ANNEX III

INFORMATION REQUIREMENTS FOR BIOCIDAL PRODUCTS

- 1. This Annex sets out the information requirements that shall be included in the dossier for the biocidal product accompanying an application for the approval of an active substance in accordance with point (b) of Article 6(1) and the dossier accompanying an application for the authorisation of a biocidal product in accordance with point (a) of Article 20(1).
- 2. The data elements set down in this Annex comprise a Core Data Set (CDS) and an Additional Data Set (ADS). The data elements belonging to the CDS are considered as the basic data which should, in principle, be provided for all biocidal products.

With regard to the ADS, the data elements to be provided for a specific biocidal product shall be determined by considering each of the ADS data elements indicated in this Annex taking into account, inter alia, the physical and chemical properties of the product, existing data, information which is part of the CDS and the types of products and the exposure patterns related to these uses.

Specific indications for the inclusion of some data elements are provided in column 1 of the Annex III table. The general considerations regarding adaptation of information requirements as set out in Annex IV to this Regulation shall also apply. In light of the importance of reducing testing on vertebrates, column 3 of the table gives specific indications for the adaptation of some of the data elements which might require the use of such tests on vertebrates.

For some of the information requirements set out in this Annex, it may be possible to satisfy these requirements based on available information of the properties of the active substance(s) contained in the product and the properties of non-active substance(s) included in the product. For non-active substances, applicants shall use the information provided to them in the context of Title IV of Regulation (EC) No 1907/2006, where relevant, and the information made available by the Agency in accordance with point (e) of Article 77(2) of that Regulation.

The relevant calculation methods used for the classification of mixtures as laid down in Regulation (EC) No 1272/2008 shall, where appropriate, be applied in the hazard assessment of the biocidal product. Such calculation methods shall not be used if, in relation to a particular hazard, synergistic and antagonistic effects between the different substances contained in the product are considered likely.

Detailed technical guidance regarding the application of this Annex and the preparation of the dossier is available on the website of the Agency.

The applicant has the obligation to initiate a pre-submission consultation. In addition to the obligation set out in Article 62(2), applicants may also consult with the competent authority that will evaluate the dossier with regard to the proposed information requirements and in particular the testing on vertebrates that the applicant proposes to carry out.

Additional information may need to be submitted if necessary to carry out the evaluation as indicated in Article 29(3) or Article 44(2).

The information submitted shall, in any case, be sufficient to support a risk assessment demonstrating that the criteria in Article 19(1)(b) are met.

3. A detailed and full description of studies conducted and of the methods used shall be included. It is important to ensure that the data available is relevant and is of sufficient quality to fulfil the requirements.

- 4. The formats made available by the Agency shall be used for submission of the dossiers. In addition, IUCLID shall be used for those parts of the dossiers to which IUCLID applies. Formats and further guidance on data requirements and dossier preparation are available on the Agency homepage.
- 5. Tests submitted for the purpose of authorisation shall be conducted according to the methods described in Regulation (EC) No 440/2008. However, if a method is inappropriate or not described, other methods shall be used which are scientifically appropriate, whenever possible internationally recognised, and their appropriateness must be justified in the application. When test methods are applied to nanomaterials, an explanation shall be provided of their scientific appropriateness for nanomaterials, and, where applicable, of the technical adaptations/adjustments that have been made in order to respond to the specific characteristics of these materials.
- 6. Tests performed should comply with the relevant requirements of protection of laboratory animals, set out in Directive 2010/63/EU and, in the case of ecotoxicological and toxicological tests, good laboratory practice, set out in Directive 2004/10/EC or other international standards recognised as being equivalent by the Commission or the Agency. Tests on physico-chemical properties and safety-relevant substance data should be performed at least according to international standards.
- 7. Where testing is done, a detailed quantitative and qualitative description (specification) of the product used for each test and its impurities must be provided.
- 8. Where test data exist that have been generated before 17 July 2012 by methods other than those laid down in Regulation (EC) No 440/2008, the adequacy of such data for the purposes of this Regulation and the need to conduct new tests according to the Regulation (EC) No 440/2008 must be decided by the competent authority of the Member State, on a case-by-case basis, taking into account, among other factors, the need to avoid unnecessary testing.
- 9. New tests involving vertebrates shall be conducted as the last available option to comply with the data requirements set out in this Annex when all the other data sources have been exhausted. In vivo testing with corrosive substances at concentration/dose levels causing corrosivity shall also be avoided.

TITLE 1

CHEMICAL PRODUCTS

Core data set and additional data set for chemical products

Information required to support the authorisation of a biocidal product is listed in the table below.

For each information requirement set down in this Annex the indications given in columns 1 and 3 of Annex II for the same information requirement shall also apply.

Column 1Information required:	Column 2All data is CDS unless indicated as ADS	Column 3Specific rules for adaptation from standard information concerning some of the information requirements that may
a Eva irritation test shall not be needed	sary where the biocidal product has been sh	own to have notential corrective properties.

a Eye-irritation test shall not be necessary where the biocidal product has been shown to have potential corrosive properties.

		require recourse to testing of vertebrates
LICANT		
Name and address, etc.		
Contact person		
Manufacturer and formulator of the biocidal product and the active substance(s) (names, addresses, including location of plant(s))		
NTITY OF THE DAL PRODUCT		
Trade name or proposed trade name		
Manufacturer's development code and number of the product, if appropriate		
Complete quantitative (g/kg, g/ l or % w/w (v/ v)) composition of the biocidal product, i.e. declaration of all active substances and non- active substances (substance or mixture according to Article 3 of Regulation (EC) No 1907/2006), which are intentionally added to the biocidal product (formulation) as well as detailed quantitative and		
	Name and address, etc. Contact person Manufacturer and formulator of the biocidal product and the active substance(s) (names, addresses, including location of plant(s)) NTITY OF THE DAL PRODUCT Trade name or proposed trade name Manufacturer's development code and number of the product, if appropriate Complete quantitative (g/kg, g/ l or % w/w (v/ v)) composition of the biocidal product, i.e. declaration of all active substances and non- active substances (substance or mixture according to Article 3 of Regulation (EC) No 1907/2006), which are intentionally added to the biocidal product (formulation) as well as detailed quantitative and	Name and address, etc. Contact person Manufacturer and formulator of the biocidal product and the active substance(s) (names, addresses, including location of plant(s)) VTITY OF THE DAL PRODUCT Trade name or proposed trade name Manufacturer's development code and number of the product, if appropriate Complete quantitative (g/kg, g/ 1 or % w/w (v/ v)) composition of the biocidal product, i.e. declaration of all active substances and non- active substances (substance or mixture according to Article 3 of Regulation (EC) No 1907/2006), which are intentionally added to the biocidal product (formulation) as

In additio informatic ingredient and, in the reaction n compositi	qualitative information on the composition of the active substance(s) contained in the biocidal product. For non-active substances, a safety data sheet in compliance with Article 31 of Regulation (EC) No 1907/2006 has to be provided. n, all relevant on on individual ts, their function e case of a nixture, the final ion of the biocidal hall be given	
	Formulation type and nature of the biocidal product, e.g. emulsifiable concentrate, wettable powder, solution	
	Where the biocidal product contains an active substance that has been manufactured in locations or according to processes or from starting materials other than those of the active substance evaluated for the purpose of approval pursuant to Article 9 of this Regulation, evidence has to be provided that technical equivalence has been established in accordance with	

a Eye-irritation test shall not be necessary where the biocidal product has been shown to have potential corrosive properties.

	Article 54 of this Regulation or has been established, following an evaluation having started before 1 September 2013, by a competent authority designated in accordance with Article 26 of Directive 98/8/EC		
CHEM TECH	SICAL, ICAL AND NICAL ERTIES		
	pearance (at 20 °C 1,3 kPa)	-	
3.1.1.	Physical state (at 20 °C and 101,3 kPa)		
3.1.2.	Colour (at 20 °C and 101,3 kPa)		
3.1.3.	Odour (at 20 °C and 101,3 kPa)		
the pH o or its dis	Acidity/alkalinity is applicable when of the biocidal product opersion in water (1) tside the pH range		
3.3.	Relative density (liquids) and bulk, tap density (solids)		
	rage stability, y and shelf-life	I	<u> </u>
	torage stability	-	
3.4.1.1.	Accelerated storage test		
3.4.1.2.	Long term storage test at ambient temperature		
a Eye-irritation test shall not be necessary where the biocidal product has been shown to have potential corrosive properties.			

		1	
	Low temperature stability test (liquids)		
the acti technic	ffects on content of ve substance and al characteristics of cidal product		
3.4.2.1.	Light		
3.4.2.2.	Temperature and humidity		
3.4.2.3.	Reactivity towards container material		
	hnical ceristics of the l product		
3.5.1.	Wettability		
3.5.2.	Suspensibility, spontaneity and dispersion stability		
3.5.3.	Wet sieve analysis and dry sieve test		
3.5.4.	Emulsifiability, re- emulsifiability and emulsion stability		
3.5.5.	Disintegration time		
3.5.6.	Particle size distribution, content of dust/fines, attrition, friability		
3.5.7.	Persistent foaming		
3.5.8.	Flowability/ Pourability/ Dustability		
3.5.9.	Burning rate — smoke generators		
a Eve-ir	ritation test shall not be necess	ary where the biocidal product has been sho	own to have potential corrosive properties.

3.5.10.	Burning completeness — smoke generators		
3.5.11.	Composition of smoke — smoke generators		
3.5.12.	Spraying pattern — aerosols		
3.5.13.	Other technical characteristics		
compa produc biocida	ysical and chemical tibility with other ets including other al products with its use is to be ised		
3.6.1.	Physical compatibility		
3.6.2.	Chemical compatibility		
3.7.	Degree of dissolution and dilution stability		
3.8.	Surface tension		
3.9.	Viscosity		
AND R	SICAL HAZARDS RESPECTIVE ACTERISTICS		
4.1.	Explosives		
4.2.	Flammable gases		
4.3.	Flammable aerosols		
4.4.	Oxidising gases		
4.5.	Gases under pressure		
a Eye-i	rritation test shall not be necess	sary where the biocidal product has been sh	own to have potential corrosive properties.

16	Elemmohla li avid-		
4.6.	Flammable liquids		
4.7.	Flammable solids		
4.8.	Self-reactive substances and mixtures		
4.9.	Pyrophoric liquids		
4.10.	Pyrophoric solids		
4.11.	Self-heating substances and mixtures		
4.12.	Substances and mixtures which in contact with water emit flammable gases		
4.13.	Oxidising liquids		
4.14.	Oxidising solids		
4.15.	Organic peroxides		
4.16.	Corrosive to metals		
	dditional physical ions of hazard		<u>.</u>
maicati	ions of nazaru		
4.17.1.	Auto-ignition temperatures of products (liquids and gases)		
4.17.2.	Relative self- ignition temperature for solids		
4.17.3.	Dust explosion hazard		
DETEC	5. METHODS OF DETECTION AND IDENTIFICATION		

a Eye-irritation test shall not be necessary where the biocidal product has been shown to have potential corrosive properties.

Analytical method including validation parameters for determining the concentration of the active substance(s), residues, relevant impurities and substances of concern in the biocidal product		
In so far as not covered by Annex II 5.2 and 5.3, analytical methods for monitoring purposes including recovery rates and the limits of determination of relevant components of the biocidal product and/or residues thereof, where relevant in or on the following:	ADS	
Soil	ADS	
Air	ADS	
Water (including drinking water) and sediment	ADS	
Animal and human body fluids and tissues	ADS	
Analytical methods for monitoring purposes including recovery rates and the limit of quantification and detection for the active substance,	ADS	wy to have potential corrective properties
	 including validation parameters for determining the concentration of the active substance(s), residues, relevant impurities and substances of concern in the biocidal product In so far as not covered by Annex II 5.2 and 5.3, analytical methods for monitoring purposes including recovery rates and the limits of determination of relevant components of the biocidal product and/or residues thereof, where relevant in or on the following: Soil Air Water (including drinking water) and sediment Analytical methods for monitoring purposes including recovery rates and the limit of quantification and detection for the active substance, 	including validation parameters for determining the concentration of the active substance(s), residues, relevant impurities and substances of concern in the biocidal productADSIn so far as not covered by Annex, II 5.2 and 5.3, analytical methods for monitoring purposes including recovery rates and the limits of determination of relevant components of the biocidal productADSSoilADSAirADSWater (including drinking water) and sedimentADSAnimal and human body fluids and tissuesADSAnalytical methods for monitoring purposes including recovery rates and the limits of determination of relevant components of the biocidal product and/or residuesADSADSADSADSAirADSADSADSAuseADSAuseADS

	and for residues thereof, in/on food of plant and animal origin or feeding stuffs and other products where relevant (not necessary if neither the active substance nor the material treated with it come into contact with food- producing animals, food of plant and animal origin or feeding stuffs)		
AGAI	TECTIVENESS NST TARGET ANISMS		
	Function, e.g. fungicide, rodenticide, insecticide, bactericide of control e.g. ng, killing, inhibiting		
6.2.	Representative organism(s) to be controlled and products, organisms or objects to be protected		
6.3.	Effects on representative target organisms		
6.4.	Likely concentration at which the active substance will be used		
6.5.	Mode of action (including time delay)		
a Eye-	a Eye-irritation test shall not be necessary where the biocidal product has been shown to have potential corrosive properties.		

		1	1
6.6.	The proposed label claims for the product and, where label claims are made, for treated articles		
6.7.	Efficacy data to support these claims, including any available standard protocols, laboratory tests or field trials used including performance standards where appropriate and relevant		
	ny known limitations	<u></u>	<u> </u>
on effi	cacy		
6.8.1.	Information on the occurrence or possible occurrence of the development of resistance and appropriate management strategies		
6.8.2.	Observations on undesirable or unintended side effects e.g. on beneficial and other non-target organisms		
6.9.	Summary and evaluation		
7. INT EXPO	ENDED USES AND SURE		
7.1. a Eye-	Field(s) of use envisaged for biocidal products and, where	sary where the biocidal product has been sho	own to have potential corrosive properties.
J -		- 1	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

7.2. Product-type 7.3. Detailed description of intended use pattern(s) for biocidal products and, where appropriate, treated articles 7.4. User e.g. industrial, trained professional, professional or general public (non- professional) 7.5. Likely tonnage to be placed on the market per year and, where relevant, for different use categories 7.6. Method of application and a description of this method 7.7. Application final concentration of the biocidal product and active substance in a treated article or in the system in which the product is to be used, e.g. cooling water, water used for heating purposes 7.8. Number and timing of applications, and where relevant, any varticular		appropriate, treated articles	
of intended use pattern(s) for biocidal products and, where appropriate, treated articles	7.2.	Product-type	
trained professional, professional or general public (non-professional) 7.5. Likely tonnage to be placed on the market per year and, where relevant, for different use categories 7.6. Method of application and a description of this method 7.7. Application rate and, if appropriate, the final concentration of the biocidal product and active substance in a treated article or in the system in which the product is to be used, e.g. cooling water, water used for heating purposes 7.8. Number and timing of applications, and where relevant, and the system is	7.3.	of intended use pattern(s) for biocidal products and, where appropriate, treated	
be placed on the market per year and, where relevant, for different use categories 7.6. Method of application and a description of this method 7.7. Application rate and, if appropriate, the final concentration of the biocidal product and active substance in a treated article or in the system in which the product is to be used, e.g. cooling water, surface water, water used for heating purposes 7.8. Number and timing of applications, and where relevant,	7.4.	trained professional, professional or general public (non-	
application and a description of this method 7.7. Application rate and, if appropriate, the final concentration of the biocidal product and active substance in a treated article or in the system in which the product is to be used, e.g. cooling water, surface water, water used for heating purposes 7.8. Number and timing of applications, and where relevant,	7.5.	be placed on the market per year and, where relevant, for different use	
 rate and, if appropriate, the final concentration of the biocidal product and active substance in a treated article or in the system in which the product is to be used, e.g. cooling water, surface water, water used for heating purposes 7.8. Number and timing of applications, and where relevant, 	7.6.	application and a description of this	
of applications, and where relevant,	7.7.	rate and, if appropriate, the final concentration of the biocidal product and active substance in a treated article or in the system in which the product is to be used, e.g. cooling water, surface water, water used	
 a Eye-irritation test shall not be necessary where the biocidal product has been shown to have potential corrosive properties. 		of applications, and where relevant, any particular	runkara tha bigaidal product has been shown to have a startish summing a

	information relating to geographical location or climatic variations including necessary waiting periods, clearance times, withdrawal periods or other precautions to protect human health, animal health and the environment	
7.9.	Proposed instructions for use	
conform	xposure data in mity with Annex VI Regulation	
7.10.1.	Information on human exposure associated with production and formulation, proposed/expected uses and disposal	
7.10.2.	Information on environmental exposure associated with production and formulation, proposed/expected uses and disposal	
7.10.3.	Information on exposure from treated articles including leaching data (either laboratory studies or model data)	
7.10.4.	Information regarding other products that the product is likely to be used together with, in particular	own to have potential corrosive properties.

the identity of the active substances in these products, if relevant, and the likelihood of any interactions 8. TOXICOLOGICAL PROFILE FOR HUMANS	
AND ANIMALS 8.1. Skin corrosion or skin irritation The assessment of this endpoint shall be carried out according to the sequential testing strategy for dermal irritation and corrosion set out in the Appendix to Test Guideline B.4. Acute Toxicity-Dermal Irritation/ Corrosion (Annex B.4. to Regulation (EC) No 440/2008)	Testing on the product/ mixture does not need to be conducted if: — there are valid data available on each of the components in the mixture sufficient to allow classification of the mixture according to the rules laid down in Directive 1999/45/EC and Regulation (EC) No 1272/2008 (CLP), and synergistic effects between any of the components are not expected
8.2. Eye irritation ^a The assessment of this endpoint shall be carried out according to the sequential testing strategy for eye irritation and corrosion as set down in the Appendix to Test Guideline B.5.Acute Toxicity: Eye Irritation/ Corrosion (Annex B.5. to Regulation (EC) No 440/2008)	Testing on the product/ mixture does not need to be conducted if: — there are valid data available on each of the components in the mixture to allow classification of the mixture according to the rules laid down in Directive 1999/45/ECand Regulation (EC) No 1272/2008 (CLP), and synergistic effects between any of the components are not expected
8.3. Skin sensitisation a Eye-irritation test shall not be necessary where the biocida	Testing on the product/ mixture does not need to be conducted if: I product has been shown to have potential corrosive properties.

2. in vivo The Mu Lymph (LLNA) where a the redu of the as first-cho for in vi If anoth sensitisa used jus	mprise the utive steps: ssment vailable animal and ive data testing rine Local Node Assay) including, ppropriate, iced variant ssay, is the pice method ivo testing.			there are valid data available on each of the components in the mixture to allow classification of the mixture according to the rules laid down in Directive 1999/45/EC and Regulation (EC) No 1272/2008 (CLP), and synergistic effects between any of the components are not expected the available information indicates that the product should be classified for skin sensitisation or corrosivity; or the substance is a strong acid (pH < 2,0) or base (pH > 11,5)
8.4. Respira sensitisa		ADS		on the product/ does not need to be ed if: there are valid data available on each of the components in the mixture to allow classification of the mixture according to the rules laid down in Directive 1999/45/EC and Regulation (EC) No 1272/2008 (CLP), and synergistic effects between any of the components are not expected
the tiere to class of mixtu acute to	cation using ed approach ification ures for xicity in	ary where the biocidal product has been sh	mixture conducte	there are valid data available on each of the components in

	Regulation (EC) No 1272/2008 is the default approach	class mixt to th down 1999 Regu 1272 and s effec of th	nixture to allow dification of the ure according e rules laid n in Directive 0/45/EC and dation (EC) No 0/2008 (CLP), synergistic ets between any e components not expected
8.5.1.	By oral route		
8.5.2.	By inhalation		
8.5.3.	By dermal route		
8.5.4.	For biocidal products that are intended to be authorised for use with other biocidal products, the risks to human health, animal health and the environment arising from the use of these product combinations shall be assessed. As an alternative to acute toxicity studies, calculations can be used. In some cases, for example where there are no valid data available of the kind set out in column 3, this may require a limited number of acute toxicity studies to be carried out using combinations of the products	conducted if: — there avail the c the n class mixt to th down 1999 Regu 1272 and s effec of th	mixture of not need to be e are valid data able on each of components in nixture to allow diffication of the ure according e rules laid n in Directive 0/45/EC and alation (EC) No 2/2008 (CLP), synergistic ets between any e components not expected
8.6.	Information on dermal absorption		
a Eye-	-	ary where the biocidal product has been shown to have potentia	l corrosive properties.

Information on dermal absorption when exposure occurs to the biocidal product. The assessment of this endpoint shall proceed using a tiered approach		
 8.7. Available toxicological data relating to: non-active substance(s) (i.e. substance(s) of concern), or a mixture that a substance(s) of concern is a component of If insufficient data are available for a non-active substance(s) and cannot be inferred through read- across or other accepted non- testing approaches, targeted test(s) described in Annex II shall be carried out for the substance(s) of concern or a mixture that a substance(s) of concern is a component of 		Testing on the product/ mixture does not need to be conducted if: — there are valid data available on each of the components in the mixture to allow classification of the mixture according to the rules laid down in Directive 1999/45/EC and Regulation (EC) No 1272/2008 (CLP)
8.8. Food and feedingstuffs studies	ADS	
8.8.1. If residues of the biocidal product remain in or on feedingstuffs for a significant period of time, then feeding and metabolism studies in livestock shall be required to permit evaluation of residues in food of animal origin	ADS	
 8.9. Effects of industrial processing and/ or domestic preparation on a Eve-irritation test shall not be necess 	ADS	own to have potential corrosive properties.

	and a rill be biocidal certain are applied nd livestock es) residue e needed			
	LOGICAL			
9. ECOTOXICO STUDIES				
to the of the produce sufficient a dece made of classiff produce 	hation relating e ecotoxicity he biocidal et which is ent to enable cision to be concerning the fication of the et is required there are lata available h of the onents in xture and fistic effects en any of mponents t expected, fication of xture can be according rules laid in Directive k5/EC, ation (EC) 07/2006 CH) and ation (EC) No 2008 (CLP)			

Where valid data on the components are not available or where synergistic effects may be expected then testing of components and/ or the biocidal product itself may be necessary		
9.2. Further Ecotoxicological studies Further studies chosen from among the endpoints referred to in Section 9 of Annex II for relevant components of the biocidal product or the biocidal product or the biocidal product itself may be required if the data on the active substance cannot give sufficient information and if there are indications of risk due to specific properties of the biocidal product		
9.3. Effects on any other specific, non-target organisms (flora and fauna) believed to be at risk	ADS	Data for the assessment of hazards to wild mammals are derived from the mammalian toxicological assessment
9.4. If the biocidal product is in the form of bait or granules the following studies may be required:	1	
9.4.1. Supervised trials to assess risks to non-target organisms under field conditions		
 9.4.2. Studies on acceptance by ingestion of the biocidal product by any non-target a Eye-irritation test shall not be necess 	sary where the biocidal product has been sh	

	organisms thought to be at risk		
9.5.	Secondary ecological effect e.g. when a large proportion of a specific habitat type is treated	ADS	
	IRONMENTAL ND BEHAVIOUR		
are appli	requirements below cable only to the components of the product		
10.1.	Foreseeable routes of entry into the environment on the basis of the use envisaged		
among the to in Sect for relevant biocidal prequired. For produce outside, we to soil, we compone may influe behaviour of the acc Data are is scienti the fate of the produce outside outside behaviour of the acc Data are us scienti	Further studies on fate and behaviour in the environment tudies chosen from te endpoints referred tion 10 of Annex II ant components of dal product or the product itself may be that are used with direct emission ater or surfaces, the nts in the product tence the fate and r (and ecotoxicity) tive substance. required unless it fically justified that of the components in the trided for the active e and other identified es of concern	ADS	
10.3.	Leaching behaviour	ADS	

a Eye-irritation test shall not be necessary where the biocidal product has been shown to have potential corrosive properties.

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10.4.	Testing for distribution and dissipation in the following:	ADS	
10.4.1.	Soil	ADS	
10.4.2.	Water and sediment	ADS	
10.4.3.	Air	ADS	
10.5.	If the biocidal product is to be sprayed near to surface waters then an overspray study may be required to assess risks to aquatic organisms or plants under field conditions	ADS	
10.6.	If the biocidal product is to be sprayed outside or if potential for large scale formation of dust is given then data on overspray behaviour may be required to assess risks to bees and non-target arthropods under field conditions	ADS	
BE AD PROTI ANIM	ASURES TO OPTED TO ECT HUMANS, ALS AND THE RONMENT	I 	·
11.1. a Eye-i	Recommended methods and precautions concerning handling, use, storage, disposal, transport or fire	ary where the biocidal product has been sh	own to have notential corrective properties

11.2.	Identity of relevant combustion products in cases of fire		
11.3.	Specific treatment in case of an accident, e.g. first- aid measures, antidotes, medical treatment if available; emergency measures to protect the environment		
of destr decont	ossibility ruction or amination following in or on the ng:		1
11.4.1.	Air		
11.4.2.	Water, including drinking water		
11.4.3.	Soil		
11.5.	Procedures for waste management of the biocidal product and its packaging for industrial use, use by trained professionals, professional users and non- professional users (e.g. possibility of reuse or recycling, neutralisation, conditions for controlled discharge, and incineration)		
11.6.	Procedures for cleaning application	ary where the biocidal product has been sh	

	equipment where relevant		
11.7.	Specify any repellents or poison control measures included in the product that are present to prevent action against non- target organisms		
	ASSIFICATION, LING, AND AGING		
of Article including the hazar statemen the provi 1999/45/ (EC) No submitte Example	labels, instructions nd safety data sheets		
12.1.	Hazard classification		
12.2.	Hazard pictogram		
12.3.	Signal word		
12.4.	Hazard statements		
12.5.	Precautionary statements including prevention, response, storage and disposal		
12.6. a Eye-irr	Proposals for safety-data sheets should be provided, where appropriate	ary where the biocidal product has been she	nwn to have potential corrosive properties

12.7.	Packaging (type, materials, size, etc.), compatibility of the product with proposed packaging materials to be included	
13.	EVALUATION AND SUMMARY	
	information	
	d from the endpoints	
	ubsection (2-12) is	
	sed, evaluated and	
a draft ri	sk assessment is	
performe	ed	

a Eye-irritation test shall not be necessary where the biocidal product has been shown to have potential corrosive properties.

Textual Amendments

F1 Inserted by Commission Delegated Regulation (EU) No 837/2013 of 25 June 2013 amending Annex III to Regulation (EU) No 528/2012 of the European Parliament and of the Council as regards the information requirements for authorisation of biocidal products (Text with EEA relevance).

TITLE 2

MICRO-ORGANISMS

Core data set and additional data set

Information required to support the authorisation of a biocidal product is listed in the table below.

For each information requirement set down in this Annex the indications given in columns 1 and 3 of Annex II for the same information requirement shall also apply.

Column 1Information Column 2All dat required: unless indicated	L
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1. APPLICANT

1.1.	Name and address	
1.2.	Contact person	

1.3. 2 IDEN	Manufacturer and formulator of the biocidal product and the micro- organism(s) (names, addresses, including location of plant(s))		
	DAL PRODUCTS	1	
2.1.	Trade name or proposed trade name		
2.2.	Manufacturer's development code and number of the biocidal product, if appropriate		
individua the final	Detailed quantitative (g/kg, g/ l or % w/w (v/ v)) and qualitative information on the constitution, composition and function of the biocidal product, e.g. micro- organism, active substance(s) and product non-active substances and any other relevant components. ant information on al ingredients and composition of the product shall be		
2.4.	Formulation type and nature of the biocidal product		
[^{F1} 2.5.	Where the biocidal product contains an active substance that has been manufactured]

in locations or according to processes or from starting materials other than those of the active substance evaluated for the purpose of approval pursuant to Article 9 of this Regulation, evidence has to be provided that technical equivalence has been established in accordance with Article 54 of this Regulation or has been established, following an evaluation having started before 1 September 2013, by a competent authority designated in accordance with Article 26 of Directive 98/8/EC

3. BIOLOGICAL, PHYSICAL, CHEMICAL AND TECHNICAL PROPERTIES OF THE BIOCIDAL PRODUCT

3.1.	Biological properties of the micro-organism in the biocidal product	
	ppearance (at 20 °C 1,3 kPa)	
3.2.1.	Colour (at 20 °C and 101,3 kPa)	
3.2.2.	Odour (at 20 °C and 101,3 kPa)	
3.3.	Acidity, alkalinity and pH value	
3.4.	Relative density	

3.5. Storage stability, stability and shelf-life

3.5.1. Effects of light 3.5.2. Effects of temperature and humidity 3.5.3. Reactivity towards the container 3.5.4. Other factors affecting stability 3.6.7. Fechnical characteristics of the biocidal product 3.6.1. Wettability 3.6.2. Suspensibility and suspension stability 3.6.3. Wet sieve analysis and dry sieve test 3.6.4. Emulsifiability, re-emulsifiability, emulsion stability 3.6.5. Particle size distribution content of dust/fines, attrition and friability 3.6.6. Persistent foaming 3.6.7. Flowability/ 3.6.8. Burning rate — smoke generators 3.6.9. Burning completeness — smoke generators			
temperature and humidity 3.5.3. Reactivity towards the container 3.5.4. Other factors affecting stability 3.6. Technical characteristics of the biocidal product 3.6.1. Wettability 3.6.2. Suspensibility and suspension stability 3.6.3. Wet sieve analysis and dry sieve test 3.6.4. Emulsifiability, re-emulsifiability, emulsion stability 3.6.5. Particle size distribution content of dust/fines, attrition and friability 3.6.6. Persistent foaming 3.6.7. Flowability/ Pourability/ Dustability 3.6.8. Burning rate — smoke generators 3.6.9. Burning completeness —	3.5.1.	Effects of light	
the container 3.5.4. Other factors affecting stability 3.6. Technical characteristics of the biocidal product 3.6.1. Wettability 3.6.2. Suspensibility and suspension stability 3.6.3. Wet sieve analysis and dry sieve test 3.6.4. Emulsifiability, re-emulsifiability, emulsion stability 3.6.5. Particle size distribution content of dust/fines, attrition and friability 3.6.6. Persistent foaming 3.6.7. Flowability/ Pourability/ Dustability 3.6.8. Burning rate — smoke generators 3.6.9. Burning completeness —	3.5.2.	temperature and	
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characteristics of the biocidal product 3.6.1. Wettability 3.6.2. Suspensibility and suspension stability 3.6.3. Wet sieve analysis and dry sieve test 3.6.4. Emulsifiability, re-emulsifiability, emulsion stability 3.6.5. Particle size distribution content of dust/fines, attrition and friability 3.6.6. Persistent foaming 3.6.7. Flowability/ Pourability/ Dustability 3.6.8. Burning rate — smoke generators 3.6.9. Burning completeness —	3.5.4.		
3.6.2. Suspensibility and suspension stability 3.6.3. Wet sieve analysis and dry sieve test 3.6.4. Emulsifiability, re-emulsifiability, emulsion stability 3.6.5. Particle size distribution content of dust/fines, attrition and friability 3.6.6. Persistent foaming 3.6.7. Flowability/ Pourability/ Dustability 3.6.8. Burning rate — smoke generators 3.6.9. Burning completeness —	charact	eristics of the	
suspension stability 3.6.3. Wet sieve analysis and dry sieve test 3.6.4. Emulsifiability, re-emulsifiability, emulsion stability 3.6.5. Particle size distribution content of dust/fines, attrition and friability 3.6.6. Persistent foaming 3.6.7. Flowability/ Pourability/ Dustability 3.6.8. Burning rate — smoke generators 3.6.9. Burning completeness —	3.6.1.	Wettability	
and dry sieve test 3.6.4. Emulsifiability, re-emulsifiability, emulsion stability 3.6.5. Particle size distribution content of dust/fines, attrition and friability 3.6.6. Persistent foaming 3.6.7. Flowability/ Pourability/ Dustability 3.6.8. Burning rate — smoke generators 3.6.9. Burning completeness —	3.6.2.		
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3.6.7. Flowability/ Pourability/ Dustability 3.6.8. Burning rate — smoke generators 3.6.9. Burning completeness —	3.6.5.	distribution content of dust/fines, attrition and	
Pourability/ Dustability 3.6.8. Burning rate — smoke generators 3.6.9. Burning completeness —	3.6.6.	Persistent foaming	
smoke generators 3.6.9. Burning completeness —	3.6.7.	Pourability/	
completeness —	3.6.8.		
	3.6.9.		

3.6.10.	Composition of smoke — smoke generators	
3.6.11.	Spraying patterns — aerosols	
3.6.12.	Other technical characteristics	
biologi with ot includi produc	ysical, chemical and cal compatibility her products ng biocidal ts with which its o be authorised or red	
3.7.1.	Physical compatibility	
3.7.2.	Chemical compatibility	
3.7.3.	Biological compatibility	
3.8.	Surface tension	
3.9.	Viscosity	
AND R	SICAL HAZARDS ESPECTIVE ACTERISITICS	
4.1.	Explosives	
4.2.	Flammable gases	
4.3.	Flammable aerosols	
4.4.	Oxidising gases	
4.5.	Gases under pressure	
4.6.	Flammable liquids	
4.7.	Flammable solids	

4.8.	Oxidising liquids		
4.9.	Oxidising solids		
4.10.	Organic peroxides		
4.11.	Corrosive to metals		
	ther physical ons of hazard		
4.12.1.	Auto-ignition temperatures of products (liquids and gases)		
4.12.2.	Relative self- ignition temperature for solids		
4.12.3.	Dust explosion hazard		
DETEC	HODS OF CTION AND IFICATION	-	
5.1.	Analytical method for determining the concentration of the micro-organism(s) and substances of concern in the biocidal product		
5.2.	Analytical methods for monitoring purposes including recovery rates and the limit of quantification and detection for the active substance, and for residues thereof, in/on food of plant and animal origin or feeding stuffs and other products where relevant (not necessary if neither the active	ADS	

	substance nor the article treated with it does not come into contact with food-producing animals, food of plant and animal origin or feeding stuffs)	
AGA	FECTIVENESS INST TARGET ANISM	
6.1.	Function and mode of control	
6.2.	Representative pest organism(s) to be controlled and products, organisms or objects to be protected	
6.3.	Effects on representative target organisms	
6.4.	Likely concentration at which micro- organism will be used	
6.5.	Mode of action	
6.6.	The proposed label claims for the product	
6.7.	Efficacy data to support these claims, including any available standard protocols, laboratory tests, or field trials used including performance standards, where appropriate and relevant	

6.8. Any other known limitations on efficacy including resistance

	9	1	· · · · · · · · · · · · · · · · · · ·
6.8.1.	Information on the occurrence or possible occurrence of the development of resistance and appropriate management strategies		
6.8.2.	Observations on undesirable or unintended side effects		
7. INT EXPO	ENDED USES AND SURE		
7.1.	Field of use envisaged		
7.2.	Product-type		
7.3.	Detailed description of intended use		
7.4.	User e.g. industrial, trained professional, professional or general public (non- professional)		
7.5.	Method of application and a description of this method		
7.6.	Application rate and if appropriate the final concentration of the biocidal product or the micro-organism active substance in a treated article or the system in which the product is to be used (e.g. in the		

	application device or bait)	
relating location including periods f necessar or other protect h	Number and timing of applications and duration of protection ticular information to the geographical or climatic variations g necessary waiting for re-entry or y withdrawal period precautions to numan health, animal and the environment	
7.8.	Proposed instructions for use	
7.9. Exj	posure data	
7.9.1.	Information on human exposure associated with the proposed/expected uses and disposal	
7.9.2.	Information on environmental exposure associated with the proposed/ expected uses and disposal	
8.	TOXICOLOGICAL PROFILE FOR HUMANS AND ANIMALS	Testing on the product/ mixture does not need to be conducted if: — there are valid data available on each of the components in the mixture to allow classification of the mixture according to the rules laid down in Directive 1999/45/ EC, Regulation (EC) No 1907/2006 (REACH) and Regulation (EC) No 1272/2008 (CLP) and synergistic

			effects between any of the components are not expected
8.1.	Skin corrosion or irritation		
8.2.	Eye irritation		
8.3.	Skin sensitisation		
8.4.	Respiratory sensitisation	ADS	
8.5.	Acute toxicity Classification using the tiered approach to classification of mixtures for acute toxicity in Regulation (EC) No 1272/2008 is the default approach		
8.5.1.	Oral		
8.5.2.	Inhalation		
8.5.3.	Dermal		
8.5.4.	Additional acute toxicity studies		
8.6.	Information on dermal absorption if required		
8.7.	Available toxicological data relating to: non-active substance(s) (i.e. substance(s) of concern), or a mixture that a substance(s) of concern is a component of If insufficient data are available for a non-active		Testing on the product/ mixture does not need to be conducted if: — there are valid data available on each of the components in the mixture to allow classification of the mixture according to the rules laid down in Directive 1999/45/ EC, Regulation (EC) No 1907/2006

substance(s) and cannot be inferred through read-across or other accepted non-testing approaches, targeted test(s) described in Annex II, shall be carried out for the substance(s) of concern or a mixture that a substance(s) of concern is a component of		(REACH) and Regulation (EC) No 1272/2008 (CLP), and synergistic effects between any of the components are not expected
8.8. Supplementary studies for combinations of biocidal products For biocidal products that are intended to be authorised for use with other biocidal products, the risks to humans, animals and the environment arising from the use of these product combinations shall be assessed. As an alternative to acute toxicity studies, calculations can be used. In some cases, for example where there are no valid data available of the kind set out in column 3, this may require a limited number of acute toxicity studies to be carried using combinations of the products		Testing on the mixture of products does not need to be conducted if: — there are valid data available on each of the components in the mixture to allow classification of the mixture according to the rules laid down in Directive 1999/45/ EC, Regulation (EC) No 1907/2006 (REACH) and Regulation (EC) No 1272/2008 (CLP), and synergistic effects between any of the components are not expected
8.9. Residues in or on treated articles, food and feedingstuffs	ADS	

9. ECOTOXICOLOGICAL STUDIES

9.1. Information relating to the ecotoxicity of the biocidal product which is sufficient to enable a decision to be

	made concerning the classification of the product is required Where there are valid data available on each of the components in the mixture and synergistic effects between any of the components are not expected, classification of the mixture can be made according to the rules laid down in Directive 1999/45/EC, Regulation (EC) No 1907/2006 (REACH) and Regulation (EC) No 1272/2008 (CLP) Where valid data on the components are not available or where synergistic effects may be expected then testing of components and/ or the biocidal product itself may be necessary	
among t to in Sec II 'Micr relevant the bioc biocidal be requi active sufficien there are due to s	Further ecotoxicological studies studies chosen from the endpoints referred ction 8 of Annex ro-organisms' for t components of idal product or the product itself may ired if the data on the ubstance cannot give nt information and if e indications of risk pecific properties of idal product	

9.3.	Effects on any other specific non-target organisms (flora and fauna) believed to be at risk	ADS	Data for the assessment of hazards to wild mammals are derived from the mammalian toxicological assessment
9.4.	If the biocidal product is in the form of bait or granules	ADS	
9.4.1.	Supervised trials to assess risks to non-target organisms under field conditions		
9.4.2.	Studies on acceptance by ingestion of the biocidal product by any non-target organisms thought to be at risk		
9.5.	Secondary ecological effect e.g. when a large proportion of a specific habitat type is treated	ADS	
	VIRONMENTAL AND BEHAVIOUR		
10.1.	Foreseeable routes of entry into the environment on the basis of the use envisaged		
informat Section 9 'Micro-or required For prod outside, '	Further studies on fate and behaviour in the environment elevant, all the ion required in 0 of Annex II organisms' may be for the product ucts that are used with direct emission vater or surfaces, the	ADS	

may infl behaviou of the ac Data are is scienti the fate of the product data provisubstance	ents in the product uence the fate and ar (and ecotoxicity) tive substance. required unless it fically justified that of the components in uct is covered by the vided for the active e and other identified es of concern		
10.3.	Leaching behaviour	ADS	
10.4.	If the biocidal product is to be sprayed outside or if potential for large scale formation of dust is given then data on overspray behaviour may be required to assess risks to bees under field conditions	ADS	
BE AD PROTE ANIMA	ASURES TO OPTED TO ECT HUMANS, ALS AND THE ONMENT		
11.1.	Recommended methods and precautions concerning: handling, storage, transport or fire		
11.2.	Measures in the case of an accident		
for dest deconta	ocedures cruction or mination of the l product and its ing		
11.3.1.	Controlled incineration		
11.3.2.	Others		

11.4.	Packaging and compatibility of the biocidal product with proposed packaging materials	
11.5.	Procedures for cleaning application equipment where relevant	
11.6.	Monitoring plan to be used for the active micro- organism and other micro-organism(s) contained in the biocidal product including handling, storage, transport and use	
LABE	ASSIFICATION, LLING AND AGING	
for use a	e labels, instructions and safety data sheets provided	
12.1.	Indication on the need for the biocidal product to carry the biohazard sign specified in Annex II to Directive 2000/54/ EC	
12.2.	Precautionary statements including prevention, response, storage and disposal	
12.3.	Proposals for safety-data sheets should be provided, where appropriate	

12.4.	Packaging (type, materials, size, etc.), compatibility of the product with proposed packaging materials to be included	
identifie in each summar	SUMMARY AND EVALUATION information ed from the endpoints subsection (2-12) is ised, evaluated and isk assessment is ed	

Changes to legislation:

There are outstanding changes not yet made to Regulation (EU) No 528/2012 of the European Parliament and of the Council. Any changes that have already been made to the legislation appear in the content and are referenced with annotations. View outstanding changes

Changes and effects yet to be applied to :

Regulation applied (with modifications) by S.I. 2023/959 reg. 4(a)Sch. 1

Changes and effects yet to be applied to the whole legislation item and associated provisions

-	
-	Annex 3 para. 4 substituted by S.I. 2019/720 Sch. 2 para. 141(3)
_	Annex 3 para. 2 words omitted by S.I. 2019/720 Sch. 2 para. 141(2)(c)
_	Annex 3 para. 2 words omitted by S.I. 2019/720 Sch. 2 para. 141(2)(d)
_	Annex 3 para. 8 words omitted by S.I. 2019/720 Sch. 2 para. 141(5)
_	Annex 3 para. 2 words substituted by S.I. 2019/720 Sch. 2 para. 141(2)(a)
_	Annex 3 para. 2 words substituted by S.I. 2019/720 Sch. 2 para. 141(2)(b)
_	Annex 3 para. 6 words substituted by S.I. 2019/720 Sch. 2 para. 141(4)
_	Annex 2 para. 4 substituted by S.I. 2019/720 Sch. 2 para. 140(3)
_	Annex 2 para. 2 words omitted by S.I. 2019/720 Sch. 2 para. 140(2)(b)
_	Annex 2 para. 8 words omitted by S.I. 2019/720 Sch. 2 para. 140(5)
_	Annex 2 para. 2 words substituted by S.I. 2019/720 Sch. 2 para. 140(2)(a)
_	Annex 2 para. 6 words substituted by S.I. 2019/720 Sch. 2 para. 140(4)
_	Annex 4 para. 1.3 words omitted by S.I. 2019/720 Sch. 2 para. 142(b)
_	Annex 4 para. 1.5 words omitted by S.I. 2019/720 Sch. 2 para. 142(c)
_	Annex 4 para. 3.1 words omitted by S.I. 2019/720 Sch. 2 para. 142(d)
_	Annex 4 para. 1.2 words substituted by S.I. 2019/720 Sch. 2 para. 142(a)
_	Annex 6 para. 10 word substituted by S.I. 2019/720 Sch. 2 para. 143(6)
_	Annex 6 para. 13 words omitted by S.I. 2019/720 Sch. 2 para. 143(9)(a)
_	Annex 6 para. 15 words omitted by S.I. 2019/720 Sch. 2 para. 143(10)
_	Annex 6 para. 1 words substituted by S.I. 2019/720 Sch. 2 para. 143(2)(a)
_	Annex 6 para. 1 words substituted by S.I. 2019/720 Sch. 2 para. 143(2)(b)
_	Annex 6 para. 6 words substituted by S.I. 2019/720 Sch. 2 para. 143(3)
_	Annex 6 para. 8 words substituted by S.I. 2019/720 Sch. 2 para. 143(4)
_	Annex 6 para. 9 words substituted by S.I. 2019/720 Sch. 2 para. 143(5)(a)
-	Annex 6 para. 9 words substituted by S.I. 2019/720 Sch. 2 para. 143(5)(b)
_	Annex 6 para. 11 words substituted by S.I. 2019/720 Sch. 2 para. 143(7)
_	Annex 6 para. 12 words substituted by S.I. 2019/720 Sch. 2 para. 143(8)
-	Annex 6 para. 13 words substituted by S.I. 2019/720 Sch. 2 para. 143(9)(b)
-	Annex 6 para. 20 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
_	Annex 6 para. 26 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
_	Annex 6 para. 36 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 48 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 50 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 51 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 52 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 53 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 55 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 56 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 57 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 58 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 59 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 60 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 62 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 64 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)

-	Annex 6 para. 66 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 67 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 68 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 69 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 71 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 72 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 73 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 74 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 75 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 77 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 78 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
-	Annex 6 para. 52 words substituted by S.I. 2019/720 Sch. 2 para. 143(12)
-	Annex 6 para. 75 words substituted by S.I. 2019/720 Sch. 2 para. 143(13)
-	Annex 6 para. 77 words substituted by S.I. 2019/720 Sch. 2 para. 143(14) (This
	amendment not applied to legislation.gov.uk. Sch. 2 para. 143(14) substituted
	immediately before IP completion day by S.I. 2020/1567, reg. 1(2), Sch. 2 para.
	39(b))
-	Annex 6 para. 77 words substituted by S.I. 2019/720, Sch. 2 para. 143(14) (as
	substituted) by S.I. 2020/1567 Sch. 2 para. 39(b)
-	Annex 6 para. 52 words substituted in earlier amending S.I. 2019/720, Sch. 2 para.
	143(12) by S.I. 2020/1567 Sch. 2 para. 39(a)
-	Art. 1(2)(c) omitted by S.I. 2019/720 Sch. 2 para. 62(3)(b)
-	Art. 2(b) words substituted by S.I. 2019/720, Sch. 2 para. 63(2)(b) (as substituted) by
	S.I. 2020/1567 Sch. 2 para. 22
-	Art. 2(c) words substituted by S.I. 2019/720, Sch. 2 para. 63(2)(c) (as substituted) by
	S.I. 2020/1567 Sch. 2 para. 22
-	Art. 2(k) substituted by S.I. 2019/720, Sch. 2 para. 63(2)(d) (as substituted) by S.I.
	2020/1567 Sch. 2 para. 22
-	Art. 3(1)(d) words inserted by S.I. 2019/720 Sch. 2 para. 64(2)(a)
-	Art. 3(1)(e) words inserted by S.I. 2019/720 Sch. 2 para. 64(2)(b)
-	Art. 3(1)(f) words omitted by S.I. 2019/720 Sch. 2 para. 64(2)(c)
-	Art. 3(1)(k) words substituted by S.I. 2019/720 Sch. 2 para. 64(2)(d) (This
	amendment not applied to legislation.gov.uk. Sch. 2 para. 64(2)(d) substituted
	immediately before IP completion day by S.I. 2020/1567, reg. 1(2), Sch. 2 para.
	23(a))
-	Art. 3(1)(k) words substituted by S.I. 2019/720, Sch. 2 para. 64(2)(d) (as substituted)
	by S.I. 2020/1567 Sch. 2 para. 23(a)
-	Art. 3(1)(m) words omitted by S.I. 2019/720 Sch. 2 para. 64(2)(e)(i)
-	Art. 3(1)(m) words omitted by S.I. 2019/720 Sch. 2 para. 64(2)(e)(ii)
-	Art. 3(1)(n) substituted by S.I. 2019/720 Sch. 2 para. 64(2)(f)
-	Art. 3(1)(n) words substituted in earlier amending provision S.I. 2019/720, Sch. 2
	para. 64(2)(f) by S.I. 2020/1567 Sch. 2 para. 23(b)
-	Art. 3(1)(o) words omitted by S.I. 2019/720 Sch. 2 para. 64(2)(g)
-	Art. 3(1)(p) words substituted by S.I. 2019/720 Sch. 2 para. 64(2)(h)(i)
-	Art. 3(1)(p) words substituted by S.I. 2019/720 Sch. 2 para. 64(2)(h)(ii)
-	Art. 3(1)(p) words substituted in earlier amending provision S.I. 2019/720, Sch. 2
	para. 64(2)(h)(ii) by S.I. 2020/1567 Sch. 2 para. 23(c)
_	Art. 3(1)(t) words inserted by S.I. 2019/720 Sch. 2 para. 64(2)(i)
_	Art. 3(1)(x) omitted by S.I. 2019/720 Sch. 2 para. 64(2)(j)
_	Art. 3(1)(af)-(ah) inserted by S.I. 2019/720 Sch. 2 para. 64(2)(k)
_	Art. 3(1)(ai) substituted in earlier amending provision S.I. 2019/720, Sch. 2 para.
	64(2)(k) by S.I. 2020/1567 Sch. 2 para. 23(d)(i)
_	Art. 3(1)(aj) substituted for point (ah) the second time it occurs in earlier amending
	provision S.I. 2019/720, Sch. 2 para. 64(2)(k) by S.I. 2020/1567 Sch. 2 para. 23(d)
	(ii)
_	Art. 3(3)-(7) substituted for Art. 3(3)(4) by S.I. 2019/720 Sch. 2 para. 64(3)
_	Art. $5(1)(d)$ words substituted by S.I. 2019/720 Sch. 2 para. $65(a)$
_	Art. 6(5)(6) inserted by S.I. 2019/720 Sch. 2 para. 66(4)

Art. 8(2A) inserted by S.I. 2019/720 Sch. 2 para. 68(5) Art. 8A inserted by S.I. 2019/720 Sch. 2 para. 69 Art. 8A word substituted in earlier amending provision S.I. 2019/720, Sch. 2 para. 69 by S.I. 2020/1567 Sch. 2 para. 24 Art. 9(1)(a) words substituted by S.I. 2019/720 Sch. 2 para. 70(2)(c) Art. 9(1)(b) words substituted by S.I. 2019/720 Sch. 2 para. 70(2)(d) Art. 9(1A) inserted by S.I. 2019/720 Sch. 2 para. 70(3) Art. 12(4) inserted by S.I. 2019/720 Sch. 2 para. 73(d) Art. 14(4)(a) words substituted by S.I. 2019/720 Sch. 2 para. 75(5)(d) Art. 14(4)(b) word substituted by S.I. 2019/720 Sch. 2 para. 75(5)(e) Art. 14(4A) inserted by S.I. 2019/720 Sch. 2 para. 75(6) Art. 14(4A) word substituted in earlier amending provision S.I. 2019/720, Sch. 2 para. 75(6) by S.I. 2020/1567 Sch. 2 para. 26 Art. 14(5A) inserted by S.I. 2019/720 Sch. 2 para. 75(8) Art. 17A inserted by S.I. 2019/720, Sch. 2 para. 78A (as inserted) by S.I. 2020/1567 Sch. 2 para. 27 Art. 19(1)(a) words substituted by S.I. 2019/720 Sch. 2 para. 80(a) Art. 19(4)(a) omitted by S.I. 2019/720 Sch. 2 para. 80(b) Art. 24A inserted by S.I. 2019/720 Sch. 2 para. 85 Art. 25(1)(a) words substituted by S.I. 2019/720 Sch. 2 para. 86(a) Art. 25(1)(a) words substituted by S.I. 2019/720 Sch. 2 para. 86(b) Art. 26(2A)-(2C) inserted by S.I. 2022/1291 reg. 2(2)(a) Art. 26(3A)(3B) inserted by S.I. 2022/1291 reg. 2(2)(c) Art. 28(3)-(7) substituted for Art. 28(3)-(5) by S.I. 2019/720 Sch. 2 para. 89(c) Art. 29(1A)(1B) inserted by S.I. 2022/1291 reg. 2(3) Art. 29(2)(a) omitted by S.I. 2019/720 Sch. 2 para. 90(4)(b) Art. 29(2)(b) omitted by S.I. 2019/720 Sch. 2 para. 90(4)(b) Art. 30(1A)-(1C) inserted by S.I. 2022/1291 reg. 2(4)(b) Art. 30(2A) inserted by S.I. 2022/1291 reg. 2(4)(d) Art. 30(4) inserted by S.I. 2022/1291 reg. 2(4)(f) Art. 55(4)(d) and semicolon omitted in earlier amending provision S.I. 2019/720, Sch. 2 para. 102 by S.I. 2020/1567 Sch. 2 para. 28(a) Art. 55(7) Art. 55(9) renumbered as Art. 55(7) in earlier amending provision S.I. 2019/720, Sch. 2 para. 102 by S.I. 2020/1567 Sch. 2 para. 28(c) Art. 55(7)(8) omitted in earlier amending provision S.I. 2019/720, Sch. 2 para. 102 by S.I. 2020/1567 Sch. 2 para. 28(b) Art. 55(7) words substituted in earlier amending provision S.I. 2019/720, Sch. 2 para. 102 by S.I. 2020/1567 Sch. 2 para. 28(d) Art. 58(9) inserted by S.I. 2019/720 Sch. 2 para. 105(6) Art. 60(4)(5) inserted by S.I. 2019/720 Sch. 2 para. 107(3) Art. 60(4)(5) words substituted in earlier amending provision S.I. 2019/720, Sch. 2 para. 107(3) by S.I. 2020/1567 Sch. 2 para. 29 Art. 69(2)(c) words omitted by S.I. 2019/720 Sch. 2 para. 115(3)(a) Art. 69(2)(o) words substituted by S.I. 2019/720 Sch. 2 para. 115(3)(b) (This amendment not applied to legislation.gov.uk. Sch. 2 para. 115(3)(b) substituted immediately before IP completion day by S.I. 2020/1567, reg. 1(2), Sch. 2 para. 32) Art. 69(2)(0) words substituted by S.I. 2019/720, Sch. 2 para. 115(3)(b) (as substituted) by S.I. 2020/1567 Sch. 2 para. 32 Art. 83A83B inserted by S.I. 2019/720 Sch. 2 para. 125 Art. 83B(1) words substituted in earlier amending provision S.I. 2019/720, Sch. 2 para. 125 by S.I. 2020/1567 Sch. 2 para. 35(a) Art. 83B(4)-(7) omitted in earlier amending provision S.I. 2019/720, Sch. 2 para. 129 by S.I. 2020/1567 Sch. 2 para. 35(b) Art. 88(2) words substituted in earlier amending provision S.I. 2019/720, Sch. 2 para. 129 by S.I. 2020/1567 Sch. 2 para. 36(a) Art. 88(3)(d) omitted in earlier amending provision S.I. 2019/720, Sch. 2 para. 129 by S.I. 2020/1567 Sch. 2 para. 36(b)

-	Art. 88(6) Art. 88(8) renumbered as Art. 88(6) in earlier amending provision S.I.
	2019/720, Sch. 2 para. 129 by S.I. 2020/1567 Sch. 2 para. 36(d)
-	Art. 88(6) words substituted in earlier amending provision S.I. 2019/720, Sch. 2
	para. 129 by S.I. 2020/1567 Sch. 2 para. 36(e)
-	Art. 88(7)(8) omitted in earlier amending provision S.I. 2019/720, Sch. 2 para. 129
	by S.I. 2020/1567 Sch. 2 para. 36(c)
-	Art. 89(7) words inserted by S.I. 2022/1291 reg. 2(5)(a)
-	Art. 89(7A)-(7C) inserted by S.I. 2022/1291 reg. 2(5)(b)
-	Art. 89(8) words substituted by S.I. 2022/1291 reg. 2(5)(c)
-	Art. 89(9) words inserted by S.I. 2022/1291 reg. 2(5)(d)
-	Art. 89(9A) inserted by S.I. 2022/1291 reg. 2(5)(e)
-	Art. 89(12) inserted by S.I. 2022/1291 reg. 2(5)(f)
-	Art. 92(1A)-(1C) inserted by S.I. 2019/720 Sch. 2 para. 133
-	Art. 93(a) word substituted by S.I. 2019/720 Sch. 2 para. 134(3)(a)
-	Art. 93(a) words substituted by S.I. 2019/720 Sch. 2 para. 134(3)(b)
-	Art. 93(b) word substituted by S.I. 2019/720 Sch. 2 para. 134(4)
-	Art. 94(1)(a) words substituted by S.I. 2019/720 Sch. 2 para. 135(2)(b)
-	Art. 94(1)(a) words substituted in earlier amending S.I. 2019/720, Sch. 2 para.
	135(2)(b) by S.I. 2020/1567 Sch. 2 para. 37
-	Art. 95(8) inserted by S.I. 2019/720 Sch. 2 para. 136(6)
-	Art. 95A-95L inserted by S.I. 2019/720 Sch. 2 para. 137 (This amendment not
	applied to legislation.gov.uk. Sch. 2 para. 137 omitted immediately before IP
	completion day by virtue of S.I. 2020/1567, reg. 1(2), Sch. 2 para. 38)
-	Art. 95A-95N inserted by S.I. 2019/720, Sch. 4 para. 2 (as inserted) by S.I.
	2020/1567 Sch. 4
-	Art. 95B(4A) inserted by S.I. 2022/1291 reg. 2(6)
-	Art. 95C(4A) inserted by S.I. 2022/1291 reg. 2(7)
-	Art. 95H(4A) inserted by S.I. 2022/1291 reg. 2(9)
-	Art. 95FA and cross-heading inserted by S.I. 2022/1291 reg. 2(8)