metabolites. Unchanged deltamethrin was the major compound in faeces.

No studies were located regarding absorption rate following inhalation exposure to animals. Consequently a default absorption value by inhalation was considered in the risk assessment.

Dermal absorption

Dermal penetration studies have been conducted in vitro in rats with deltamethrin as an oil/water emulsion (EW) and as an emulsifiable concentrate (EC) in rat and human skin and in an in vivo study in rats. The results of these studies indicated that dermal absorption was somewhat lower for the EW 15 than for the EC 25. The Decis EC 25 formulation may be considered to be a worst case with regard to K-Othrine formulations. The main difference which is relevant to skin absorption is the solvent (water in K-Othrine SC formulations versus light aromatic solvent in Decis EC 25). The content of aromatic solvent is expected to enhance the degree of dermal absorption in comparison with K-Othrine formulations. For the solid formulations of deltamethrin a lower dermal absorption is expected since water and certain solvents favour.

Using data obtained in the dermal absorption studies on Decis EC 25 formulation, the dermal absorption of deltamethrin in man was estimated to 1.19% for the concentrate and 1.89% for the a.s. when diluted in the spray solution. The value of 2% (maximum dermal absorption) was used in the risk assessment.

Acute toxicity, irritation and corrosivity, sensitisation

Deltamethrin was considered of high acute toxicity by the oral and inhalation route (LD_{50} rat: 87 mg/kg bw; LC_{50} rat: 0.6 mg/L), while the acute dermal toxicity of deltamethrin was low (LD_{50} rat: >2000 mg/kg bw). Clinical signs of systemic toxicity, poor condition and neurotoxicity were observed in rats after oral and inhalation administration. Skin and eye irritation and pathological changes (enlarged inguinal and mandibular lymph nodes, and pulmonary congestion) were noted in addition after administration via the inhalation route. No clinical signs were noted in rats after dermal application.

The vehicle has a great influence on the LD_{50} . Sesame oil as vehicle shows less toxicity than polyethylene glycol. Aqueous suspensions are significantly less toxic than formulations in oils.

Deltamethrin was not irritating according to skin- and eye irritation studies in rabbits, and no sensitising potential was found in tests according to GPMT (Guinea Pig Maximisation Test) or Buehler.

Repeated dose toxicity (short-term toxicity)

The short-term oral toxicity of deltamethrin was investigated in rats (90-day studies) and dogs (90-day studies; one-year study). In both species, the nervous system was the main target organ. Reduced bodyweight gain was also noted in both species.

The lowest relevant NOAEL for short-term toxicity was 1 mg/kg bw/day obtained in the 90-day (gelatine capsules, vehicle: PEG 200) and 1-year oral (gelatine capsules, vehicle: none) toxicity studies in dogs based on clinical signs of neurotoxicity noted in both sexes at the dose level of \geq 2.5 mg/kg bw/day.

In addition, the repeated dose toxicity was investigated in rats after dermal exposure (21-day toxicity study in rats) where dermal irritation was noted, and inhalation exposure (14-day toxicity study in rats) where clinical

signs (irritative and neurotoxic) and reduced bodyweight gain were noted. Scratching was noted in all treated groups in the inhalation toxicity study. This effect was considered to be related to the irritant nature of deltamethrin but may also be due to the neurotoxic nature of the substance (an indirect consequence of parasthesia).

Genotoxicity

The genotoxic potential of deltamethrin was investigated in a battery of tests in vitro (assays for gene mutations, chromosomal aberrations and DNA effects). All tests were negative.

Based on the weight of evidence from this full in vitro package and the results of the carcinogenicity studies, it was concluded that deltamethrin is not mutagenic.

Chronic toxicity (long-term toxicity) and carcinogenicity

The long term toxicity of deltamethrin was studied in rats and mice. No evidence of carcinogenic potential of deltamethrin was found in the rat or the mouse. In both species the nervous system was the target organ. The liver was another target organ in the rat.

Lowest relevant NOAEL for long-term toxicity was 1 mg/kg bw/day obtained in the 2-year chronic toxicity/carcinogenicity (feeding) study in the rat based on liver effects (histopathological changes) noted at the dose level of 5 mg/kg bw/day and above. In addition clinical signs of neurotoxicity were noted at higher doses.

Reproductive toxicity

Reproductive toxicity of deltamethrin was investigated in a two-generation study in rats. Developmental toxicity was investigated in rats, mice and rabbits. The mouse study was considered acceptable but of restricted quality due to low number of pregnant animals used in each test groups.

No effect on mating performance or fertility was noted in the rat two-generation (feeding) study. Clinical signs (indicating neurotoxic effects), reduced body growth and histopathological changes (gastric erosions) were noted in adult rats. In offsprings reduced pup body weights, increased pup deaths (F1 generation) and reduced lactation index (F1 generation) were noted at maternal toxic doses.

No developmental toxicity was noted in rats or rabbits at maternal toxic doses. Increased incidence of supernumerary ribs was noted in the offspring of mouse at doses with maternal toxicity.

Lowest relevant developmental LOAEL was 3 mg/kg bw/day based on a statistically significant increase in the occurrence of supernumerary ribs noted in mice at \geq 3 mg/kg bw/day.

Neurotoxicity

The neurotoxicity of deltamethrin was investigated in standard toxicity studies with the rat (acute neurotoxicity study; subchronic neurotoxicity study; developmental neurotoxicity (DNT) study) and in experimental (non GLP) studies in rats and mice. No studies on acute delayed neurotoxicity were submitted (not required).

The NOAEL for acute neurotoxicity in adult CD-rats was 5 mg/kg bw, while the NOAEL for subchronic neurotoxicity in adult CD rats was 4 mg/kg bw/day. In both studies the NOAEL was based on signs of neurotoxicity noted at 15 mg/kg bw/day and above, and mortalities and reduced bodyweight gain noted at higher dose levels.

The NOAEL for developmental neurotoxicity in Wistar rats was 6.78 mg/kg bw/day based on reduced bodyweight gain, increased incidence of vocalizations with handling (males only) and delayed balanopreputial separation noted in offsprings at a dose with maternal toxicity (16.1 mg/kg bw/day).

The DNT study follows the OECD guideline no. 426 in that way that some exposure to the pups was demonstrated in the pilot study. However, the view of RMS is that there might be some uncertainty in the DNT study protocol in those cases where direct dosing of pups has not been considered and the exposure level in offspring is not clear. No blood analyses were taken and the offspring dose level might be very low. The effects noted in the pups of the high dose group (decreased body weight and body weight gain, delayed sexual maturation in males) are not sufficient evidence to support exposure to the pups during the brain growth spurt period since these effects in the offspring could be due to maternal toxicity or exposure in utero. Furthermore, there is a concern for the lack of data for the most sensitive strain. Comparing data from standard neurotoxicity studies the Wistar rat used in the DNT study seems to be a less sensitive strain with

regard to neurotoxicity of deltamethrin. There were no clinical signs of neurotoxicity reported for adult Wistar rats administered deltamethrin via the diet at doses up to 16.1 mg/kg bw/day (noted in the DNT study), whereas clinical signs of neurotoxicity were evident in the CD rat at a dose level of 14 mg/kg bw/day (noted in the 13-week neurotoxicity study). The choice of strain used in the deltamethrin DNT study might therefore be questioned.

Due to the uncertainties mentioned above the RMS originally proposed (draft CAR) to use an extra safety factor of 3 in the risk assessment of deltamethrin. The Technical Meeting I in 2010 reached an agreement that where uncertainties are perceived by the RMS of a pyrethroid on the DNT studies (especially negative studies), these uncertainties should be formally expressed in the CAR. The TM also agreed that the currently available evidence does not support the use of an extra assessment factor to cover for the perceived uncertainties on DNT in the dossier of deltamethrin.

During the Technical Meeting II in 2010, it was decided to use the document on survey of DNT studies for pyrethroids prepared by the Netherlands as basis for the assessment of this category of substances. The conclusions of this survey were:

Possible DNT effects induced by pyrethroids are covered by the AELs set on neurotoxicity in the acute neurotoxicity and medium-term studies since DNT effects from acceptable OECD TG 426 performed studies are taking place at higher LOAELs than other neurotoxicological effects.

The DNT effects are also covered by the AELs set for long-term exposure (based on neurotoxic or other critical endpoints).

As neurotoxic effects are critical effects after acute or medium-term exposure and the available data indicate that DNT effects are induced at higher LOAELs, it is unlikely that, in the absence of DNT studies, the potential DNT effects are not covered by AELs set on neurotoxic effects observed in acute and medium-term studies. It was concluded that additional DNT studies according to OECD TG 426, if such a study is not present, is not necessary.

The RMS respects the decision of TM although the view of RMS is still that there might be some uncertainty in the DNT study protocol and the most sensitive strain has not been used in the DNT study.

Medical data

Medical data from manufacturing, formulating and packaging plants indicate that transitory skin sensations were the most prevalent finding (paraesthesia, transient local burning, tingling, pickling sensations, itching, numbness of the facial skin – erythema in some cases). Cases of intoxications (mostly occupational due to inappropriate handling of products) have been reported. Two cases of occupational acute deltamethrin poisoning died of convulsions and another died of pulmonary oedema. No late sequeala of pyrethroid poisoning have been described in the scientific literature. There is no specific antidote for pyrethroids. Any treatment can only be symptomatic.

Other test(s) related to the exposure of humans The trans-deltamethrin isomer has been tested for oral acute toxicity and mutagenicity (Ames test). The results of these studies showed that the acute oral toxicity of the trans-deltamethrin does not exceed the acute oral toxicity of the parent compound cis deltamethrin and no

genotoxicity potential was found according to the Ames test.

In a study where food commodities (covered and uncovered) were exposed to an environment in which a deltamethrin based product was applied as a general surface treatment showed that the use of deltamethrin products will not contaminate food stuffs when spray is applied downwards. Spraying overhead or direct transfer of residues from treated spaces was not investigated in this study. However, no exposure of food stuffs is expected during and after crack and crevice treatment of food handling areas with the deltamethrin product when label instructions are followed.

Biocidal products

The acute toxicity of K-Othrine SC 26.25, SC 7.5 and DP 0.05 by oral, dermal and inhalation exposure is low. The acute toxicity of K-Othrine WG 250 by oral and dermal route is low, whereas the acute toxicity by

inhalation route is moderately; therefore K-Othrine WG 250 should be classified as "Harmful" and assigned the risk phrase R20 ("Harmful by inhalation").

The products are not irritating to skin or eyes, and are not sensitising to skin.

Tolerable exposure

The reference values, (acute/medium term and long term AELs) derived for deltamethrin were obtained from studies in dogs since the data submitted demonstrated that the dog was the most sensitive species to the toxicity of deltamethrin. In addition a safety factor of 100 was applied taking into account a factor for inter- and intraspecies differences of 100 (10×10).

Acceptable daily intake (ADI)

Setting of an ADI is not considered necessary since no exposure of foodstuffs should occur during and after treatment of food handling areas with deltamethrin when product label instructions are followed.

Acute reference dose (ARfD)

Setting of an ARfD is not considered necessary since no exposure of foodstuffs should occur when product label instructions are followed, and risk of contamination of drinking water is not considered.

Acceptable exposure levels (AELs)

AEL (acute): An AEL of 0.0075 mg/kg bw/day was derived based on the NOAEL (1 mg/kg bw/day) obtained in a 13-week dog study after taking an oral absorption of 75% and a safety factor of 100 into account. In the study neurotoxic effects occurred early after dosing.

AEL (medium-term): An AEL of 0.0075 mg/kg bw/day was derived based on the NOAEL (1 mg/kg bw/day) obtained in the 13-week and 1-year dog studies after taking an oral absorption of 75% and a safety factor of 100 into account.

AEL (long-term): An AEL of 0.0075 mg/kg bw/day was derived based on the NOAEL (1 mg/kg bw/day) obtained in the 1-year dog study after taking an oral absorption of 75% and a safety factor of 100 into account.

Maximum acceptable concentration in drinking water

According to Council Directive 98/83/EC relating to the quality of water intended for human consumption, the maximum admissible concentration for pesticides in drinking water is 0.1µg/l for substances considered separately.

2.7.1.2 Toxicology of the substance(s) of concern

The biocidal product contains no substance of concern.

2.7.1.3 Toxicology of the biocidal product

Toxicological data have been submitted on the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI.

The basis for the health assessment of the biocidal product is laid out in Annex 5 "Toxicology – biocidal product."

2.7.1.3.1 Percutaneous absorption

A default dermal absorption value of 10% has been used for risk assessment considering the physicochemical properties of the active substance (log Pow > 4 and molecular weight > 500 g/mol), in accordance with the recommandations of EFSA guidance on dermal absorption¹⁰.

2.7.1.3.2 Acute toxicity

Acute oral and acute dermal toxicity of the product was tested.

In the acute oral toxicity study (OECD 423), rats were exposed to a single dose of 2000 mg/kg bw.

No mortality was observed during the test.

An increased salivation was noted in 1 animal (1/6) 30 minutes after the test item administration. A noisy respiration was noted in 1 animal (1/6) at 4 and 24 hours post-dose. The animals recovered a normal behaviour at 48 hours post-dose.

The macroscopic examination of the animals at the end of the study did not reveal treatment related changes.

The LD50 of the test item SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is higher than 2000 mg/kg body weight by oral route in the rat.

In the dermal acute toxicity study (OECD 402), rats were exposed to a single dose of 2000 mg/kg bw.

No mortality occurred during the study. Neither cutaneous reactions nor systemic clinical signs related to the administration of the test item were observed.

The macroscopic examination of the animals at the end of the study did not reveal treatment-related changes.

The dermal LD_{50} of the test article SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is higher than 2000 mg/kg bw in rats.

Based on the results, no classification is required for SANITERPEN INSECTICIDE DK PRET A L'EMPLOI.

No acute inhalation toxicity study was generated for SANITERPEN INSECTICIDE DK PRET A L'EMPLOI. The product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI does not contain ingredient classified for health effects resulting from an acute exposure by inhalation. Therefore, according to the classification rules in Directive 1999/45/EC, no classification regarding acute inhalation toxicity is warranted for the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI.

2.7.1.3.3 Irritation and corrosivity

In the skin irritation study (OECD 404) in rabbits (3 animals), a slight erythema (score 0.7/0.0/0.7) and a very slight oedema (score 0.7/0.0/0.0) was noted on the treated area of 3 animals respectively, mean at 24h-48h-72h.

The erythematous and oedematous reactions were totally reversible between D3 and D8.

A slight dryness was noted in 1 animal between D3 and D8.

Based on the results, no classification is required for SANITERPEN INSECTICIDE DK PRET A L'EMPLOI.

 $^{^{10}}$ Guidance on dermal absorption, EFSA, 2012 $\,$

In the eye irritation study (OECD 405) in rabbits (3 animals), the observed ocular reactions have been moderate to severe and totally reversible in the three animals.

A moderate redness (score 1.3/1.7/1.3) was noted in the 3 rabbits respectively, mean at 24h-48h-72h and totally reversible between D4 and D8.

A moderate to severe chemosis (score 1.0/2.0/0.3) was associated and noted in the 3 rabbits respectively, mean at 24h-48h-72h and totally reversible between D2 and D4.

A moderate corneal opacity (score 2.0/1.7/0.7) was noted in the 3 rabbits respectively, mean at 24h-48h-72h and totally reversible between D2 and D4.

A congestion of the iris (score 0.0/0.3/0.3) was noted in the 3 rabbits respectively, mean at 24h-48h-72h and totally reversible between D1 and D2.

Based on the results, classification as **Eye Irrit.2 – H319** is required for SANITERPEN INSECTICIDE DK PRET A L'EMPLOI, according to the CLP regulation criteria.

2.7.1.3.4 Sensitisation

Two LLNA studies (LLNA with cell count and LLNA BrdU) in mice were submitted.

Negative skin reactions were observed under the experimental conditions.

The non radioactive cell count LLNA is not currently validated and no guideline is available. The EC determination is not possible; therefore, the first LLNA study submitted has not been accepted.

In the second study, a LLNA BrdU (OECD 442B), no mortality and no signs of systemic toxicity were noted in the test and control animals during the test.

No Stimulation Index of more than 1.6 was recorded with the test concentrations.

The Stimulation Index (SI) calculated by pooled approach was respectively 1.21, 0.93 and 0.88 for the treated groups at 25%, 50% and 100% respectively.

No cutaneous reactions were noted during the study.

No significant increase in ear thickness and in ear weight was noted in animals treated with test item.

Based on these results, no classification is required for SANITERPEN INSECTICIDE DK PRET A L'EMPLOI.

2.7.1.3.5 Other studies

No other studies submitted.

2.7.2 Human exposure assessment

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is an insecticide and acaricide containing 0.02% (w/w) deltamethrine as active substance. It is intended to be use by professional only to control flying and crawling insects and mites.

The product is a RTU product applied by spraying, only indoors in animal houses and shelters (small farming areas or association for animal protection).

The application rate is claimed to be 50 mL of product/ $1m^2$.

2.7.2.1 Identification of main paths of human exposure towards active substance from its use in biocidal product

Exposure path	Industrial use	Professional use	General public	via the environment
Inhalation	na	yes	yes	na
Dermal	na	yes	yes	na
Oral	na	na	na	na

Table 3: Main paths of human exposure

na: not applicable

2.7.2.2 Direct exposure as a result of use of the active substance in biocidal product

2.7.2.2.1 Exposure of professional users

In Annex 6, Safety for professional operators", the results of the exposure calculations for the active substance and the substance of concern for the professional user are laid out.

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is a RTU product supplied in a bottle with a hand sprayer, therefore dermal and inhalation exposure can occur only during application phase. Moreover, exposure during the cleaning of equipment is not expected.

Professional exposure during the application phase has been assessed using ConsExpo (for details, see Annex 6).

Tier	Inhalation exposure	Dermal exposure	Oral (non respirable) exposure	Total exposure		
PPE	Systemic dose	Systemic dose	Systemic dose	Systemic dose		
	mg a.i. / kg bw /day	y bw mg a.i. / kg bw /day /day		mg a.i. / kg bw /day		
Task – time frame:		Application – 120 minutes daily				
Tier 1: Without PPE	1.02 x 10 ⁻⁴	1.84 x 10 ⁻³	4.73 x 10 ⁻⁵	0.0024		

Table 4: Exposure assessment for professional users

2.7.2.2.2 Exposure of non-professional users

In Annex 7 "Safety for non-professional operators and the general public", the results of the exposure calculations for the active substance and the substance of concern for the non-professional user and the general public are laid out.

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is for professional use only.

2.7.2.3 Indirect exposure as a result of use of the active substance in biocidal product

The general public (adult, child and infant) can be potentially exposed to SANITERPEN INSECTICIDE DK PRET A L'EMPLOI *via* the oral, dermal and inhalation routes.

As SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is applied by professionals in animal houses and shelters, child and infant would not come into contact with freshly treated surfaces, so exposure is not calculated.

The exposure estimation values have been calculated on the basis of these scenarios:

- Adult and child Inhalation of volatilised residues, indoor (chronic);
- Adult and child Dermal exposure to treated surfaces (chronic).

For the scenario "Inhalation of volatilised residues by an adult" the saturated Vapour Concentration (SVC) has been calculated using the following parameters:

- Vapour pressure of deltamethrin: 1.24 x 10⁻⁸ Pa;
- The gas constant R: 8.31 J.K.mol⁻¹;
- Temperature: 293 K;
- Molecular Weight of deltamethrin: 505.2 g/mol;
- Body weight: 60kg (adult) and 23.9 kg (child) (HEEG opinion on default human factor values)¹¹;
- Inhalation rate: 16 m³/24h (adult) and 12 m³/24h (child) (HEEG opinion on default human factor values).

For the scenario "Dermal exposure of an adult to treated surfaces", the dermal exposure value has been calculated using the following parameters:

- Concentration of active substance in the product after dilution: 0.02%;
- Application rate: 50 mL/m²;
- Density of the product: 0.947;
- Transfer coefficient of dislogebable residues (dried fluids on various type of surface): 18% (TNsG on Human exposure, 2008);
- Hand surface (only palms of both hands): 410 cm²(adult) and 213.9 cm² (child) (HEEG opinion on default human factor values);
- Dermal absorption value: 10%.

As SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is applied by professionals in animal houses and shelters, infant would not come into contact with freshly treated surfaces, so exposure is not calculated.

Table 5: Secondary exposure assessment

	Inhalation exposure	Dermal exposure	Oral exposure	Total exposure
Scenario	Systemic dose	Systemic dose	Systemic dose	Systemic dose
	mg a.i. / kg bw /day	mg a.i. / kg bw /day	mg a.i. / kg bw /day	mg a.i. / kg bw /day
		Chronic exposure		

¹¹ HEEG opinion on default human factor values for use in exposure assessemnts for biocidal product, endorsed at TM II 2013.

	Inhalation exposure	Dermal exposure	Oral exposure	Total exposure	
Scenario	Systemic dose	Systemic dose	Systemic dose	Systemic dose	
	mg a.i. / kg bw /day	mg a.i. / kg bw /day	mg a.i. / kg bw /day	mg a.i. / kg bw /day	
Adult – Inhalation of volatilised residues, indoor	6.9 x 10 ⁻⁷	na	na	6.9 x 10 ⁻⁷	
Child – Inhalation of volatilised residues, indoor	1.3 x 10 ⁻⁶	na	na	1.3 x 10 ⁻⁶	
Adult – Dermal exposure with treated surface, indoor	na	1.2 x 10 ⁻⁴	na	1.2 x 10 ⁻⁴	
Child – Dermal exposure with treated surface, indoor	na	1.5 x 10 ⁻⁴	na	1.5 x 10 ⁻⁴	

2.7.2.4 Indirect exposure via residues in food

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is intended to be used only in empty animals' houses and shelters excluding thoses used by animals for human consumption. So as housing or transport vehicles used for livestock will not be treated, exposure of livestock and human exposure via food of animal origin was not assessed in this report.

2.7.3 Risk assessment for human health

The estimated exposures for the professional users are compared to the systemic AEL of deltamethrine set in the Assessment Report (0.0075 mg/kg bw/day for short, medium and long-term exposures).

2.7.3.1 Risk for direct exposure

2.7.3.1.1 Professional users

Based on the risk assessment of the active substance, the risk for professional users resulting from the intended use is acceptable for SANITERPEN INSECTICIDE DK PRET A L'EMPLOI.

Scénario	%AEL	Risk				
Application – 120 minutes						
Tier 1 Without EPI	0.0075	0.0024	32	Acceptable		

Table 6: Summary of risk characterisation for professionals

2.7.3.1.2 Non-professional users

The product is for professional use only.

2.7.3.2 Risk for indirect exposure

Based on the risk assessment of the active substance, no unacceptable risk has been identified for every indirect exposure considered scenarios. For details, see Excel document in separate appendix.

Table 7: Summar	y of risk characterisation for ge	eneral public
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Scénario	AEL (mg/kg bw/d)	Exposure (mg/kg bw/d)	%AEL	Risk					
	Chronic exposure								
Adult – Inhalation of volatilised residues, indoor	0.0075	6.9 x 10 ⁻⁷	0.01	Acceptable					
Child – Inhalation of volatilised residues, indoor	0.0075	1.3 x 10 ⁻⁶	0.02	Acceptable					
Adult – Dermal exposure with treated surface, indoor	0.0075	1.2 x 10 ⁻⁴	1.6	Acceptable					
Child – Dermal exposure with treated surface, indoor	0.0075	1.5 x 10 ⁻⁴	2.03	Acceptable					

2.7.3.3 Risk for consumers via residues

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is intended only in empty animals' houses and shelters excluding thoses used by animals for human consumption. So as housing or transport vehicles used for livestock will not treated, exposure of livestock and human exposure via food of animal origin was not assessed in this report.

2.7.4 Summary of risks characterisation of the product for human health

Risks related to the use of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI by professionals are considered acceptable during spray application. Risk related to secondary exposure is also considered acceptable.

Regarding the intended used, exposure of livestock and human exposure via food of animal origin are not expected. Therefore, a dietary risk assessment is not required.

Risk mitigation measures linked to risk assessment human health

- Apply strict hygiene measures: do not eat, drink or smoke during handling of the product and wash hands after use of the product;
- Keep out of the reach of children.

Risk mitigation measures linked to risk assessment for consumers

- Do not use on surfaces likely to be in direct contact with animals intended for food consumption, food, feed or drinks.

Required information linked to risk assessment for human health and consumers

None.

Emergency

- Inhalation: Remove victim to fresh air and keep at rest in a position comfortable for breathing. Get medical attention if symptoms occur, show this container or label.
- Skin contact: Remove contaminated clothing and shoes. Wash contaminated skin with soap and water. Get medical attention if symptoms occur.
- Eye contact: Immediately flush with plenty of water, occasionally lifting the upper and lower eyelids. Check for and remove any contact lenses. Continue to rinse with warm water for at least 10 minutes. Get medical attention if irritation or vision impairment occur.
- Ingestion: Wash out mouth with water. Get medical attention if symptoms occur, show this container or label.
- Do not drink or induce vomiting in case of consciousness alteration.
- Note to physician: Treat symptomatically. Contact poison treatment specialist immediately if large quantities have been ingested or inhaled.

Disposal considerations

None.

2.8 Risk assessment for the environment

The summary of information on the active substance properties is carried out with the data from the Competent Authority Report (CAR) of deltamethrin supplied by Bayer Environmental Science SAS. (Competent Authority Report According to Directive 98/8/EC, Active substance in Biocidal Products, deltamethrin CAS 52918-63-5, Product Type 18 (Insecticides, acaricides and products to control other arthropods), RMS Sweden, May 2011).

2.8.1 Fate and distribution in the environment of the active substance deltamethrin

2.8.1.1 Degradation

2.8.1.1.1 Abiotic degradation

2.8.1.1.1.1 Hydrolysis in function of pH

For the active substance deltamethrin, no significant hydrolysis was observed at pH 5 and 7. At pH 9, however, the hydrolysis was significant with a half-life of 2.5 days at 25°C (7 days at 12°C). At pH 8, half-life was 31 days at 23°C (75 days at 12°C).

2.8.1.1.1.2 Photolysis in water

Direct photochemical reactions do not occur at a rate that makes this a significant route of degradation of deltamethrin under natural conditions in water.

2.8.1.1.1.3 Photolysis in soil

In soil, direct and indirect photochemical reactions may contribute to the degradation of deltamethrin, but other routes of transformation account for the major loss of parent compound.

2.8.1.1.1.4 Photodegradation in air

The photo-oxidative degradation of deltamethrin in air was estimated by a structural activity relationship (QSAR) method using the Atmospheric Oxidation Program v1.91 (AOPWIN). Half-life for reaction with OH-radicals was calculated to 16.4 hours.

2.8.1.1.2 Biotic degradation

2.8.1.1.2.1 <u>Aquatic compartment</u>

• Ready biodegradation / inherent biodegradation

Deltamethrin is not readily biodegradable under OECD 301F Test (degradation of 0% after 28 days).

• Degradation in water/sediment system

A higher tier water/sediment simulation study in two different water/sediment systems has been conducted and provides information on aerobic aquatic degradation of deltamethrin. 60% of the applied radioactivity was found in the sediments immediately after application. The total system degradation DT_{50} of deltamethrin in two different water/sediment systems was estimated to 85 and 267 days at 12°C, and the dissipation DT_{50} in sediment to 104 and 253 days at 12°C. The difference in degradation rate between the two systems probably reflects difference in amount of fine-textured material and amount of organic matter.

2.8.1.1.2.2 Degradation in STP

No study on the degradation of the active substance in STP has been submitted in the CAR of deltamethrin.

2.8.1.1.2.3 <u>Terrestrial compartment</u>

• Aerobic degradation

Four laboratory studies on degradation in soils have been submitted in the CAR of deltamethrin, and one further study presented calculations of rate of degradation for the relevant metabolite Br_2CA (> 10%) based on data from the four laboratory studies. In one additional study the rate of degradation for deltamethrin and its relevant metabolite were re-calculated using more appropriate approaches than in the original studies.

Deltamethrin is relatively rapidly degraded in soil, with a geometric mean DT_{50} value of 48 days at 12°C. The main metabolite of deltamethrin was Br_2CA . It was detected in available studies, up to 23% of applied radioactivity after about 2 weeks of incubation. No other metabolites were detected at levels of > 10% of applied radioactivity. When normalised to 12°C, the geometric mean of DT_{50} value for Br_2CA was 5.6 days.

• Anaerobic degradation

No study on the anaerobic degradation of deltamethrin in soil has been submitted in the CAR.

2.8.1.2 Distribution

Deltamethrin is very strongly adsorbed to soil and other organic matter, with an arithmetic mean Koc value of 408 250 L.kg⁻¹. The relevant metabolite is more mobile with an arithmetic mean Koc value of 25.6 L.Kg⁻¹.

2.8.1.3 Accumulation

The bioaccumulation of ¹⁴C-deltamethrin was investigated in bluegill sunfish (*Lepomis macrochirus*). The BCF_{fish} values obtained were 310, 2800 and 1400 L.Kg⁻¹ for edible, non-edible and whole body tissue, respectively. After the 14-day depuration period 70, 75 and 76% of the ¹⁴C-residues had been eliminated from the edible, non-edible and whole body tissue, respectively. The biological half-life was 4.3 days for whole body tissue.

No experimental data are available for terrestrial bioconcentration. Therefore, the terrestrial BCF have been estimated using a linear Quantitative Structure Activity Relationship (QSAR) model and the log P_{ow} of deltamethrin of 4.6 at 25°C. The BCF_{earthworm} was 483 L.kg⁻¹ (according to TGDII Equation 82d).

2.8.1.4 Behaviour in air

Due to its low vapour pressure, deltamethrin is not expected to volatilise to air from plants and soil at significant levels, which was confirmed in a wind tunnel study. However, the calculated Henry's law constant is 1.252×10^{-3} Pa.m³.mole⁻¹, indicating that deltamethrin has a tendency to volatilise from water. If present in air, the data on indirect photo-oxidation indicate a rapid degradation when reacting with hydroxyl radicals.

2.8.2 Effects on environmental organisms for active substance deltamethrin

No new ecotoxicological information on the active substance deltamethrin has been submitted in the product dossier compared to the CAR.

2.8.2.1 Aquatic compartment (including water, sediment and STP)

2.8.2.1.1 Aquatic organisms

The table below summarises all the data available for the active substance deltamethrin. The metabolite Br2CA is considered not relevant for the aquatic compartment. Moreover it has been demonstrated in the CAR that the risk assessment for the metabolite, Br2CA was covered by the risk assessment for deltamethrin.

Test item	Species	Guideline	Endpoints	Toxicity [µg.L ⁻¹]	Reference				
Fish	Fish								
Deltamethrin	Onchorhynchus mykiss	OECD 203	LC ₅₀ – 96h Flow-through conditions	0.26 ¹	A.7.4.1.1/02				
	Pimephales promelas	US EPA 72-5	NOEC – 260d	0.017 ¹	A.7.4.3.2/02				
Invertebrates	Invertebrates								
Deltamethrin	Gammarus fasciatus	US EPA	LC ₅₀ – 96h Flow-through conditions	0.0003 ¹	A.7.4.1.2/02				

Table 8: Existing endpoints for aquatic organisms

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Test item	Species	Guideline	Endpoints	Toxicity [µg.L ⁻¹]	Reference				
	Daphnia magna	OECD 211	NOEC – 21d Flow-through conditions	0.0041 ¹	A.7.4.3.4/01				
	Chironomus riparius	BBA 1995 NOEC – 28d		0.0035 ¹	A.7.4.3.5.1/01				
Algae	Algae								
Deltamethrin	Chlorella vulgaris	Brazilian method D.4.1	$EbC_{50} - 96h$ $ErC_{50} - 96h$ NOErC Static conditions	>0.47E03 ¹ >0.47 E03 ¹ 0.47 E03	A.7.4.1.3/02				
Higher tier stu	Higher tier studies								
Deltamethrin	water flea	Mesocosm guidance ³	NOEC Mesocosm conditions	0.0048 ²	A.7.4.3.5.3				

¹ measured concentrations ² nominal concentrations

³ OECD 2004 "Simulated Freshwater Lentic Field Tests (Outdoor Microcosms and Mesocosms)"

Additional endpoints: not relevant

Justification of PNEC_{water}

According to the TGD for Risk Assessment (2003), and using the lowest chronic laboratory NOEC value (3.5 ng.L⁻¹) and an assessment factor of 5 (considering that the test organism had been identified as the most sensitive), the PNEC_{water} is 0.7 ng L^{-1} .

2.8.2.1.2 Sediment dwelling organisms

Justification of PNEC_{sediment}

The PNEC_{sediment} is estimated from PNEC_{water} in using the Equilibrium Partitioning Method (according to the TGD for Risk Assessment (2003)) with the mean Koc value for deltamethrin of 408 250 L.Kg⁻¹. The PNEC_{sediment} is 6.2 µg kg wwt⁻¹.

2.8.2.1.3 STP micro-organisms

The table below summarises the data available for the active substance deltamethrin.

		Species/ Exposu		Exposure	Result [mg a.s.L ⁻¹]			
Test item	Guideline	•	design	•	NOEC	EC ₅₀	EC ₈₀	reference
Deltamethrin	OECD 209	Activated sludge	Respiration inhibition	3h		>300	-	A.7.4.1.4/01
Deltamethrin	OECD 209	Activated sludge	Respiration inhibition	3h	>0.3	>0.3		A.7.4.1.4/02

Table 9: Existing endpoints for STP micro-organisms

Additional endpoints: not relevant

Justification of PNEC_{STP micororganisms}

According to the TGD for Risk Assessment (2003), and taking into account that deltamethrin had no significant effect at the highest tested concentration (NOEC $\ge 0.3 \text{ mg L}^{-1}$), an assessment factor of 10 can be applied. Thus, the PNEC_{microorganisms} is 30 µg.L⁻¹.

2.8.2.2 Atmosphere

Significant exposure of the environment via air is not expected.

Due to its low vapour pressure, deltamethrin is not expected to volatilise to air from plants and soil at significant levels, which was confirmed in a wind tunnel study. However, the calculated Henry's law constant is 1.252×10^{-3} Pa.m³.mole⁻¹, indicating that deltamethrin has a tendency to volatilise from water. If present in air, the data on indirect photo-oxidation indicate a rapid degradation when reacting with hydroxyl radicals (DT₅₀ reaction with OH-radicals = 16.4 hours). It is thus considered that it is not likely that significant volatilisation will occur after use of deltamethrin.

2.8.2.3 Terrestrial compartment

The table below summarises all the data available for the active substance deltamethrin and its relevant metabolite, Br_2CA .

Test item	Guideline/ Test method	Species inoculums	Endpoint / type of test	Exposure design duration	Results	reference
ACUTE						
Deltamethri n	OCDE 207	Eisenia fetida	LC50 _{mortality}	14d - Artificial soil	> 1290 mg/kg ⁻¹ _{dw}	A.7.5.1.2/0 1
CHRONIC						
Br ₂ CA	SECOFAS E (1996)	Hypoaspis aculeifer	NOEC _{mortality} Br ₂ CA mixed with LUFA 2.1 soil	14d	10 mg/kg ⁻¹ _{dw soil}	A.7.5.2.1/0 1
	BBA VI 2-2	Eisenia fetida	NOEC _{reproduction}	56d - Artificial soil	0.78 mg/kg ⁻¹ _{dw soil}	A.7.5.2.1/0 2
Deltamethri	ISO 11267	Folsomia candida	NOEC _{mortality}	28d - Artificial soil	1.25 mg/kg ⁻¹ _{dw soil}	A.7.5.2.1/0 3
n	Hypoaspis ring-test (SETAC, 2005)	Hypoaspis aculeifer	NOEC _{mortality} and NOEC _{reproduction}	16d - Artificial soil	1.78 mg/kg ⁻¹ _{dw soil}	A.7.5.2.1/0 4
	BBA VI, 1-1	Microorganism s	NOEC- Effect on aerobic respiration in 2 soils	28/56-d	>0.50 mg/kg ⁻¹ _{dw} _{soil} equivalent to > 375 g/ha	A.7.5.1.1/0 1

Table 10: Toxicity so soil organisms

Test item	Guideline/ Test method	Species inoculums	Endpoint / type of test	Exposure design duration	Results	reference
	BBA VI, 1-1	Microorganism s	NOEC - Effect on N cycle in 2 soils	28d	>0.50 mg/kg ⁻¹ _{dw} _{soil} equivalent to > 375 g/ha	A.7.5.1.1/0 2

Additional endpoints: not relevant.

Justification of PNEC_{soil}

Due to the lack of effects in the tests on micro-organisms and chronic toxicity to earthworms, the PNEC is based on the NOEC from the reproduction test on springtails.

The results are converted to standard soil which is defined as a soil with an organic matter content of 3.4% using the following equation:

$$\begin{split} \text{NOEC}_{\text{standard}} &= \text{NOEC}_{\text{exp}} \times \text{F}_{\text{om, soil standard}} / \text{F}_{\text{om, soil exp}} \text{ (TGD, part II, Eq. 71)} \\ & \text{With} \qquad \text{NOEC}_{\text{exp}} = 1.25 \text{ mg.kg}^{-1} \text{ dry soil} \\ & \text{F}_{\text{om, soil standard}} = 3.4 \% \end{split}$$

 $F_{om, soil exp} = 5 \%$

Then, NOEC_{standard} = 0.85 mg.kg^{-1} dry soil

An assessment factor of 10 can be applied. Thus, the following $PNEC_{soil}$ is derived:

 $PNEC_{soil} = 85 \ \mu g.kg^{-1} \ dry \ soil (75 \ \mu g.kg^{-1} \ wet \ soil)$

No $PNEC_{soil}$ was derived for the metabolite Br_2CA , since toxicity results show that the parent compound is more toxic and more persistent than this metabolite. Therefore the risk assessment of this metabolite is covered by the active substance for the soil compartment.

2.8.2.4 Effects on honeybees

No data, the exposure of deltamethrin to honeybees is expected to be very limited.

2.8.2.5 Non compartment specific effect relevant to the food chain

The table below summarises the data available for the active substance deltamethrin:

Test item	Guideline/Te st method	Species	Test/ Duration	Results	reference		
Birds	Birds						
Deltamethrin	US EPA FIFRA E 71-1	Bobwhite quail (Colinus virginianus)	Acute oral LD ₅₀	>2250 mg.kg ⁻¹ _{bw}	A.7.5.3.1.1/01		
	Conducted	Mallard duck	Acute oral	> 4640 mg.kg ⁻¹ _{bw}	A.7.5.3.1.1/02		

Table 11: Toxicity to birds and mammals

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Test item	Guideline/Te st method	Species	Test/ Duration		Results	reference		
	before an appropriate guideline	(Anas platyrhynchos)	LD ₅₀		LD ₅₀			
	US EPA 71-2 / OECD 205	Bobwhite quail (Colinus virginianus)	Dietary 5-day LC ₅₀		> 5620 mg/kg ⁻¹ _{diet}	A.7.5.3.1.2/01		
	US EPA 71-2 / OECD 205	Mallard duck (Anas platyrhynchos)	Dietary 5-day LC ₅₀		8039 mg/kg ⁻¹ _{diet}	A.7.5.3.1.2/02		
	US EPA 71-4; OECD 206	Bobwhite quail (Colinus virginianus)	Reproduction 22-week NOEC	n	> 450 mg/kg ⁻¹ _{diet} (55 mg.kg ⁻¹ _{bw} d ⁻¹)	A.7.5.3.1.3/01		
	US EPA 71-4; OECD 206	Mallard duck (Anas platyrhynchos)	Reproduction 22-week NOEC	on	> 450 mg/kg ⁻¹ _{diet} (70 mg.kg ⁻¹ _{bw} d ⁻¹)	A.7.5.3.1.3/02		
Mammals								
Deltamethrin	OECD 401	rat	LD ₅₀	Oral	95 mg.kg ⁻¹ _{bw} (males) 87 mg.kg ⁻¹ _{bw} (females)	A6.1.1/01		
	OECD 416	rat	NOAEL	Oral	80 ppm	A6.8.2/01		

Justification of PNECoral,bird and PNECoral,mammal for secondary poisoning

The PNEC_{bird} and the PNEC_{mammals} calculations are based on a long-term toxicity / reproduction study with bird and on a 2 generation toxicity test on rat respectively. According to the TGD for Risk Assessment (2003), an assessment factor of 30 for bird and mammal can be applied. Thus, the following PNEC_{oral} are derived:

 $PNEC_{oral,bird} = 15 \text{ mg.kg}^{-1}_{diet}$ $PNEC_{oral,mammal} = 2.67 \text{ mg.kg}^{-1}_{diet}$

2.8.2.6

Table 12: Summary of PNECs of the active substance deltamethrin

Summary of PNECs of the active substance deltamethrin

Compartment	Species	Endpoint	Safety factor	PNEC		
Surface water	Chironomus riparius	NOEC – 28d = 3.5 n.gL ⁻¹	5	0.7 ng.L ⁻¹		
Sediment	6.2 μg.kg ⁻¹ ww sediment (equilibrium partitioning)					
Microorganisms (STP)	Activated sludge	NOEC ≥ 0.3 mg L^{-1}	10	30 µg.L ⁻¹		

Soil	Folsomia candida	NOECstandard = 0.85 mg.kg ⁻¹ _{dry soil}	10	75 μg.kg ⁻¹ wet soil
Bird	Colinus virginianus Anas platyrhynchos	NOEC > mg/kg ⁻¹ _{diet}	30	15 mg.kg ⁻¹ diet
Mammal	Rat	NOAEL = 80 ppm	30	2.67 mg.kg ⁻¹ diet

2.8.2.7 PBT and ED Assessment

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According to the PBT assessment in TGD, criteria for substance to be persistent (P) and very persistent (vP) are fulfilled when:

- T 1/2 in freshwater sediment > 120 days for P;
- T 1/2 in freshwater sediment > 180 days for vP.

Results of a simulation test on two different water/sediment systems show that deltamethrin partitions very rapidly to sediment. The degradation half-lives of deltamethrin in both systems were 85 and 267 days at 12°C (degradation in the whole water/sediment systems).

It can be concluded that in one system the DT_{50} normalised to 12°C exceeds the P- and the vP-criteria for freshwater sediment – while in the other system the DT_{50} normalised to 12°C does not exceed any of P / vP criteria. With one value below the criteria and one value above, it cannot be concluded whether deltamethrin should be classified as persistent in the sediment compartment or not.

Laboratory data from four different soils were available; the DT_{50} s ranged from 31 to 74 days (12°C). Hence, none of the half-lives for degradation in soil exceeds the P-criterion for soil (i.e. > 120 days). It is concluded that deltamethrin cannot be classified as persistent in the soil compartment.

Both environmental compartments for which there is adequate data available (i.e., sediment and soil) are considered to represent relevant environmental compartments since emissions may occur to both systems, via release from STP or application of sludge. Based on the data above **it can be concluded that deltamethrin potentially fulfils the criteria for persistence**.

According to the PBT assessment in TGD, a substance is considered to fulfill the B criterion when the bioconcentration factor (BCF) exceeds a value of 2 000 L/kg. In a BCF study done with *Lepomis macrochirus*, the steady-state BCF for uptake of deltamethrin estimated in whole fish was 1400 L/kg. Considering this result, **deltamethrin is not selected according to the B criterion**.

According to the PBT assessment in TGD, the toxicity criterion is fulfilled when the chronic NOEC for aquatic organism is less than 0.01 mg.L⁻¹ or when the substance is toxic to mammals and classified as Very Toxic or Toxic after oral dosing. Based on ecotoxicity freshwater data on water flea, NOEC = 4.8 ng.L^{-1} , **T criterion is fulfilled**.

As the B criterion is not fulfilled and only the T criterion is clearly fulfilled, deltamethrin is not classified as a PBT.

2.8.3 Effects on environmental organisms for biocidal product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

The applicant did not provide ecotoxicological data about the biocidal product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI. The risk assessment for the product is based on the data obtained from the active substance deltamethrin (Competent Authority Report According to Directive 98/8/EC, Active substance in

Biocidal Products, deltamethrin CAS 52918-63-5, Product Type 18 (Insecticides, acaricides and products to control other arthropods), RMS Sweden, May 2011).

No classified compound other than the active substance is present in the biocidal product. Therefore, FR CA considered that the effects of deltamethrin outweigh those of the non-active components of the product and that the effects assessment for the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI can be extrapolated from the effects assessment of the active substance deltamethrin.

2.8.3.1 Aquatic compartment (including water, sediment and STP)

2.8.3.1.1 Aquatic organisms

No additional data. Refer to section 2.8.2.1.

2.8.3.1.2 Sediment dwelling organisms

No additional data. Refer to section 2.8.2.1.

2.8.3.1.3 STP micro-organisms

No additional data. Refer to section 2.8.2.1.

2.8.3.2 Atmosphere

No additional data. Refer to section 2.8.2.2.

2.8.3.3 Terrestrial compartment

No additional data. Refer to section 2.8.2.3.

2.8.3.4 Non compartment specific effect relevant to the food chain

No additional data. Refer to section 2.8.2.5.2.8.2.5

2.8.3.5 Summary of PNECs

No additional data. Refer to section 2.8.2.6.

2.8.4 Environmental exposure assessment

The product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI contains 0.2% w/w of deltamethrin. It is used by professionals only. The product is applied by spraying and is intended to control insects and mites. SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is applied only indoors in empty animal houses and shelters.

Targeted animal houses and shelters are small houses which are used in rural areas by farmers, hunters, agricultural workers, or houses and shelters used for small farming (dog farming, ornamental chicken farming), or by small associations working for the protection of the animals.

Animals which are intended to be protected with the product are not intended to be used for consumption. Only pets (i.e. horses, dogs, rabbits...) environment is intended to be treated.

The product is not intended to be used in big animal farming or industrial farming. The packaging is not intended for such an activity (3 x 60mL, 1L, 5L).

The intended uses of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI can be covered in one environmental exposure scenario as the profile of users and location of use are the same for all the pets' environment location which are treated with the product.

The recommended application dose is $5 \text{ mL}_{product.m}^2$ or $10 \text{ mg}_{a.s..m}^2$. The efficacy against target insects lasts for 3 months after application in field conditions. During winter there is no insect infestation due to climatic condition. The product shall be applied only 3 times per year maximum.

2.8.4.1 Assessment of exposure to the environment: Professional Use of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI for treatment of pet's environment in rural areas

The environmental risk assessment is made according to the ESD n°18 "Emission scenario document for insecticides, acaricides and products to control other arthropods for household and professional uses" (2008).

The ESD n°18 relates to consumer uses (in households) and professional uses (in other buildings). Nonprofessional users are excluded in the scenario presented below as the product is intended to be used by professional only.

"Other buildings" shall be clearly defined in our case. Buildings which are treated are only:

- Animal housing in rural areas (for example horse shelter, ornamental chickens shelter, dog shelters);
- Animal shelters for small animal farming intended for pet production or pet protection (dog kennels, rabbit houses).

Another ESD on the use of insecticides is available: ESD n°14 "Emission scenario document for Stables and Manure Storage Systems" which covers insecticide applications in industrial farming (production of livestock, poultry...). It was considered that this ESD was not fully adapted for the uses of SANITERPEN INSECTICIDE PRET A L'EMPLOI described above (small animal housing, generally up to 10 pets).

Details on use of the product.

Number of application and dose

Three applications per year every 3 months are required to manage insect infestation with SANITERPEN INSECTICIDE DK PRET A L'EMPLOI. The application dose is 50 mL $_{product}/m^2$, corresponding to 10 mg $_{a.s.}/m^2$ (considering 0.2% deltamethrin in the product). During application, the surfaces are treated only once with the product.

Surface

The maximum surface to be treated per application is proposed by the applicant to be 350 m². This value is conservative as far as the product is sold in a 750mL format which cans covers $15m^2$.

No literature exists on the areas of specific type of building which is targeted by SANITERPEN INSECTICIDE DK PRET A L'EMPLOI but usual surfaces (floor, walls and ceilings) which are encountered in rural areas are:

- Approximately 200 m² for horse shelters
- Approximately 150 m² for kennels
- Approximately 90 m² for rabbits hutches
- Approximately 100 m² for chicken coops.

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is is relevant only for the very small surfaces to be treated. Ready to-use products avoid any exposure resulting from mixing/loading phases.

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is intended to be used by professionals only. Limitation of application to a maximum surface of 350 m² is a risk management measure clearly stated on the label of the product: "SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is designed to be used on a max. surface of 350 m² per application. If treatment requires a higher surface treatment, another solution shall be investigated'.

Equipment

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is intended to be used only inside its original packaging which is a low pressure hand sprayer.

Operating instructions

Different risk mitigations are proposed by the applicant before the use of the product. They are reported below in order to have a clear view of the operating instructions to include in the exposure assessment:

Before using SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

- Buildings, houses and shelters shall be cleared from all animals, animal food and water. Textiles (pillows, blankets...) shall be taken away and washed thoroughly.
- Floor and walls, including previously treated surfaces, shall be mechanically cleaned from dusts using vacuum system or brush and dusts shall be collected and treated as a waste.
- Depending on the case, risk management measures regarding resistance (see section 6 of the IUCLID dossier) shall be considered.
- Surface to be treated shall be determined and calculated regarding specific cases (pets to be protected, insects to be controlled). In all case it should be limited to 350 m².

Application

- Application shall be made with the original packaging. Surfaces shall be treated only once during the application.

After application of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

- No cleaning of the equipment is required.
- SANITERPEN INSECTICIDE PRET A L'EMPLOI must be kept inside the original packaging. After application treated surfaces shall not be cleaned (cleaning will take place before the next application).
- Animals shall be re-introduced only when the surfaces are dried (max. 24hours). Textiles shall be reintroduced also only when animals are re-introduced.

Details on the Exposure assessment

In the calculations, default values (according to the TGD) and updated data from MOTA (Manual of Technical Agreements – version 6) were used, unless submitted data were available in the dossier. All deviations are detailed and justifications are provided below. Calculations are based on the maximum amount used per application as a worst-case scenario.

The assessment has been carried out for the **application phase** and for the subsequent **cleaning phase**, as it is indicated by the applicant that the surface must be cleaned before a new application (but not between applications). In his proposal, the applicant carried out an environmental exposure assessment in considering specific operating instructions. The treated surfaces shall be mechanically cleaned using vacuum system or brush. And in any case, never throw rinsing water in the waste water system or surface. Consequently in the applicant evaluation, a release to wastewater was only taken into account from applicator emission. Nevertheless, in order to evaluate the relevance of these operating instructions, an exposure assessment with emission to wastewater from floor and treated surfaces has been added.

To complete the assessment, a potential emission into the environment *via* a potential **application of manure** (for example to cover horse manure application) on arable land and grassland has been taken into account. It should be noted that this scenario taken from the ESD n°14 "Emission scenario document for Stables and

Manure Storage Systems" is considered as a very worst case approach as the emission calculations form this document cover industrial farming.

Release to STP via wastewater after cleaning of treatment surfaces

- Number of buildings per STP and simultaneity factor

The applicant proposed a value of 100 pets-housing per STP combined with a simultaneity factor of 0.82% (1 to 11 applications per year according to the ESD) considering the maximal intended application number of 3 per year. These proposals were considered acceptable. Nevertheless, this leads to less than on building cleaned per day, which is not realistic. Consequently, one building cleaned per day at the STP scale has been considered in the environmental exposure assessment for the use of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI product.

- Cleaning efficiency (Spray-Surface)

A cleaning efficiency of 50% was used as proposed for a spray-surface application in the ESD n°18 for household insecticides (Table 3.3-8)

Parameter	Symbol	Value	Unit	Source		
Product Information						
Product Name	(-)	SANITERPEN INSECTICIDE DK PRET A L'EMPLOI	(-)	(-)		
Active Ingredient	(-)	Deltamethrin	(-)	(-)		
Fraction of deltamethrin in product	Fai	0.002	(-)	Input		
Application dose of Product	Q _{prod}	50	mL.m ⁻²	Input		
Treatment Rate of deltamethrin	Qai	0.01	g _{as} .m ⁻²	Output		
Quantity of commercial product used for the preparation, building	Qprod, prep	1.75E+04	g	Output		
Product formulation and container type	(-)	Liquid 5 liters (D= 45/63 mm)	(-)	Pick-list		
Treatment Sub-category	(-)	Surface spray / Total Surface	(-)	Pick-list		
User Type	(-)	Professional	(-)	Pick-list		
Area of treated surface, larger building	AREAtreated	350	m ²	Calculated		
Indoor application by spraying - Application						
Number of applications per day, building Fraction emitted to air Fraction emitted to applicator Fraction emitted to floor Emission to air Emission to applicator Emission to floor	N _{appl} Fappli,air Fappli,applicator Fappli,floor+treated Eapplication,air Eapplication,applicator Eapplication,floor+treated	1 0.02 0.006 0.974 7.00E-05 2.10E-05 3.41E-03	(-) (-) (-) Kg.d ⁻¹ Kg.d ⁻¹ Kg.d ⁻¹	Default Default Default Output Output Output Output		
Indoor application by spraying – Cleaning be	fore the next applicati	ion				
Fraction emitted to wastewater from applicator (washable coveralls) Fraction emitted to wastewater from floor / treated surface during the cleaning step Number of animal housings per STP	Fapplicator,ww Ftreated surface,ww Nbuildings/STP	1 1 1	(-) (-) (-)	Default		
Cleaning efficiency (Spray-Surface)	F _{CE}	0.5	(-)	Default		
Release to waste water for one building and one application						
Emission from applicator to waste water	E _{applicator,ww}	2.10E-05	Kg.d⁻¹	Output		
$E_{applicator,ww} = E_{application,applicator} \times F_{applicator,ww}$						
Emission from floor/treated to waste water during the cleaning step	E treated surface, ww	1.70E-03	Kg.d ⁻¹	Output		

Table 13: Release to wastewater for the application and the cleaning phases

Parameter		Symbol	Value	Unit	Source
	$E_{treated,ww} = E_{application,floor+t}$	rreated $ imes F_{treated}$ surfa	$_{ce,ww} \times N_{buildings/STP} \times F_{CE}$		

Release to soil via application of manure

The emission estimation is based on the ESD n°14 "Emission scenario document for Stables and Manure Storage Systems", which is considered as a very worst case approach as the emission calculations from this document cover industrial farming. The scenario is calculated for the worst case scenario "veal calves" described in the ESD, considering default values for this animal category. The input parameters and the results of the emission calculation are reported below, considering three applications per year maximum without taking into account of winter, as there is no insect infestation during winter due to climatic condition, which leads to a treatment interval of 3 months.

Table 14: Emissions to soil via manure spreading - Input parameters for the emission scenario PT18 - Disinfection of animal houses by spraying – professional use (veal calf)

Variable/parameter	Unit	Symbol	S/D/O	Value
INPUT				
Type of animal house	[-]	cat-subcat (i1)	D	Veal Calves
Type of application	[-]	appway (i3)	D	Spraying
Area of the housing : Wall and floor	[m ²]	AREA	D	330
Content of a.i. in product	[g.L ⁻¹]	Fbioc	S	0.2
Amount of) product prescribed to be used per m ²	[L.m ⁻²]	Vprod	S	5.00E-02
Dilution factor	[g.L ⁻¹]	F _{dil}	D	1
Fraction of a.i. released	[-]	F _{slurry/manure}	D	0.5
OUTPUT				
Amount of a.i. to be used for one application	[kg]	Qai-prescr	0	3.30E-03
Amount of a.i. in slurry/manure after one application	[kg]	Qaislurry/manure t0	0	1.65E-03
CALCULATIONS				
Emission to manure/ slurry :				
Qai-prescr _{i1,i2,i3} = 10^{-3} * Fbioc * Vprod * F _{dil} * AF	REA i1			
Qai _{manure} = Qai-prescr * F _{slurry/manure}				
Number of disinfectant applications in one year	[-]	Napp-bioc	D	3
Biocide application interval	[d]	Tbioc-int	D/O	92
Number of manure applications - grassland	[-]	Nlapp-grass	D	4/1*
Number of manure applications - arable land	[-]	Nlapp-arab	D	1
Manure application time interval for grassland	[d]	Tgr-int	D	53
Manure application time interval for arable land	[d]	Tar-int	D	212
Number of animals	[-]	Nanimal i1	D	80
Amount of nitrogen per animal	[kg.d ⁻¹]	Qnitrog i1	D	2.38E-02
IF NITROGEN IMMISSION STANDARDS ARE APPLIED				
Nitrogen immission standard for one year - grassland	[kg.ha ⁻¹]	$Q_{N,grassland}$	D	170
Nitrogen immission standard for one year - arable land	[kg.ha ⁻¹]	$Q_{N,arable_land}$	D	170
Mixing depth with soil - grassland	[m]	$DEPTH_{grassland}$	D	0.05
Mixing depth with soil - arable land	[m]	DEPTHarable_land	D	0.2
Density of wet bulk soil	[kg.m ⁻³]	RHO _{soilwet}	D	1700
Intermediate Calculations		N		
Number of biocide applications - grassland	[-]	Napp-manure	0	1

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Variable/parameter	Unit	Symbol	S/D/O	Value	
Number of biocide applications - arable land	[-]	Napp-manure arable land	0	3	
Amount of active ingredient in manure - grassland	[kg]	Qai-grass _{i1,i2,i3,i4}	0	1.65E-03	
Amount of active ingredient in manure - arable land	[kg]	Qai-arab _{i1,i2,i3,i4}	0	4.95E-03	
Amount of nitrogen - grassland	[kg]	Qnitrog-grass _{i1,i4}	0	1.01E+02	
Amount of nitrogen - arable land	[kg]	Qnitrog-arab _{i1,i4}	0	4.04E+02	
OUTPUTS					
Soil exposure					
Initial concentration of a.i. in soil - nitrogen - grassland	[mg.kg ⁻¹ wwt]	PIECgrs-N _{i1,i2,i3,i4}	0	3.27E-03 (1 appl) 8.17E-04 (4 appl.)	
Initial concentration of a.i. in soil - nitrogen - arable land	[mg.kg ⁻¹ wwt]	PIECars-N _{i1,i2,i3,i4}	0	6.13E-04	
CALCULATIONS Intermediate Calculations Napp-manure (for grassland and arable land):					
If Tbioc-int > Tgr/ar-int, then Napp-manure = 1 If Tbioc-int < Tgr/ar-int, then Napp-manure = ROUND (Tgr/ar-int/Tbioc-int)					
$\begin{array}{l} \mbox{Qai-grass}_{i1,i2,i3,i4} = \mbox{Qai}_{manure (DEG)} * \mbox{Napp-manure}_{gr} \\ \mbox{Qai-arab}_{i1,i2,i3,i4} = \mbox{Qai}_{manure (DEG)} * \mbox{Napp-manure}_{ar} \end{array}$					
Qnitrog-grass _{i1,i4} = Nanimal _{i1} * Qnitrog _{i1} * Tgr-int _{i2}					
$Qnitrog-arab_{i1,i4} = Nanimal_{i1} * Qnitrog_{i1} * Tar-int_{i2}$					
End calculations					
Soil exposure	_				
$PIECgrs - N_{i1,i2,i3,i4} = \frac{100 \times Qai - grass_i}{Qnitrog - grass_{i1,i4} \times Nl_{app-grass_{i1}}}$	1,i2,i3,i4 × Q _N ,grass	sland and × RHOs oil:			
$100 \times Qai$ - and	ss − − − grassi	unu · · · · · · · · · · · · · · · · · ·			
$PIECars - N_{i1,i2,i3,i4} = \frac{100 \times Qai - arab_{i1,i2}}{Qnitrog - arab_{i1,i4} \times Nl_{app-arab}}$,i3,i4 ^ YN,arable- × DEPTH _{arab-la}	^{land} nd × RHOsoil _{wet}			

* The ESD indicates applications on grassland in 4 times; nevertheless a worst case application in only one go was considered here for the soil contamination. For surface water contamination via run-off, it was deemed more realistic to take the PEC value after only one application on 4.

2.8.4.2 PEC calculations

Emission to the environment can occur either through release to STP, or through release to manure. These two ways of emission induce different environmental concentrations.

According to the deltamethrin dossier, the following distribution characteristics in the STP (determined from the SimpleTreat 3.1 Model) and the following physicochemical parameters have been used. Moreover calculations have been adjusted to consider a SLUDGE RATE value of 710 kg.d⁻¹ instead of 790 kg.d⁻¹ integrated in the EUSES program.

Fate	% of residue
to air	0.0
to water	9.6
to sludge	90.4
degraded	0.0
Total	100.0

Physico-chemical parameter	Value
Organic carbon-water partition coefficient	408 250 L kg-1
Henry's law constant	1.252 x 10-3 Pa.m3.mol-1
Rate constant for biodegradation in soil	1.44E-02 (12°C)

No PEC was derived for the relevant metabolite Br_2CA in the aquatic or terrestrial compartment, since toxicity results show that the parent compound is more toxic and more persistent than this metabolite. Nevertheless, predicted concentrations in porewater have been estimated for this metabolite, considering a Koc value of 25.61 L.Kg⁻¹ and a DT₅₀ in soil at 12°C of 5.6 days.

2.8.4.2.1 Aquatic compartment (surface water, sediment, STP)

Two ways of aquatic contamination are foreseen after application and cleaning of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI:

- via the STP, if the surfaces are cleaned with water,
- via the contaminated manure application on agricultural soil and potential run-off to waterbodies.

RELEASE VIA THE WASTE WATER (STP)

The concentrations of deltamethrin in the STP effluent, in surface water and in sediment are calculated according to the TGD equations considering the emissions to waste water calculated from indoor applications of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI in empty animal houses and shelters.

Symbol	Parameter	Value	Unit	Reference
E _{applicator,ww}	Emission from applicator to waste water	2.10E-05	[kg.d ⁻¹]	Output
PEC _{STP}	PEC in the treated wastewater	1.01E-06	[mg.L ⁻¹]	TGD Eq. 33
PEClocalwater	PEC in water during emission episode	6.25E-08	[mg.L ⁻¹]	TGD Eq. 45
PEClocal _{sed}	PEC in sediment during emission episode	5.55E-04	[mg.kg ⁻¹ _{wwt}]	TGD Eq. 50

Table 16: PECs in the aquatic compartment - Emission from applicator to STP

Table 17: PECs in aquatic compartment - Emission from surface cleaning phase to STP

Symbol	Parameter	Value	Unit	Reference
E treated surface, ww	Emission from floor/treated to waste water during the cleaning step	1.70E-03	[kg.d ⁻¹]	Output
PEC _{STP}	PEC in the treated wastewater	8.18E-05	[mg.L ⁻¹]	TGD Eq. 33
PEClocal _{water}	PEC in water during emission episode	5.07E-06	[mg.L ⁻¹]	TGD Eq. 45
PEClocal _{sed}	PEC in sediment during emission episode	4.50E-02	[mg.kg ⁻¹ _{wwt}]	TGD Eq. 50

RELEASE VIA THE MANURE SPREADING

The PEClocal_{water} following the run-off after manure/slurry spreading onto soil are calculated as shown below: PIEC grs-ars- surface water N = PIEC grs-ars- groundwater N / DILUTIONrun-off. A dilution factor of 10 was considered according to the ESD n°14. Result for the worst case scenario (housing of veal calves) is shown in the following table.

PEC soil values considered to carry out the calculation (initial PEC soil) are presented in Table 18. It is worth noting that for surface water contamination via run-off, it was deemed more realistic to take the PEC value after only one application for grassland.

Symbol	Parameter	Value		Unit
Veal calves (Del	tamethrin)	Grassland	Arable land	
PEClocal _{water} Deltamethrin	PEC in water during emission episode	1.14E-08	8.54E-09	[mg.L ⁻¹]

2.8.4.2.2 Atmospheric compartment

Significant exposure of the environment via air is not expected.

Due to its low vapour pressure, deltamethrin is not expected to volatilise to air from plants and soil at significant levels, which was confirmed in a wind tunnel study. However, the calculated Henry's law constant is 1.252×10^{-3} Pa.m³.mole⁻¹, indicating that deltamethrin has a tendency to volatilise from water. If present in air, the data on indirect photo-oxidation indicate a rapid degradation when reacting with hydroxyl radicals (DT₅₀ reaction with OH-radicals = 16.4 hours).

2.8.4.2.3 Terrestrial compartment (soil and groundwater)

RELEASE VIA THE WASTE WATER

The concentrations in agricultural soil, following the spreading of contaminated STP sludge, are calculated according to the TGD equations considering the emission rates to wastewater (E_{ww}). A degradation of deltamethrin in soil (DT_{50} value of 48 days at 12°C) is taken into account. To estimate PECs in porewater for the relevant metabolite Br₂CA, a Koc value of 25.61 L.Kg⁻¹ and a DT_{50} in soil at 12°C of 5.6 days have been considered. Initial concentrations of Br₂CA in soil following application of sewage sludge to land were estimated on the worst-case assumption that the metabolite is formed in the sludge at a quantity equivalent to 100% of the parent (adjusted to take into account the molecular weights of the compounds, F=0.59).

According to the TGD, PECs groundwater were derived from the twa value of PECs soil over 180 days.

Symbol	Parameter	Value	Unit	Reference
E _{applicator,ww}	Emission from applicator to waste water	2.10E-05	Kg.d⁻¹	Output
PEC _{local soil}	PEC soil 30d	3.21E-05	[mg.kg ⁻¹ _{wwt}]	TGD Eq. 60
PEC local soil porewater Deltamethrin		1.96E-06	[ug] ⁻¹]	
PEC local soil porewater B _{r2CA}	PEC in porewater (agricultural.soil) 180d	1.79E-03	· [µg.L ⁻¹]	TGD Eq. 67

Table 19: PECs in Terrestrial compartment - Release via the STP - Emission from applicator

Table 20: PECs in Terrestrial compartment – Release via the STP - Emission from surface during cleaning phase

Symbol	Parameter	Value	Unit	Reference
E treated surface, ww	Emission from floor/treated to waste water during the cleaning step	1.70E-03	Kg.d⁻¹	Output
PEC _{local soil}	PEC soil 30d	2.61E-03	[mg.kg ⁻¹ _{wwt}]	TGD Eq. 60
PEC local soil porewater Deltamethrin	DEC	1.59E-04	[µg.L ⁻¹]	TGD Eq. 67
PEC local soil porewater B _{r2CA}	PEC in porewater (agricultural.soil) 180d	1.45E-01	[µy.∟]	160 Eq. 67

RELEASE VIA THE MANURE SPREADING

Soils are exposed SANITERPEN INSECTICIDE DK PRET A L'EMPLOI by manure used in agriculture. The concentrations in soil in the case of an immission standard for nitrogen and land application on arable land or grassland (PIECars-N_i / PIECgrs-N_i) are summarised in the table below for the worst case scenario, "housing of veal calves".

Table 21: Overview on the calculated PEC for the soil compartment (intial and twa over 180d) – Manure/slurry spreading

Symbol	Parameter	Value		Unit	Deferrer
Emission from floo	or/treated to manure	eated to manure Grassland-N Arable Land-		Unit	Reference
PIEC _{local soil} Deltamethrin	PEC soil initial	3.27E-03 8.17E-04*	6.13E-04	- [mg.kg ⁻¹ _{wwt}]	TGD Eq. 60
PIEC _{local soil} B _{r2CA}		1.93E-03 4.82E-04*	3.61E-04		
PIEC local soil 180d Deltamethrin		1.17E-03 2.92E-04*	2.19E-04	Ima ka ⁻¹ 1	
PEC local soil 180d B _{r2CA}	PEC in soil 180d	8.64E-05 2.16E-05*	1.62E-05	· [mg.kg ⁻¹ _{wwt}]	TGD Eq. 67

* For one application only on grassland

Application on soil of manure containing deltamethrin and Br_2CA can result in groundwater contamination. Therefore, PEC porewater values are derived based on PT 18 ESD calculations from concentrations of the active substance in soil based on the nitrogen immission standard for arable/grassland soil and leading to concentrations in porewater for soils. The time-weighted averaged PEC values (over 180 days) of deltamethrin and Br_2CA are used to define groundwater concentrations. The concentrations in porewater after application on arable land and grassland are summarised in the table below.

Table 22: Overview on the calculated PEC groundwater initial – Manure/slurry spreading

Veal calves (Deltamethrin)	4.55E-04 1.14E-04*	8.54E-05
Veal calves (Br ₂ CA)	3.39 8.46E-01*	6.35E-01

* For one application only on grassland

Table 23: Overview on the calculated PEC groundwater 180 days – Manure/slurry spreading

RELEASE VIA THE MANURE SPREADING	PEC local soil porewater Grassland [µg.L ⁻¹]	PEC _{local soil porewater} Arable Land [µg.L ⁻¹]
Veal calves (Deltamethrin)	1.63E-04	3.05E-05
Veal calves (Br ₂ CA)	1.52E-01	2.84E-02

2.8.4.2.4 Non-compartmental-specific exposure relevant to the food chain (secondary poisoning)

The product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is an insecticide applied only indoors.

Primary poisoning, i.e. the direct consumption of insecticide by non-target animals like birds, mammals or honeybees, is a topic only for outdoor uses. Consequently, risk of primary poisoning is not relevant and has not been assessed.

The concentration of deltamethrin in contaminated food is calculated in order to address the risk of secondary poisoning to top predators via the aquatic food chain (i.e. fish-eating birds and mammals). Biomagnification may also occur via the terrestrial food chain and the risk concerning secondary poisoning for worm-eating birds and mammals is also assessed.

RELEASE VIA THE WASTE WATER

The PEC oral, fish value and the PEC oral, earthworm are presented in the Tables below.

Table 24: Overview on the calculated local PECoral fish and local PECoral earthworm - Releas	e to waste
water - Emission from applicator	

Symbol	Parameter	Value	Unit	Reference
E treated surface, ww	Emission from floor/treated to waste water during the cleaning step	2.10E-05	Kg.d⁻¹	Output
PEC _{oral, predator}	Predicted Environmental Concentration in food (fish)	4.38E-05	[mg.kg ⁻¹]	TGD Eq. 76
PEC _{oral, predator}	Predicted Environmental Concentration in food (earthworm)	1.15E-04	[mg.kg ⁻¹]	TGD Eq. 81

Table 25: Overview on the calculated local PECoral _{fish} and local PECoral _{earthworm} – Release to waste water - Emission from surface cleaning phase

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Symbol	Parameter	Value	Unit	Reference
E treated surface, ww	Emission from floor/treated to waste water during the cleaning step	1.70E-03	Kg.d⁻¹	Output
PEC _{oral, predator}	Predicted Environmental Concentration in food (fish)	3.54E-03	[mg.kg ⁻¹]	TGD Eq. 76
PEC _{oral, predator}	Predicted Environmental Concentration in food (earthworm)	9.27E-05	[mg.kg ⁻¹]	TGD Eq. 81

RELEASE VIA THE MANURE SPREADING

The PEC _{oral,fish} value and the PEC _{oral,earthworm} are presented in the Tables below for the worst case scenario, housing of veal calves.

Table 26: Overview on the calculated local PECoral _{fish} and local PECoral _{earthworm} – Release via the manure spreading

RELEASE VIA THE MANURE SPREADING	Grassland	Arable Land
Veal calves		
PEC _{fish} [mg.kg ⁻¹]	7.97E-06	5.98E-06
PEC _{earthworm} [mg.kg ⁻¹]	9.46E-05	1.77E-05

2.8.5 Risk characterisation for the environment

Risk characterization for the environment is done quantitatively by comparing predicted environmental concentrations (PEC) and the concentrations below which effects on organism will not occur (PNEC) according to the Technical Guidance Document (TGD, 2003) and Emission Scenario Documents for PT18. The environmental risk characterization has been carried out for deltamethrin. For indoor uses of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI in empty animal houses and shelters, risks following indirect emissions via the STP and via manure applications on agricultural land are characterized for all compartments.

2.8.5.1 Professional Use of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI for treatment of pet's environment in rural areas - INDOOR application in empty animal houses and shelters

Calculations are based on the maximum amount applied per day as a worst-case scenario, as detailed in the environmental exposure assessment section. The scenario includes the emission of active substance to environmental compartments when applying the product and during cleaning events. According to the intended use, the main route of entry into the environment is assumed to be indirect to the aquatic compartment, via STP effluents, during the preparation/application phase and the cleaning phase.

To complete the assessment, a potential emission into the environment *via* application of manure (for example to cover horse manure application) on arable land and grassland have been taking into account considering a maximum of 3 insecticide applications per year. It should be noted that this scenario taken from the ESD n°14 "Emission scenario document for Stables and Manure Storage Systems" is considered as a very worst case approach as the emission calculations form this document cover industrial farming.

RELEASE VIA THE WASTE WATER

The table below summarizes the PEC/PNEC ratios for the aquatic compartment (including STP, surface water and sediment) and terrestrial compartment (including soil and groundwater), the threshold values for groundwater being < $0.1 \ \mu g.L^{-1}$.

Table 27: Risk characterization for indirect emissions (*via* the STP) - Indoor application in empty animal houses and shelters of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

	PEC	PEC/PNEC	Risks		
Emission from a	Emission from applicator				
STP	PNEC _{STEP microorganisms} =	= 3.00E-02 mg.L ⁻¹			
[mg.L ⁻¹]	1.01E-06	3.37E-05	Acceptable		
Surface water	PNEC _{surface water} = 0.70	ng L ⁻¹			
[mg.L ⁻¹]	6.25E-08	8.93E-02	Acceptable		
Sediment	PNEC _{sediment} = 6.20 µg	-1 •Kg _{wwt sediment}			
[mg.kg _{wwt} ⁻¹]	5.55E-04	8.95E-02	Acceptable		
Soil	PNEC _{soil} = 0.075 mg.k	9w soil			
[mg.kg _{wwt} -1]	3.21E-05	4.28E-04	Acceptable		
Threshold value = $0.1 \ \mu g.L^{-1}$					
Groundwater	Deltamethrin	< 0.1 µg.L ⁻¹	Accontable		
	BR ₂ CA	< 0.1 µg.L ⁻¹	Acceptable		
Secondary Pois.	PNEC _{oral mammal} 2.67 n	ng kg _{diet} ⁻¹			
Terrestrial food chain [mg kg _{diet} ⁻¹]	1.15E-04	4.31E-05	Acceptable		
Aquatic food chain [mg kg _{diet} ⁻¹]	4.38E-05	1.64E-05	Acceptable		
Emission from surface cleaning phase					
STP PNEC _{STEP microorganisms} = 3.00E-02 mg.L ⁻¹		= 3.00E-02 mg.L ⁻¹			
[mg.L ⁻¹]	8.18E-05	2.73E-03	Acceptable		
Surface water $PNEC_{surface water} = 0.70 \text{ ng L}^{-1}$		ng L ⁻¹			
[mg.L ⁻¹]	5.07E-06	7.24E+00	Unacceptable		
Sediment	PNEC _{sediment} = 6.20 μg	-1 -1 -1			
[mg.kg _{wwt} ⁻¹]	4.50E-02	7.26E+00	Unacceptable		
Soil	$PNEC_{soil} = 0.075 \text{ mg.kg}_{w soil}^{-1}$				

[mg.kg _{wwt} ⁻¹]	2.61E-03	3.48E-02	Acceptable
	Threshold value = $0.1 \ \mu g.L^{-1}$		
Groundwater	Deltamethrin	< 0.1 µg.L ⁻¹	Acceptable
	BR ₂ CA	> 0.1 µg.L ⁻¹	Unacceptable
Secondary Pois.	PNEC _{oral mammal} 2.67 mg kg _{die} ⁻¹		
Terrestrial food chain [mg kg _{diet} ⁻¹]	9.27E-05	3.47E-05	Acceptable
Aquatic food chain [mg kg _{diet} ⁻¹]	3.54E-03	1.33E-06	Acceptable

RELEASE VIA THE MANURE SPREADING

The tables below summarize the PEC/PNEC ratios for terrestrial compartment (including soil and groundwater), the threshold values for groundwater being < $0.1 \ \mu g.L^{-1}$.

Table 28: Risk characterization for terrestrial compartment - Indoor application in empty animal houses and shelters of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

<u>Release via the manure</u> spreading <u>Veal calves</u>	PEC	PEC/PNEC	Risks
Grassland			
Surface water	$PNEC_{surface water} = 0.70 \text{ ng L}^{-1}$		
[mg.L ⁻¹]	1.14E-08	1.63E-02	Acceptable
Soil	PNECsoil = 0.075 mg.kgw soil ⁻¹		
[mg.kg _{wwt} ⁻¹]	3.27E-03	4.36E-02	Acceptable
Threshold value = $0.1 \ \mu g.L^{-1}$			
Groundwater	Deltamethrin	< 0.1 µg.L ⁻¹	Acceptable
	BR₂CA	> 0.1 µg.L ⁻¹ (0.152 µg.L ⁻¹)	Acceptable
Secondary Pois.	PNEC $_{\text{oral mammal}} = 2.67 \text{ mg kg}_{\text{diet}}^{-1}$		
Terrestrial food chain [mg kg _{diet} ⁻¹]	9.46E-05 3.54E-05 Acceptable		Acceptable
Aquatic food chain [mg kg _{diet} ⁻¹]	7.97E-06	2.99E-06	Acceptable

Arable Land			
Surface water	$PNEC_{surface water} = 0.70 \text{ ng L}^{-1}$		
[mg.L ⁻¹]	8.54E-09	1.22E-02	Acceptable
Soil	PNECsoil = 0.075 mg.kgw soil ⁻¹		
[mg.kg _{wwt} -1]	6.13E-04	8.17E-03	Acceptable
	Threshold value = 0.1 µg.L ⁻¹		
Groundwater	Deltamethrin	< 0.1 µg.L ⁻¹	Acceptable
	BR ₂ CA	< 0.1 µg.L ⁻¹	Acceptable
Secondary Pois.	PNEC _{oral mammal =} 2.67 mg kg _{diet} $^{-1}$		
Terrestrial food chain [mg kg _{diet} ⁻¹]	1.77E-05 6.64E-06 Acceptable		Acceptable
Aquatic food chain [mg kg _{diet} ⁻¹]	5.98E-06 2.24E-06 Acceptable		Acceptable

2.8.6 Conclusions for the environment

The product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is applied by professionals only indoors in empty animal houses and shelters.

Targeted animal houses and shelters are small houses which are used in rural areas by farmers, hunters, agricultural workers, or houses and shelters used for small farming (dog farming, ornamental chicken farming), or by small associations working for the protection of the animals.

The risk assessment has been carried out for the **application phase** and for the subsequent **cleaning phase**, as it is indicated by the applicant that the surface must be cleaned before a new application (but not between applications). Consequently a release to wastewater from floor and treated surfaces was taking into account. To complete the assessment, a potential emission into the environment *via* a potential **application of manure** (for example to cover horse manure application) on arable land and grassland has been taking into account. It should be noted that this scenario taken from the ESD n°14 "Emission scenario document for Stables and Manure Storage Systems" is considered as a very worst case approach as the emission calculations form this document cover industrial farming.

Risks are acceptable for the application phases from the applicator for all environmental compartments.

Concerning the **cleaning phase**, risks are acceptable for STP and soil compartments. On the other hand, risks are <u>unacceptable for surface water</u>, sediment and groundwater, taking into account the intended dose rate and a cleaning efficiency of 50% with water.

Considering a potential **application of manure** (for example to cover horse manure application) on arable land and grassland, risks are acceptable considering that the proposed scenario is a very worst case approach.

In conclusion, risks are acceptable taking into account the intended dose rate and with respect to the operating instructions proposed by the applicant and presented below leading to **no releases at all to the STP via the wastewater during the cleaning phase of the treated surfaces**:

Risk mitigation measures linked to environmental risk assessment

Before using SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

- Floor and walls, including previously treated surfaces, shall be mechanically cleaned from dusts using vacuum system or brush and dusts shall be collected and treated as a waste.
- Surface to be treated shall be determined and calculated regarding specific cases (pets to be protected, insects to be controlled). In all case it should be limited to 350 m².

After application of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

- After application the sprayer equipment shall be rinsed 3 times with water and rinsing water shall be collected and treated as a waste.
- The product must be kept inside the original packaging.
- In any case, never throw rinsing water or the product in the waste water system or surface water.

After application treated surfaces shall not be cleaned (dry cleaning will take place before the next application).

Disposal considerations

- Dispose of unused product, its packaging and all other waste in accordance with local regulations.
- Do not discharge unused product into water courses, into pipes (sink, toilets...) nor down the drains.

2.9 Risk assessment for companion and ornamental animals

The product is applied only indoors in animal houses and shelters to protect animals from target pests. Only animals not intented for human consumption are considered.

No data were submitted by the applicant to assess the risk for animals living in treated premises and few data in the CAR can be used to address this issue. Moreover, existing scenarios are designed to assess the exposure of animals intended for human consumption (assessment of dietary risk) and the exposure of wild animals (primary and secondary poisoning). No scenario and no guidance is available to assess the risks for companion animals and ornamental animals (pets).

However, some elements were considered to estimate the risk for these animals, based on guidance on the assessment of exposure of farm animals intended for human consumption and on extrapolation from method used for human risk assessment.

Exposure by ingestion (animals licking treated surfaces, grooming, ingestion of dead insects, ingestion of contaminated feed and drinks) and by dermal contact is considered.

2.9.1 Toxicological reference values

Deltamethrin is used in veterinary medicines authorised in EU. National and European databases were consulted. Roughly 10-12 medicines based on deltamethrin are authorised for dermal application. The doses are the following:

- 100 to 225 mg/animal for cattle;
- 75 mg/animal for adult sheeps;
- 25 mg/animal for young sheeps below 10 kg;
- 750 mg to 1 g in collars for dogs.

Moreover, on the basis of the studies assessed in the CAR, the following reference values were determined:

<u>Dogs</u>: for human, an AEL of 0.0075 mg/kg bw/day was derived based on the NOAEL (1 mg/kg bw/day) obtained in the 1-year dog study, taking into account an oral absorption of 75% and a safety factor of 100. A specific AEL for dogs can be derived considering an intra-species assessment factor of 1, leading to an AEL for dog of 0.075 mg/kg bw/d.

- <u>Rabbits</u>: a NOEL_{maternal} of 10 mg/kg bw/d has been observed in a developmental toxicity study in rabbits. This value can be used to derive a specific AEL for this species considering an intra-species assessment factor of 1 and an oral absorption of 75%. An AEL value of 0.75 mg/kg bw/d is calculated for rabbits.
- <u>Birds</u>: a PNEC of 15 mg/kg diet was derived from a study on *Colinus virginianus*, with a safety factor if 30.

2.9.2 Exposure assessment for horses, rabbits and birds

Exposure of animals has been evaluated on the basis of the draft "Guidance on estimating livestock exposure to active substances used in biocidal product" (version CA-Dec10-Doc.6.2.b). The calculator developed by BfR and available online has been used. The application rate of 10 mg deltamethrin/m² was used.

Step 1: Screening scenario:

- Hypothesis: the entire amount of biocidal product applied is taken up by animals regardless of the route of exposure;
- Default values are taken from the draft guidance document;
- Screening scenario: surface treatment of animal housing (floor and wall of stable without partitions);
- Animal are present in the premises during treatment.

Step 2 - realistic worst case:

Exposure can be refined considering each relevant route of exposure (oral-animal licking surfaces, oralingestion of dead insects (for poultry), dermal-rubbing against surfaces). Moreover, according to the applicant, animals are not present during application and all feed and toughs are taken out.

No exposure or negligible exposure is expected from licking surfaces, for horses and rabbits. Licking is also not relevant for birds.

Input values for realistic worst case scenario are reported in Table 29.

Results are reported in Table 30.

Factor	Value (source: draft guidance document)
Oral exposure through ingestion of flies	
Fly consumption	10 flies/day (default value)
Consumption of B.P. (spray deposit) by flies	3.5 mg product/d
Concentration of a.s. in BP applied on surfaces	0.2 g/L
a.s. consumption by flies	0.0007 mg/fly/d
Feed intake	Broiler: 0.12 kg dry matter/d (default value)
	Turkey: 0.35 kg dry matter/d (default value)
Dermal exposure through rubbing against sur	faces
Emission factor for spraying	1 (worst case)
Body surface area in contact with surface	Horse: 1.62 m ² (default value)
Body weight	Broiler: 1.7 kg (default value)
	Turkey: 7 kg (default value)
	Haraa, 400 kg (dafault value)
	Horse: 400 kg (default value)

Table 29: Input values for realistic worst case scenario

Table 30: Exposure of animals for surface treatment of animal housing (floor and wall of stable without
partitions) – screening scenario and realistic worst case scenario

		Exposure (mg a.s./kg bw/day)		
Animal Spec	ies			Dermal - Rubbing against surfaces
Broilers		Nd	0.0412	Nd
Broilers	free range, litter floor	0.4706	Nd	Nd
Broilers	parent broilers, free range (grating floor)	0.5042	Nd	Nd
Broilers	parent broilers in rearing, free range (grating floor)	0.4902	Nd	Nd
Laying hen		Nd	0.0368	Nd
Laying hen	battery	0.2757	Nd	Nd
Laying hen	free range (litter floor)	1.0684	Nd	Nd
Laying hen	free range (grating floor)	0.4795	Nd	Nd
Turkey		Nd	0.0100	Nd
Horse		Nd	Nd	0.04050
Rabbit		0.6720	Nd	Nd

Nd: not determined

It should be noted that these results are based on parameters relevant for farming of animals intended for human comsumption. They may be not representative of the conditions in companion animals breeding installations.

Estimates for the risk characterisation are presented in Table 31.

	Exposure (mg/kg bw/d)	AEL or PNEC	Risk ratio
Broiler	0,0041	15 mg/kg diet Corresponding to 1.8 mg/d or 1.1 mg/kg bw/d	0,004
Turkey	0,0010	15 mg/kg diet Corresponding to 5.25 mg/d or 0.75 mg/kg bw/d	0,001
Rabbits	0.6720	0.75 mg/kg bw/d	0.9
Horses	0.0405	No AEL	Not determined

	Exposure is higher than human AEL and lower than dog AEL. Exposure is also lower than medicinal doses for veterinary medicines for cattle and sheeps, within the same order of magnitude.
--	--

2.9.3 Exposure assessment and risk characterisation for dogs

Exposure by ingestion (animals licking treated surfaces) was assessed by reverse scenario, in order to determine the maximum treated surface area that a dog has to lick to reach the AEL.

The following parameters have been considered:

- Concentration of active substance in the diluted product (applied on surfaces): 0.02%;
- Application rate: 50 mL product/m²;
- Density value: 0.947;
- Oral absorption value: 75%;
- Transfer coefficient : 100% (from TNsG 2008); _
- Body weight of a puppy: 5 kg;
- $AEL_{dog} = 0.075 \text{ mg/kg bw/d}$

A maximum surface area of 0.05 m^2 has to be licked by a puppy to reach the AEL. This value can be considered realistic, leading to an unacceptable risk.

2.9.4 Pharmacovigilance data

Veterinary pharmacovigilance available data in France cannot be used to determine hypersensitivity of a species to deltamethrin.

However, it could be noted that the most adverse effects have been reported for cats and equines (29 cases for cats and 17 for horses since 2002). The main clinical symptoms are:

- For cats: hypersalivation, vomiting, ataxia, tremors, and (likely related to overdosing) death;
- For horses: hyperthermia, itch and agitation.

Furthermore, bibliographic data reveal that cats are particularly sensitive to pyrethroids, due to a lack of glucuronide conjugation enzymes and other not yet elucidated reason(s)^{12,13,14}. Although sensitivity to permethrin is more documented, without any further data, it is recommended that the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is not used to treat premises were cats are housed, as well as other species that may display a particular sensitivity to deltamethrin.

¹² Gfeller, R.G., Messonnier, S.P., 2004. Handbook of Small Animal Toxicology and Poisonings, second ed. Mosby, St. Louis, MO, USA.

¹³ Anadón A., Martínez-Larrañaga M.R., Martínez M.A., 2008. Use and abuse of pyrethrins and synthetic pyrethroids in veterinary medicine. The Veterinary Journal 182 (2009) 7–20. ¹⁴ Beugnet F., Franc M., 2012. Insecticide and acaricide molecules and/or combinations to prevent pet infestation by ectoparasites.

Trends in Parasitology, July 2012, Vol. 28, No. 7.

2.9.5 Conclusions for companion and ornamental animals

In conclusion, secondary exposure to companion animals with treated surfaces cannot be precisely assessed. However, considering the presented estimations, this exposure cannot be considered negligible. In order to limit this exposure, the following instructions and risk mitigation measures must be respected.

Risk mitigation measures linked to risk assessment for animals

- Do not use in premises where cats or other animals with particular sensitivity to pyrethrinoïds are housed.
- Alternate products containing active substances with different mode of action than pyrethroïds.
- Apply only during a fallowing period in animal shelters/housings (empty premises).
- Do not apply on surfaces likely to be licked by animals.
- Prior treatment, remove the bowls, feeders, drinkers and textiles from animal sleeping areas. If they cannot be removed, empty and cover them with a plastic sheet for the duration of the treatment.
- Wait complete drying of the treated surfaces after the end of the treatment, before allowing animals to re-enter.

2.10 Measures to protect man, animals and the environment

See Summary of Product Characteristics (SPC)

3 PROPOSAL FOR DECISION

Conclusions of efficacy and risk assessment

Risk assessment for Physico-chemical properties

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is a ready-to-use product containing deltamethrin (0.02 % (w/w)) spray application. It is not highly flammable, not auto-flammable, not explosive and does not have oxidizing properties.

The product is stable for 14 days at 54°C and 2 years at ambient temperature.

The product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI is compatible with PET packagings

Summary of efficacy assessment

The efficacy level of the product SANITERPEN INSECTICIDE DK PRET A L'EMPLOI (0.02 % w/w deltamethrin) is satisfactory for the uses proposed in Annex 0b.

Summary of risks characterisation of the product for human health

Risks related to the use of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI by professionals are considered acceptable during spray application. Risk related to secondary exposure is also considered acceptable.

Summary of risks characterisation of the product for consumer

Regarding the intended used, exposure of livestock and human exposure via food of animal origin are not expected. Dietary risk assessment is not required.

Summary of risks characterisation of the product for the environment

Risks are acceptable for the application phases from the applicator for all environmental compartments.

Concerning the cleaning phase, risks are acceptable for STP and soil compartments. On the other hand, risks are <u>unacceptable for surface water</u>, sediment and groundwater, taking into account the intended dose rate and a cleaning efficiency of 50% with water.

Considering a potential application of manure (for example to cover horse manure application) on arable land and grassland, risks are acceptable considering that the proposed scenario is a very worst case approach.

In conclusion, risks are acceptable taking into account the intended dose rate and with respect to the operating instructions proposed by the applicant and presented below leading to no releases at all to the STP via the wastewater during the cleaning phase of the treated surfaces:

Summary of risks characterisation of the product for companion and ornamental animals

Secondary exposure to companion animals with treated surfaces cannot be precisely assessed. However, considering the presented estimations, this exposure cannot be considered negligible. In order to limit this exposure, the following instructions and risk mitigation measures must be respected.

Risk mitigation measures and conditions of use

The uses proposed for authorisation are detailed in Annex 0B

Conditions of use linked to efficacy assessment

To ensure a satisfactory level of efficacy and avoid the development of resistance in susceptible insect populations, the following recommendations have to be implemented:

- Always read the label or leaflet before use and respect follow all the instructions provided.

- Take into account the life cycle and characteristics of target insects to adapt treatments. In particular, target the most susceptible stage of the pest, timing of applications and areas to be treated.
- Adopt integrated pest management methods such as the combination of chemical, physical control methods and other public health measures, taking into account local specificities (climatic conditions, target species, conditions of use, etc).
- Alternate products containing active substances with different mode of action, (to remove resistant individuals from the population).
- The users should inform if the treatment is ineffective and report straightforward to the registration holder.

Risk mitigation measures for human health and environment

- Shake before use.
- Apply strict hygiene measures: do not eat, drink or smoke during handling of the product and wash hands after use of the product;
- Keep out of the reach of children.
- Do not use on surfaces likely to be in direct contact with animals intended for human consumption, food, feed or drinks.
- Floor and walls, including previously treated surfaces, shall be mechanically cleaned from dusts using vacuum system or brush and dusts shall be collected and treated as a waste.
- Surface to be treated shall be determined and calculated regarding specific cases (pets to be protected, insects to be controlled). In all case it should be limited to 350 m².
- The product must be kept inside the original packaging.
- In any case, never throw rinsing water or the product in the waste water system or surface water.
- After application treated surfaces shall not be cleaned (dry cleaning will take place before the next application).

Risk mitigation measures for animals

- Do not use in premises where cats or other animals with particular sensitivity to pyrethrinoïds are housed.
- Alternate products containing active substances with different mode of action than pyrethroïds.
- Apply only during a fallowing period in animal shelters/housings (empty premises).
- Do not apply on surfaces likely to be licked by animals.
- Prior treatment, remove the bowls, feeders, drinkers and textiles from animal sleeping areas. If they cannot be removed, empty and cover them with a plastic sheet for the duration of the treatment.
- Wait complete drying of the treated surfaces after the end of the treatment, before allowing animals to re-enter.

Emergency (not assessed by Anses)

- **Inhalation:** Remove victim to fresh air and keep at rest in a position comfortable for breathing. Get medical attention if symptoms occur, show this container or label.
- **Skin contact**: Remove contaminated clothing and shoes. Wash contaminated skin with soap and water. Get medical attention if symptoms occur.
- **Eye contact**: Immediately flush with plenty of water, occasionally lifting the upper and lower eyelids. Check for and remove any contact lenses. Continue to rinse with warm water for at least 10 minutes. Get medical attention if irritation or vision impairment occurs.
- **Ingestion:** Wash out mouth with water. Get medical attention if symptoms occur, show this container or label.Do not drink or induce vomiting in case of consciousness alteration.
- **Note to physician:** Treat symptomatically. Contact poison treatment specialist immediately if large quantities have been ingested or inhaled.

Disposal

- Dispose of unused product, its packaging and all other waste in accordance with local regulations.

- Do not discharge unused product into water courses, into pipes (sink, toilets...) nor down the drains.

Recommendations to be taken into account by the authorisation holder

The authorization holder should report any observed resistance incidents to the Competent Authorities (CA) or other appointed bodies involved in resistance management.

Information required post-authorisation

Required information linked to efficacy assessment

- Suitable information (as semi-field or field tests) demonstrating the efficacy of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI against target organisms mosquitoes (*Culex pipiens* and *Aedes aegypti*) and fleas (*Pulicidae*), will need to be provided in support of the authorisation, within one year.
- Establish a baseline and monitor levels of effectiveness on populations in key areas (at least one survey per year) in order to detect any significant changes in susceptibility to active substance. Information from resistance monitoring programs allows early detection of problems and gives information for correct decision making.

4 APPENDICES

Annex 0a: Practical use claimed by the applicant

 Table 32: Practical use claimed by the applicant

Name of the product and type of formulation (gel, paste, spray, dust, powder, fumigation)	Target organisms (common species and genus) and development stages (eggs, larvae, nymph, adults)*	User category (professional/non professional)*	Application aim	Area of use (indoor, outdoor, and field of use)	Method of application	Application rate (expressed in g/m ³ , g/m ² , ml/m ²) Maximum and minimum dosage (if appropriate)	Mode of action including time delay (kill, knockdown)	Time delay of residual efficacy if indirect or surface treatment (hours, days, weeks and months)	Time delay for human , food and animals reentrance after treatment (if appropriate)	Frequency and duration of application	Dosage and applications requirements (exposure time, ventilation, temperature,)	Package Individual packaging (yes/no)**	Primary packaging *** : type : bulk, individual wrapping/ nature: bucket, bottle, sachet/ material: paper, polyethylene/ sizes	Secondary packaging	Accepted and authorized by the RMS (yes/no)
SANITERPEN INSECTICIDE DK PRET A L'EMPLOIFormulation: ready-for-use liquid (AL).	Dermanyssidae : Dermanyssus gallinae (Poultry red mite) Eggs, larvae and adults	Professional	To kill insect to protect health of animals and humans	To be used indoors only in empty Animal houses/ shelters	Spra ying	50mL/m2 350m2 max. per application	Knockdo wn + Residual efficacy of 3 months.	Residual efficacy of three months.	Animals can re- enter the treated areas when the surfaces are dried. 24hours per default.	One applicat ion every 3 months Max 3 applicat ions per year.	Applicati on shall be made only with the original packagin g.	Yes	750mL PET bottle with a hand sprayer	-	

Culicidae: Culex sp, Aedes sp. (Mosquitoes) Adults	Professional	To kill insect to protect health of animals and humans	To be used indoors only in empty Animal houses/ shelters	Spra ying	50mL/m2 350m2 max. per application	Knockdo wn + Residual efficacy of 3 months.	Residual efficacy three months.	of	Animals can re- enter the treated areas when the surfaces are dried. 24hours per default.	One applicat ion every 3 months Max 3 applicat ions per year.	Applicati	Yes	750mL PET bottle with a hand sprayer	-	
Muscidae: <i>Musca domestica</i> (House fly) Adults	Professional	To kill insect to protect health of animals and humans	To be used indoors only in empty Animal houses/ shelters	Spra ying	50mL/m2 350m2 max. per application	Knockdo wn + Residual efficacy of 3 months.	Residual efficacy three months.	of	Animals can re- enter the treated areas when the surfaces are dried. 24hours per default.	One applicat ion every 3 months Max 3 applicat ions per year.	Applicati on shall be made only with the original packagin g.	Yes	750mL PET bottle with a hand sprayer	-	
Pulicidae: (Fleas) Larvae and adults	Professional	To kill insect to protect health of animals and humans	To be used indoors only in empty Animal houses/ shelters	Spra ying	50mL/m2 350m2 max. per application	Knockdo wn + Residual efficacy of 3 months.	Residual efficacy three months.	of	Animals can re- enter the treated areas when the surfaces are dried. 24hours per default.	One applicat ion every 3 months Max 3 applicat ions per year.	Applicati on shall be made only with the original packagin g.	Yes	750mL PET bottle with a hand sprayer	-	

Annex 0b: Proposed uses for authorisation

This table reflects the results of the risk assessment. In case of differences between the uses suggested by Anses to be authorised and the uses contained in the decision taken by the French ministry, only the original and signed decision has a legal value.

Table 33: Proposed uses for authorisation

Name of the product	User category (professional/non professional)*	Type of formulation (grains, powder, paste, block)*	Target organism (rat, mice…)*	Dosage claimed expressed in g/bait point, for high and low infestation (if appropriate)	Area of use (sewers, in and around buildings, indoor only, open areas, waste dumps,)*	Methods of application of the bait (ex: pre-filled secured bait box)	Primary packaging: bulk, individual wrapping	Authorizatio n
SANITERPEN INSECTICIDE PRET A L'EMPLOI Formulation: ready-for-use liquid (AL).	Professional uses	Ready to use bottle with a hand sprayer	House flies, adults (<i>Musca</i> <i>domestica</i>) Mosquitoes, adults <i>genus Culex*,</i> <i>genus Aedes*</i> Fleas (<i>Pulicidae</i>)*, adults	Application rate : 50 mL/m ² Corresponding to 100 % v/v The product is ready to use	To be used indoors only in empty Animal houses/shelters	One application every 3 months. Max 3 applications per year on porous and non-porous surfaces	750 ml PET bottle with a hand sprayer	Yes

In the absence of sufficient supporting data on mosquitoes (genus Culex and Aedes) and fleas (Pulicidae), suitable information (as semi-field or field tests) demonstrating the efficacy of SANITERPEN INSECTICIDE DK PRET A L'EMPLOI against these target organisms, will need to be provided in support of the authorisation, within one year.

Annex 1: Summary of product characteristics

See separated file.

Annex 2: List of studies reviewed

List of <u>new data¹⁵</u> submitted in support of the evaluation of the active substance

List of <u>new data</u> submitted in support of the evaluation of the biocidal product

Section No Refere		Author	Year	Title	Owner of data		Letter of access		Data protection claimed		ential es for lation
Doc IIIB						Yes	No	Yes	No	Yes	No
B3.1.1 B3.1.2 B3.2 B3.4.1.1 B3.7	B.3.1	C. Da Costa, B. Demangel	2010	Accelerated storage procedure for 14 days at $54 \pm 2^{\circ}$ C and physico-chemical tests before and after storage on SANITERPEN INSECTICIDE EFFET CHOC, Défitraces, Report No.09-901011-006							
B3.1.1 B3.1.2 B3.2 B3.4.1.2 B3.7	B.3.2	B. Demangel	2013a	Physico-chemical tests and chemical stability after a storage procedure for 2 years at $20 \pm 2^{\circ}$ C on SANITERPEN INSECTICIDE EFFET CHOC, Défitraces, Report No.09-901011-007	Action Pin		x	x			
B3.1.3	B.3.3	B. Demangel	2013b	SANITERPEN INSECTICIDE DK EXTRA PRÊT A L'EMPLOI, Défitraces,Study No.13-901011-015	Action Pin		x	x			
B3.3 B3.4.1.3 B3.5.7 B3.8 B3.9 B4.6	B.3.4	Da Costa	2009a	Physico chemical tests on SANITERPEN INSECTICIDE EFFET CHOC, Défitraces, Report No.09-901011-005	Action Pin		x	x			

¹⁵ Data which have not been already submitted for the purpose of the Annex I inclusion.

Section No	Refere nce No	Author	Year	Title	Owner of data	Letter of access	Data protectio claime	Essential studies for evaluation
B3.5.6	B.3.5	N. Buton	2010	Prestations pour mesures granulométriques sur aérosols, Malvern Instruments SARL, Report 12/07/2010	Action Pin	x	x	
B3.5.6	B 3 6	B. Demangel	2013c	Analytical certificate SANITERPEN INSECTICIDE DK PRÊT A L'EMPLOI, batch 27214-B1/95001, Défitraces	Action Pin	x	x	
B4.1 B4.13 B4.17.1	B 4 1	Da Costa	2009b	Physico chemical tests on SANITERPEN INSECTICIDE DK, Défitraces, Report No.09-901011-001	Action Pin	x	x	
B5.1	B5.1/01	H. Ricau	2010a	Validation of the analytical method for the determination of deltamethrin in SANITERPEN INSECTICIDE EFFET CHOC, in compliance with SANCO/3030/99 rev.4 EU, Défitraces, Report No.09-901011-008	Action Pin	x	x	
B5.1	B5.1/01	H. Ricau	2010b	Validation of the analytical method for the determination of deltamethrin in SANITERPEN INSECTICIDE DK, in compliance with SANCO/3030/99 rev.4 EU, Défitraces, Report No.09-901011-004	Action Pin	x	x	
B6.5	B6.5	N. Huguet	2013	Deltamethrin mode of action.	Action Pin	X	X	Х
B6.7	B6.7/0 1	B. Serrano	2013	Laboratory masurement of the effectiveness of an insecticide speciality intended for the destruction of insects in farm buildings, animal accomodatin, transport and farming equipment - SANITERPEN INSECTICIDE DK Prêt à l'emploi. Laboratoire TEC. Assay No.1558c/112, 2013/04.	Action Pin	x	x	x
B6.7	B6.7/0 2	B. Serrano	2011	Mesure en laboratoire de l'efficacité d'une spécialité insecticide vis-à-vis des mouches. Laboratoire TEC. Assay	Action Pin	x	x	x

Section No Reference No				Title	Owner of data	Letter of access		Data protection claimed		studi	ential les for uation
				No.1296b/1008. 2011/02.							
B6.7	B6.7/0 3	B. Serrano	2014	Laboratory measurement of the effectiveness of an insecticide speciality intended for the control of insects in farm buildings, animal accomodations, transport and farming equipment - Efficacy against flies <i>Musca domestica</i> . Laboratoires TEC, November 2014, Assay N° 1826a/0914	Action Pin		x	x		x	
B6.7	B6.7/0 3	B. Serrano	2012	Field testing of an insecticide speciality intended to control flies in breeding premises. Laboratoires TEC, November 2014, Assay N° 1826a/0914	Action Pin		x	x		x	
B6.7	B6.7/0 4	B. Serrano	2014	Simulated use trial of the efficacy of insecticide products intended to control insects in breeding or food storage premises - Trial against fleas and mosquitoes (Aedes sp. +Culex sp.). Laboratoire TEC, November 2014, assay N°1826b/0914	Action Pin		x	x		x	
B6.8	B6.8/0 1	N. Liu and X. Yue	2000	Insecticide resistance and cross-resistance in the house fly (Diptera: Muscidae). Journal of Economic Entomology, 93 (4): 1269-1275.	Public		х		х	x	
B6.8	B6.8/0 2	Y.L. Konan, A.A. Koffi, J.M.C. Doannio and F. Darriet	2003	Résistance de <i>Culex quinquefasciatus</i> (Say, 1823) à la deltaméthrine et l'utilisation de la moustiquaire imprégnée en milieu urbain de Bouaké, Côte d'Ivoire. Bulletin de la Société de Pathologie Exotique, 92 (2): 128-129.	Public		x		х	x	
B6.8	B6.8/0 3	H.F. Tahir, A. Butt and S.Y. Khan	2009	Response of <i>Culex quinquefasciatus</i> to deltamethrin in Lahore district. Journal of Parasitology and Vector Biology, 1 (3): 19-24.	Public		x		x	x	
B6.8	B6.8/0	R.L. Bossard, N.C.	1998	Review of insecticide resistance in cat fleas	Public		Х		Х	Х	

Section No	Refere nce No	Author	Year	Title	Owner of data	Lette acce		prote	ata ection med	studi	ential es for lation
	4	Hinkle and M.K. Rust		(Siphonaptera: Pulicidae). Journal of Medical Entomology, 35 (4): 415-422.							
B6.8	B6.8/0 5	M. Marangi, M.A. Cafiero, G. Capelli, A. Camarda, O.A.E. Sparagano and A. Giangaspero	2009	Evaluation of the poultry red mite, Dermanyssus gallinae (Acari: Dermanyssidae) susceptbility to some acaricides in field populations from Italy. Experimental and Applied Acarology, 48: 11-18.	Public		x		х	x	
B6.8	B6.8/0 6	F. Beugnet, C. Chauve M. Gauthey and L. Beert	1997	Resistance of the red poultry mite to pyrethroids in France. Veterinary Record, 140: 577-579.	Public		x		x	x	
B8.1	B8.1	F. Richeux	2009a	SANITERPEN INSECTICIDE Effet choc: Skin irritation test in the rabbit. Phycher Bio Développement, Report number IC-OCDE-PH-09/0152	Action Pin		х	x		x	
B8.2	B8.2	F. Richeux	2009b	SANITERPEN INSECTICIDE effet choc: Eye irritation test in the rabbit. Phycher Bio Développement, Report number IO-OCDE-PH-09/0152	Action Pin		х	x		x	
B8.3	B8.3/0 1	F. Richeux	2009c	SANITERPEN INSECTICIDE effet choc, Local Lymph Node Assay in the mouse. Phycher Bio Développement, Report number LLNA-PH-09/0152	Action Pin		х	x		x	
B8.3	B8.3/0 2	F. Richeux	2009d	SANITERPEN INSECTICIDE effet choc: Acute oral toxicity in the rat – Acute toxic class method. Phycher Bio Développement, Report number TAO423-PH-09/152	Action Pin		х	x		x	
B8.5. 3	B8.5.3	F. Richeux	2009e	SANITERPEN INSECTICIDE effet choc: Acute dermal toxicity in the rat. Phycher Bio Développement, Report number TAD-PH-09/0152	Action Pin		х	x		x	
B10.1 B10.2	B10.1		2011	Deltamethrin (PT18) Assessment report, Finalised in the Standing Committee on	Public		X		x		

Section No	Refere nce No	Author	Year	Title	Owner of data	er of Letter of access		Data protection claimed		Essenti studies f evaluatio	
B10.4				Biocidal Products at its meeting on 6 May 2011 in view of its inclusion in Annex I to Directive 98/8/EC							
B10.4	B10.4	Ambrosi Scientific Consulting	2013	SANITERPEN DK Prêt à l'emploi EPI Suite Results For CAS 52918-63-5 (Deltamethrin)	No Public		x	x			
IIB	IIB 3.3	EUSES		Full report EUSES calculations	Action Pin		Х	Х			

Annex 3: Analytical methods residues – active substance

Deltamethrin

Date: 31/03/2015

Table 34: Matrix, action levels, relevant residue and reference

matrix	limit	relevant residue	reference or comment
	IIIIII	Televant Tesidue	
plant products	-	deltamethrin	No exposure expected
food of animal origin	0.02 mg/kg	deltamethrin	Confirmatory method is
-			required at AS reapproval
soil	0.1 µg/kg	deltamethrin	Confirmatory method is
			required at AS reapproval
drinking water	3 ng/L	deltamethrin	Confirmatory method is
-	-		required at AS reapproval
surface water	3 ng/L	deltamethrin	Confirmatory method is
	5		required at AS reapproval
air	2.25 μg/m ³	deltamethrin	-
body fluids /	20 ng/L	deltamethrin	Confirmatory method is
tissues	J. J		required at AS reapproval

Matrix	Method	Limit of quantification	Reference
Soil	LC-MS/MS	0.1 µg/kg***	Brumhard, B. (2005a)
Air	GC-ECD	0.27 μg/m ³	Class, T. (2001a)
	GC-ECD	3 ng/L***	Class, T. (2001b)
Water	LC-MS/MS	5 ng/L*	Brumhard, B. (2005b)
	GC-ECD	50 ng/L*	Martens, R. (1999)
Bland	GC-MS multi- method for 20 ng/L*** pyrethroids		Ramesh, A. & Ravi, P.E. (2004)
Blood	GC-MS	200 µg/L**	Frenzel, T. et al (1998) Frenzel, T. et al (2000) Brennecke, R. (1998)
Muscle, fat, liver/kidney, eggs	GC-ECD	0.02 mg/kg***	Martens, R. (2000)
Milk	GC-ECD	0.02 mg/L***	Martens, R. (2000)
	GC-ECD	0.02 mg/kg for rice, flour, bread, meat, candy, butter, banana cream pie and lettuce	Silvoy, J.J. (1993a)
Plants	LC-MS/MS	 0.02 mg/kg for edible material for barley, broccoli, corn, melon, lettuce, olive, pepper, sugar beet, tobacco, tomato, wheat and zucchini 0.05 mg/kg for non-edible materials for barley, broccoli, corn, melon, lettuce, olive, pepper, sugar beet, tobacco, tomato, wheat and zucchini 	Zimmer D. & Philipowski C. (2004)

* The LOQ is not low enough to cover the concentration having an effect on the most sensitive aquatic organisms (NOEC: 4.8 ng/L, from mesocosm study; see Doc II-A section 4).
 ** The LOQ (200 μg/l) is not in compliance with the general requirement in Regulation (EU) No 528/2012 (i.e. 50

μg/l). *** Confirmatory methods is required to update this dossier

Annex 4 : Toxicology and metabolism –active substance

Threshold Limits and other Values for Human Health Risk Assessment

Date: 19/02/2015

Summary			
	Value	Study	SF
AEL long-term	0.0075 mg/kg bw/d	13-week dog study	100
AEL medium-term	0.0075 mg/kg bw/d	13-week and 1- year dog studies	100
AEL acute	0.0075 mg/kg bw/d	1-year dog study	100
Inhalative absorption		100%	
Oral absorption		75%	
Dermal absorption		2%	
Classification			
with regard to (according to the criteri	toxicological data a in Reg. 1272/2008)	Acute tox. 3* - H301 Acute tox. 3* - H331	
		No specific limit concentrations	

Annex 5 : Toxicology – biocidal product

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

Date:	19/02/2015
Date.	13/02/2013

General information

Formulation Type Active substance(s) (incl. content) Category RTU 0.02% w/w

Acute toxicity, irritancy and skin sensitisation	n of the preparation (Annex IIIB, point 6.1, 6.2, 6.3)
Rat LD50 oral (OECD 420)	>2000mg/kg bw/d
Rat LD50 dermal (OECD 402)	>2000mg/kg bw/d
Rat LC50 inhalation (OECD 403)	None
Skin irritation (OECD 404)	Not irritant for skin
Eye irritation (OECD 405)	Irritant for eyes
Skin sensitisation (OECD 429; LLNA)	Not skin sensitizer
Additional toxicological information (e.g. Anr	nex IIIB, point 6.5, 6.7)
Short-term toxicity studies	none
Toxicological data on active substance(s)	none
(not tested with the preparation)	
	none
Toxicological data on non-active substance(s)	none

Furthe	r toxicological	Information
i uitite	i lonicologica	mormation

(not tested with the preparation)

Classification and labelling proposed for the preparation with regard to toxicological properties (Annex IIIB, point 9)	
Regulation 1272/2008/EC	Eye Irrit.2 – H319

none none

Product Assessment Report – SANITERPEN INSECTICIDE DK PRET A L'EMPLOI -Deltamethrin

Annex 6 : Safety for professional operators

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

Date: 19/02/2015

Exposure assessment

Exposure scenarios for intended uses (Annex IIIB, point 6.6)

Primary exposure of professionals

	Component CAS		Actual Dermal Total [mg/kg/d]	Inhalation Exposure [mg/m³]	Oral (non respirable) exposure [mg/kg/d]	Model
Mixing / Loading and application						
Tier 1	Deltamethrine	52918-63-5	1.84 x 10 ⁻³	1.02 x 10 ⁻⁴	4.73 x 10 ⁻⁵	ConsExpo

Risk assessment

Component	CAS	AEL [mg/kg/d]	Absorp inhalati on	tion [%] dermal	Total syst exposure [mg/kg bw/d]	% AEL	Risk	
Mixing / Loading and application								
Deltamethrine	52918-63-5	0.0075	100	10	0.0024	32	Accepta ble	

Secondary exposure of general public

	Component	CAS	Inhalation Exposure [mg/m³]	Actual Dermal Total [mg/kg/d]	Oral exposure [mg/kg/d]
		Chronic	exposure		
Adult – Inhalation of volatilised residues, indoor	Deltamethrine	52918-63-5	6.9 x 10 ⁻⁷	na	na
Child – Inhalation of volatilised residues, indoor	of volatilised Deltamethrine		1.3 x 10 ⁻⁶	na	na
Adult – Dermal exposure with treated surface, indoor	Deltamethrine	52918-63-5	na	1.2 x 10⁻⁴	na
Child – Dermal exposure with treated surface, indoor	Deltamethrine	52918-63-5	na	1.5 x 10 ⁻⁴	na

Risk assessment

Product Assessment Report – SANITERPEN INSECTICIDE DK PRET A L'EMPLOI -Deltamethrin

	Component	AEL [mg/kg/	Absorp	tion [%]	Total syst	% AEL	Risk
	Component	[mg/kg/ d]	oral	dermal	exposure [mg/kg bw/d]	% AEL	KI5K
	Chronic exposure						
Adult – Inhalation of volatilised residues, indoor	Deltamethrine	0.0075	75	10	6.9 x 10 ⁻⁷	0.01	Acceptable
Child – Inhalation of volatilised residues, indoor	Deltamethrine	0.0075	75	10	1.3 x 10 ⁻⁶	0.02	Acceptable
Adult – Dermal exposure with treated surface, indoor	Deltamethrine	0.0075	75	10	1.2 x 10 ⁻⁴	1.6	Acceptable
Child – Dermal exposure with treated surface, indoor	Deltamethrine	0.0075	75	10	1.5 x 10 ⁻⁴	2.03	Acceptable

ConsExpo 4.1 report – Primary exposure for professionnals

file name: Report date: 16/03/2015

Product

SANITERPEN DK PAE

Compound

Compound name : CAS number : molecular weight vapour pressure KOW General Exposure Data	delatmethrin 52918-63-5 505 1,28E-8 4,6	g/mol Pascal 10Log
exposure frequency body weight	365 60	1/year kilogram
Inhalation model: Exposure to spray		
weight fraction compound exposure duration room volume ventilation rate mass generation rate spray duration airborn fraction weight fraction non-volatile density non-volatile room height inhalation cut-off diameter non-respirable uptake fraction Spraying away from exposed person	0,02 240 58 0,5 0,75 120 0,2 1 1,8 2,5 15 75	% minute m3 1/hr g/sec minute fraction fraction fraction g/cm3 meter micrometer %
uptake fraction inhalation rate	100 1,25	% m3/hour
Dermal model: Direct dermal contact with	product : constant rate	
weight fraction compound contact rate release duration Uptake model: fraction	0,02 46 120	% mg/min minute
uptake fraction	10	%
Output		70

Inhalation (point estimates)

Product Assessment Report – SANITERPEN INSECTICIDE DK PRET A L'EMPLOI -Deltamethrin

	inhalation mean event concentration : inhalation mean concentration on day of e inhalation air concentration year average : inhalation acute (internal) dose : inhalation chronic (internal) dose :	•	0,00123 0,000205 0,000205 mg/kg mg/kg/day	mg/m3 mg/m3 mg/m3/day
Der	mal : point estimates			
	dermal load : dermal external dose : dermal acute (internal) dose : dermal chronic (internal) dose :	- 0,0184 0,00184 0,00184	mg/cm2 mg/kg mg/kg mg/kg/day	
<u>Ora</u>	I non-respirable: point estimates			
	oral external dose : oral acute (internal) dose : oral chronic (internal) dose :	0,000473 0,000473 0,000473	mg/kg mg/kg mg/kg/day	
Inte	grated (point estimates)			
	total external dose: total acute dose (internal): total chronic dose (internal):	0,0191 0,00242 0,00241	mg/kg mg/kg mg/kg/day	

Annex 7 : Safety for non-professional operators and the general public

SANITERPEN INSECTICIDE DK PRET A L'EMPLOI

Date: 19/02/2015

The product is for professional use only.

Annex 8: Residue behaviour

Deltamethrin

Date:19/02/2015

Regarding the intended used, exposure of livestock and human exposure via food of animal origin are not expected. Dietary risk assessment is not required.

Test substance	Test organisms	Test system / Concentrations applied / exposure time	Test conditions	Test results: effects, mode of action, resistance	Reference	RI
SANITERPEN INSECTICIDE DK PRET A I'EMPLOI	House fly (Musca domestica) nymphs + eggs Mosquito (Culex Pipiens) adults Poultry red mite (Dermanyssus gallinae) nymphs + adults Cat flea (Ctenocephalides felis) Nymphs + adults	Laboratory test	Room of 60 m ³ Average conditions of a warehouse with typical surfaces treated measured 15 cm x 15 cm (concrete, untreated wood, plaster panel and ceramic tiles) Constant temperature between 20 et 25 °C Relative humidity $65\% \pm 5\%$ Photoperiod: 16h light / 8 h darkness	100 % mortality was achieved within 24 hours Residual activity up to 3 months after application on both porous and non-porous surfaces	Serrano (TEC) (2013) 1558c/1112R	1
SANITERPEN INSECTICIDE EFFET CHOC PAE	House fly (Musca domestica) adults	Laboratory test Internal method derived from CEB n°135 2 hours of exposure time 50 ml/m ²	Ceramic tiles are treated with the product and after drying put in contact with house flies for 2 hours. The mortality of the insects are recording every 10 minutes until 2 hours of contact and finally after 24 and 48 hours. The same procedure is also applied with product stored after 3, 6, 12, 18, 24, 30 and 36 months. Constant temperature 20 °C \pm 1 °C. Relative humidity 60% \pm 5% Photoperiod: 16h light / 8 h dark	achieved within 24 hours even after 36 months of	Anonymous (TEC) (2011) 1296b/1008R	2

Annex 9: Efficacy of the active substance from its use in the biocidal product (*)

Test substance	Test organisms	Test system / Concentrations applied / exposure time	Test conditions	Test results: effects, mode of action, resistance	Reference	RI
SANITERPEN INSECTICIDE DK PRET A I'EMPLOI	<u>House fly</u> (Musca domestica) adults <u>Mosquitoes</u> (Aedes aegypti) adults	Laboratory test CEB n°135 1 hour of exposure time 50 ml/m ²	Room of 60 m ³ Average conditions of a warehouse with typical surfaces treated (concrete and ceramic tiles) Constant temperature between 20 and 25 °C Relative humidity $65\% \pm 5\%$ Photoperiod: 16h light / 8 h dark	100 % mortality was achieved within 24 hours Residual activity up to 3 months after application on both porous and non-porous surfaces	Serrano (TEC) (2014) 1826a/0914R	1
SANITERPEN INSECTICIDE DK PRET A I'EMPLOI	<u>Mosquitoes</u> (Aedes aegypti) Adult (<i>Culex pipiens</i>) Adult <u>Cat flea</u> (<i>Cténocephalides felis</i>) adult	Semi-field study BSI 4172 part 1&2 method 50 ml /m ²	Room of 30 m ³ (12 m ² floor) 1 typical surface measured 6 cm ² (ceramic tiles) Constant temperature 25 °C \pm 2°C Relative humidity 65% \pm 5% Dead insects are count 7 days after treatment	, , , , , , , , , , , , , , , , , , ,	Serrano (TEC) (2014) 1826b/0914R	2
SANITERPEN INSECTICIDE DK	House fly (<i>Musca domestica</i>) adults	Field study CEB n°107. 10 % v/v of diluted product, 50 ml/m ²	After a confirmation of the level of infestation, the product is applied on vertical surface by spraying. The reduction of	the treated breeding premises until 90 % from the first week and 95 % 2 weeks after the treatment and still 98 % after 3 months. The efficacy demonstrated is similar to the reference		1

Test substance	est organisms	Test system / Concentrations applied / exposure time	Test conditions	Test results: effects, mode of action, resistance	Reference	RI
SANITERPEN INSECTICIDE DK PAE (0.02% deltamethr SANITERPEN INSECTICIDE DK PAE Without deltameth	<u>Sand fly</u> (<u>Phlebotomus</u> <u>duboscqi)</u> <u>Adults</u>	Laboratory test ASTM E 654-96 Direct spraying Dose: 1.57 ml on the 0.0314 m ² test surface = 50 ml/m ² .	Room of 30 m3 (12 m ² floor) Constant temperature 25°C +/- 2°C Relative humidity 65% +/- 5% Insects/mites were held into a glass ring of 20 cm diameter (0.0314 m ²) and 10 cm height. A steel grid was placed on the top in case of flying insects. 4 replicates Expression of results: KT 100 = time of exposure to knockdown 100% of insects/mites. Mortality after 24 hours. Mortality after 72 hours. Results of both products are compared.	The untreated control gave a low mortality (< 10%), the trial is validated. The marketed product SANITERPEN INSECTICIDE DK PAE has proved a fast (KT100 < 15 min) and a complete insecticide efficacy (mortality 24 hours = 100%) against all the pests in testing; The same product but without deltamethrin gave no significant efficacy (<10%) against all the pests in testing;	Serrano, (TEC), (2015) 1948c/0615R a	2