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ANNEX II

INFORMATION REQUIREMENTS FOR ACTIVE SUBSTANCES

TITLE 1

CHEMICAL SUBSTANCES

Core data set and additional data set for active substances

Information required to support the approval of an active substance is listed in the table below.

Conditions for not requiring a specific test that are set out in the appropriate test methods in the Regulation (EC) No 440/2008 and are not repeated in column 3, also apply.

re	olumn 1Information quired	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 require recourse to testing of vertebrates
1.	APPLICANT		
1.1	. Name and address		
1.2	Contact person		
1.3	manufacturer (name, address and location of manufacturing plant(s))		
	IDENTITY OF THE CTIVE SUBSTANCE		
info Sec ena to l tec doc nec	r the active substance, the formation given in this ction shall be sufficient to able the active substance be identified. If it is not hnically possible or if it es not appear scientifically cessary to give information one or more of the items		
a	The information provided should be manufactured, if different.	for the purified active substance of stated	specification or for the active substance as
b	The information provided should be	for the purified active substance of stated	specification.
c	OJ L 20, 26.1.1980, p. 43.		
d	OJ L 372, 27.12.2006, p. 19.		
e	OJ L 348, 24.12.2008, p. 84.		

OJ L 348, 24.12.2008, p. 84.

	ow, the reasons shall be arly stated		
2.1	Common name proposed or accepted by ISO and synonyms (usual name, trade name, abbreviation)		
2.2	Chemical name (IUPAC and CA nomenclature or other international chemical name(s))		
2.3	Manufacturer's development code number(s)		
2.4	. CAS number plus EC, INDEX and CIPAC numbers		
2.5	Molecular and structural formula (including SMILES notation, if available and appropriate)		
2.6	Information on optical activity and full details of any isomeric composition (if applicable and appropriate)		
2.7	. Molar mass		
2.8	Method of manufacture (syntheses		
a	The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.		pecification or for the active substance as
b	The information provided should be	for the purified active substance of stated sp	pecification.
c	OJ L 20, 26.1.1980, p. 43.		
d	OJ L 372, 27.12.2006, p. 19.		

substa inforr startir and so includ specif	ng suppliers, cations mmercial	
of pur active manu kg, g/ v) as a provid inclus	ication ty of the substance as actured in g/ or %w/w (v/ ppropriate, ing vely the and lower	
any ir additi by-pro of syr optica degra produ substa unstal and er etc. of un-rea	hesis, isomers, ation tts (if the nce is le) un-reacted d-groups polymers and cted starting als of UVC-	
of at l repres batche active includinform	ation on	
	t of the provided should be for the purified active substance of stated specification or for the active substance	as

a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.

b The information provided should be for the purified active substance of stated specification.

c OJ L 20, 26.1.1980, p. 43.

d OJ L 372, 27.12.2006, p. 19.

e OJ L 348, 24.12.2008, p. 84.

	impurities referred to in 2.10.			
2.12.	The origin of the natural active substance or the precursor(s) of the active substance, e.g. an extract of a flower			
AND PRO	IYSICAL CHEMICAL PERTIES OF THE IVE SUBSTANCE			
3.1. A	Appearance ^a			
3.1.1.	Aggregate state (at 20 °C and 101,3 kPa)			
3.1.2.	Physical state (i.e. viscous, crystalline, powder) (at 20 °C and 101,3 kPa)			
3.1.3.	Colour (at 20 °C and 101,3 kPa)			
3.1.4.	Odour (at 20 °C and 101,3 kPa)			
3.2.	Melting/freezing point ^b			
3.3.	Acidity, alkalinity			
3.4.	Boiling point ^b			
3.5.	Relative Density ^b			
3.6.	Absorption spectra data (UV/VIS, IR, NMR) and a			
	e information provided should be nufactured, if different.	for the purified active substance of stated s	pecification or for the active substance as	
b Th	e information provided should be	for the purified active substance of stated s	pecification.	
c OJ	OJ L 20, 26.1.1980, p. 43.			
d OJ	OJ L 372, 27.12.2006, p. 19.			
e OJ	e OJ L 348, 24.12.2008, p. 84.			

		mass spectrum,		
		molar extinction		
		coefficient at relevant		
		wavelengths, where		
		relevant ^b		
3.	7. Vaj	pour pressure ^b		
3 7	7.1.	Henry's law		
٥.,		constant must		
		always be stated for		
		solids and liquids if		
		it can be calculated		
3.8	3.	Surface tension ^b		
3.9).	Water solubility ^b		
3.1	Λ	Partition coefficient		
J. 1	0.	(n-octanol/		
		water) and its pH		
		dependency ^b		
3.1	1	Thermal stability,		
J.1		identity of		
		breakdown		
		products ^b		
3.1	2.	Reactivity towards		
		container material		
2 1	2	D: : /:	ADS	
3.1	3.	Dissociation constant		
		Constant		
3.1	4.	Granulometry		
3.1	5	Viscosity	ADS	
J. 1	. J.	Viscosity	ADC	
3.1	6.	Solubility in	ADS	
		organic solvents,		
		including effect		
a		nformation provided should be factured, if different.	for the purified active substance of stated s	pecification or for the active substance as
b	The information provided should be for the purified active substance of stated specification.			
c		20, 26.1.1980, p. 43.		
d	OJ L	372, 27.12.2006, p. 19.		
e	OJ L 348, 24.12.2008, p. 84.			

	of temperature on solubility ^b	
3.17.	solvents used in biocidal products and identity of relevant breakdown products ^a	ADS
ANI	HYSICAL HAZARDS D RESPECTIVE ARACTERISTICS	
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.	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	
	The information provided should be nanufactured, if different.	for the purified active substance of stated specification or for the active substance as
b T	The information provided should be	for the purified active substance of stated specification.
c (OJ L 20, 26.1.1980, p. 43.	
d C	OJ L 372, 27.12.2006, p. 19.	
e (OJ L 348, 24.12.2008, p. 84.	

	emit flammable gases		
4.13.	Oxidising liquids		
4.14.	Oxidising solids		
4.15.	Organic peroxides		
4.16.	Corrosive to metals		
	dditional physical ors for hazards		
4.17.1.	Auto-ignition temperature (liquids and gases)		
4.17.2.	Relative self ignition temperature for solids		
4.17.3.	Dust explosion hazard		
DETEC	HODS OF CTION AND IFICATION		
relevant	Analytical methods including validation parameters for the determination of active substance as manufactured and where appropriate, for relevant residues, isomers and impurities of the active substance and additives (e.g. stabilisers) urities other than impurities this only	for the provided active relations of the land	
	a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.		

- b The information provided should be for the purified active substance of stated specification.
- c OJ L 20, 26.1.1980, p. 43.
- d OJ L 372, 27.12.2006, p. 19.
- OJ L 348, 24.12.2008, p. 84.

$\geq 1 \text{ g/}$	<u> </u>		
5.2. Analytical methods for monitoring purposes including recovery rates and the limits of quantification and detection for the active substance, and for residues thereof in/on the following where relevant			
5.2.1.	Soil		
5.2.2.	Air		
5.2.3.	Water (surface, drinking etc.) and sediment		
5.2.4.	Animal and human body fluids and tissues		
5.3.	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 substance nor articles treated with it come into contact with food-	ADS	
		for the purified active substance of stated sp	pecification or for the active substance as
		for the purified active substance of stated sp	pecification.
c O.	J L 20, 26.1.1980, p. 43.		
d O.	J L 372, 27.12.2006, p. 19.		
e O.	J L 348, 24.12.2008, p. 84.		

	producing animals, food of plant or	
	animal origin or feeding stuffs)	
AC	EFFECTIVENESS GAINST TARGET RGANISMS	
6.1.	Function, e.g. fungicide, rodenticide, insecticide, bactericide and mode of control e.g. attracting, killing, inhibiting	
6.2.	Representative organism(s) to be controlled and products, organisms or objects to be protected	
6.3.	Effects on representative target organism(s)	
6.4.	Likely concentration at which the active substance will be used in products and, where appropriate, in treated articles	
6.5.	Mode of action (including time delay)	
6.6.	Efficacy data to support these claims on biocidal	
a	The information provided should be manufactured, if different.	for the purified active substance of stated specification or for the active substance as
b	The information provided should be	for the purified active substance of stated specification.
c	OJ L 20, 26.1.1980, p. 43.	
d	OJ L 372, 27.12.2006, p. 19.	
e	OLL 348 24 12 2008 p. 84	

	products and, where label claims are made, on treated articles, including any available standard protocols, laboratory tests or field trials used including performance standards where appropriate		
	. Any known limitations efficacy		
6.7.	1. Information on the occurrence or possible occurrence of the development of resistance and appropriate management strategies		
6.7.	undesirable or unintended side- effects, e.g. on beneficial and other non-target organisms		
	NTENDED USES AND POSURE		
7.1.	Field of use(s) envisaged for biocidal products and, where appropriate, treated articles		
7.2.	Product-type(s)		
a	The information provided should be manufactured, if different.	e for the purified active substance of stated sp	pecification or for the active substance as
b	The information provided should be	e for the purified active substance of stated sp	pecification.
c	OJ L 20, 26.1.1980, p. 43.		
d	OJ L 372, 27.12.2006, p. 19.		
e	OJ L 348, 24.12.2008, p. 84.		

OJ L 348, 24.12.2008, p. 84.

7.3.	Detailed description of the intended use pattern(s) including in treated articles			
7.4.	Users 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 the envisaged major use categories			
confo	Exposure data in ormity with Annex VI s Regulation			
7.6.1.	Information on human exposure associated with the intended uses and disposal of the active substance			
7.6.2.	Information on environmental exposure associated with the intended uses and disposal of the active substance			
7.6.3.	Information on exposure of food-producing animals and food and feeding stuffs associated with the			
	The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.		pecification or for the active substance as	
b The	The information provided should be for the purified active substance of stated specification.			
	OJ L 20, 26.1.1980, p. 43.			
	*	OJ L 372, 27.12.2006, p. 19.		

	intended uses of the active substance		
7.6.4.	Information on exposure from treated articles including leaching data (either laboratory studies or model data)		
PROFI HUMA ANIM	KICOLOGICAL ILE FOR AN AND AL INCLUDING BOLISM		
endpoin according testing sirritation set out in Test Gu Toxicity Corrosio	Skin irritation or skin corrosion essment of this at shall be carried out ng to the sequential strategy for dermal and corrosion in the Appendix to ideline B.4. Acute y-Dermal Irritation/on (Annex B.4. lation (EC) No		
endpoin according testing sirritation set dow to Test C Toxicity Corrosio	Eye irritation essment of this at shall be carried out ng to the sequential strategy for eye n and corrosion as n in the Appendix Guideline B.5.Acute y: Eye Irritation/ on (Annex B.5. lation (EC) No		
8.3.	Skin sensitisation		Step 2 does not need to be conducted if:
	information provided should be afactured, if different.	for the purified active substance of stated s	pecification or for the active substance as
b The i	b The information provided should be for the purified active substance of stated specification.		
c OJ L	c OJ L 20, 26.1.1980, p. 43.		
d OJ L	d OJ L 372, 27.12.2006, p. 19.		
e OJ L	348, 24.12.2008, p. 84.		

endpoin	essment of this at shall comprise the ng consecutive steps: an assessment of the available human, animal and alternative data in vivo testing The Murine Local Lymph Node Assay (LLNA) including, where appropriate, the reduced variant of the assay, is the first-choice method for in vivo testing. If another skin sensitisation test is used justification shall be provided			the available information indicates that the substance 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.	Respiratory sensitisation	ADS		
8.5. M	utagenicity		1	
endpoin	essment of this at shall comprise the shall comprise the shall consecutive steps: an assessment of the available in vivo genotoxicity data an in vitro test for gene mutations in bacteria, an in vitro cytogenicity test in mammalian cells and an in vitro gene mutation test in mammalian cells are required appropriate in vivo genotoxicity studies shall be considered in case of a positive result in any of the			
a The i	-	for the purified active substance of stated s	pecification of	or for the active substance as
	ifactured, if different.	101 and parinted delive substance of stated s	Politication	or the deare substance ds
b The i	The information provided should be for the purified active substance of stated specification.			
c OJ L	20, 26.1.1980, p. 43.			
d OJ L	I OJ L 372, 27.12.2006, p. 19.			
e OJ L	348, 24.12.2008, p. 84.			

8.5.1. 8.5.2. 8.5.3.	in vitro genotoxicity studies In vitro gene mutation study in bacteria In vitro cytogenicity study in mammalian cells In vitro gene mutation study in mammalian cells		
endpoin	In vivo genotoxicity study essment of this at shall comprise the ng consecutive steps: If there is a positive result in any of the in vitro genotoxicity studies and there are no results available from an in vivo study already, an appropriate in vivo somatic cell genotoxicity study shall be proposed/conducted by the applicant If either of the in vitro gene mutation tests is positive, an in vivo test to investigate unscheduled DNA synthesis shall be conducted A second in vivo somatic cell test may be necessary, depending on the	ADS	The study/ies do(es) not generally need to be conducted if: — the results are negative for the three in vitro tests and if no metabolites of concern are formed in mammals or valid in vivo micronucleus data is generated within a repeat dose study and the in vivo micronucleus test is the appropriate test to be conducted to address this information requirement the substance is known to be carcinogenic category 1A or 1B or mutagenic category 1A, 1B or 2.
a The information provided should be for the purified active substance of stated specification or for the active substance as			

a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.

b The information provided should be for the purified active substance of stated specification.

c OJ L 20, 26.1.1980, p. 43.

d OJ L 372, 27.12.2006, p. 19.

e OJ L 348, 24.12.2008, p. 84.

results, quality and relevance of all the available data If there is a positive result from an in vivo somatic cell study available, the potential for germ cell mutagenicity should be considered on the basis of all available data, including toxicokinetic evidence to demonstrate that the substance reached the tested organ. If no clear conclusions about germ cell mutagenicity can be made, additional investigations shall be considered		
8.7. Acute toxicity In addition to the oral route of administration (8.7.1), for substances other than gases, the information mentioned under 8.7.2 to 8.7.3 shall be provided for at least one other route of administration — The choice for the second route will depend on the nature of the substance and the likely route of human exposure — Gases and volatile liquids should be administered by the inhalation route		The study/ies do(es) not generally need to be conducted if: — the substance is classified as corrosive to the skin
	for the purified active substance of stated sp	pecification or for the active substance as

- manufactured, if different.
- The information provided should be for the purified active substance of stated specification.
- c OJ L 20, 26.1.1980, p. 43.
- d OJ L 372, 27.12.2006, p. 19.
- OJ L 348, 24.12.2008, p. 84.

OJ L 20, 26.1.1980, p. 43. OJ L 372, 27.12.2006, p. 19. OJ L 348, 24.12.2008, p. 84.

be provide either the or inhalati is the only of exposu humans the oral test in considered a new dentoxicity stocarried out in vitro de penetratio	re is the then on for coute need ed. If dermal on route re to then an hay be d. Before mal acute tudy is t, an termal n study 28) should ted to likely e and tmal ofility y be al nces routes of ation are	
8.7.1. By oral ro The Acute Toxic Cl Method is the prefe method for the dete of this endpoint	ass rred	The study need not be conducted if: — the substance is a gas or a highly volatile substance
8.7.2. By inhalar route is appropriate exposure of humans inhalation is likely account: — the vapour of the sub	ation if s via taking into	
a The information prov manufactured, if diffe		substance of stated specification or for the active substance as
b The information prov	rided should be for the purified active	substance of stated specification.

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	volatile substance			
	has vapour pressure			
	$> 1 \times 10^{-2} \text{ Pa at } 20$			
	°C) and/or			
	the active substance			
	is a powder			
	containing			
	a significant			
	proportion (e.g.			
	1 % on a weight			
	basis) of particles			
	with particle size			
	MMAD < 50			
	micrometers or			
	the active substance			
	is included in			
	products that			
	are powders or			
	are applied in			
	a manner that			
	generates exposure			
	to aerosols, particles			
	or droplets of an			
	inhalable size			
	(MMAD < 50)			
	micrometers)			
_	the Acute Toxic			
	Class Method			
	is the preferred			
	method for the			
	determination of			
	this endpoint			
8.7.3.	By dermal route			
Testing l	by the dermal route is			
necessar	y only if:			
_	inhalation of			
	the substance is			
	unlikely, or			
	skin contact in			
	production and/or			
	use is likely, and			
	either			
	the physicochemical			
	and toxicological			
a The ir	nformation provided should be	for the purified active substance of stated s	pecification or for the active substance as	
	manufactured, if different.			

The information provided should be for the purified active substance of stated specification.

c

OJ L 20, 26.1.1980, p. 43.

OJ L 372, 27.12.2006, p. 19.

OJ L 348, 24.12.2008, p. 84.

_	properties suggest potential for a significant rate of absorption through the skin, or the results of an in vitro dermal penetration study (OECD 428) demonstrate high dermal absorption and bioavailability		
met	Toxicokinetics and tabolism studies in mmals		
The meta proverate the trelevinche meta rate	toxicokinetics and abolism studies should ide basic data about the and extent of absorption, issue distribution and the vant metabolic pathway ading the degree of abolism, the routes and of excretion and the vant metabolites		
be recourse and concern furth required	toxicokinetic and metabolism studies in mammals itional studies might equired based on the ome of the toxicokinetic metabolism study flucted in rat. These her studies shall be ired if: there is evidence that metabolism in the rat is not relevant for human exposure route-to-route extrapolation from oral to dermal/	ADS	
	The information provided should be manufactured, if different.	for the purified active substance of stated s	pecification or for the active substance as
b	The information provided should be	for the purified active substance of stated s	pecification.
c	OJ L 20, 26.1.1980, p. 43.		
d	OJ L 372, 27.12.2006, p. 19.		
e	OJ L 348, 24.12.2008, p. 84.		

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8.9. Repeated dose toxicity In general, only one route of administration is necessary and the oral route is the preferred route. However, in some cases it may be necessary to evaluate more than one route of exposure. For the evaluation of the safety of consumers in relation to active substances that may end up in food or feed, it is necessary to conduct toxicity studies by the oral route Testing by the dermal route shall be considered if: skin contact in production and/or use is likely, and inhalation of the substance is unlikely, and one of the following conditions is met: (i) toxicity is observed in an acute dermal toxicity test at lower doses than 4. The information provided should be for the purified active substance of stated specification or for the active substance as	inhalation exposure is not feasible Where it is considered appropriate to obtain information on dermal absorption, the assessment of this endpoint shall proceed using a tiered approach for assessment of dermal absorption		
manufactured, if different.	In general, only one route of administration is necessary and the oral route is the preferred route. However, in some cases it may be necessary to evaluate more than one route of exposure. For the evaluation of the safety of consumers in relation to active substances that may end up in food or feed, it is necessary to conduct toxicity studies by the oral route Testing by the dermal route shall be considered if: — skin contact in production and/or use is likely, and — inhalation of the substance is unlikely, and — one of the following conditions is met: (i) toxicity is observed in an acute dermal toxicity test at lower doses than The information provided should be	for the purified active substance of stated s	study (28 or 90 days) does not need to be conducted if: — a substance undergoes immediate disintegration and there are sufficient data on the cleavage products for systemic and local effects and no synergistic effects are expected, or — relevant human exposure can be excluded in accordance with Section 3 of Annex IV In order to reduce testing carried out on vertebrates and in particular the need for free-standing single- endpoint studies, the design of the repeated dose toxicity studies shall take account of the possibility to explore several endpoints within the framework of one study

The information provided should be for the purified active substance of stated specification.

OJ L 20, 26.1.1980, p. 43.

OJ L 372, 27.12.2006, p. 19. OJ L 348, 24.12.2008, p. 84.

c d 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. (See end of Document for details) View outstanding changes

in the oral toxicity test, or (ii) information or test data indicate dermal absorption is comparable or higher than oral absorption, or (iii) dermal toxicity is recognised for structurally related substances and for example is observed at lower doses than in the oral toxicity test or dermal absorption comparable or higher than oral absorption Testing by the inhalation route shall be considered if: exposure of humans via inhalation is likely taking into account the vapour pressure of the substance (volatile

- a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.
- **b** The information provided should be for the purified active substance of stated specification.
- **c** OJ L 20, 26.1.1980, p. 43.
- **d** OJ L 372, 27.12.2006, p. 19.

substances and

e OJ L 348, 24.12.2008, p. 84.

manufactured, if different.

OJ L 20, 26.1.1980, p. 43.

OJ L 372, 27.12.2006, p. 19.
OJ L 348, 24.12.2008, p. 84.

b

d

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_	gases have vapour pressure $> 1 \times 10^{-2}$ Pa at 20 °C), and/or there is the possibility of exposure to aerosols, particles or droplets of an inhalable size (MMAD < 50 micrometers)	
8.9.1.	Short-term repeated dose toxicity study (28 days), preferred species is rat	The short-term toxicity study (28 days) does not need to be conducted if: (i) a reliable sub- chronic (90 day) study is available, provided that the most appropriate species, dosage, solvent and route of administration were used, (ii) the frequency and duration of human exposure indicates that a longer term study is appropriate and one of the following conditions is met: — other available data indicate that the substance may have a dangerous property that cannot be detected in a short-

The information provided should be for the purified active substance of stated specification.

OJ L 348, 24.12.2008, p. 84.

8.9.2. Sub-chronic repeated dose toxicity study but which are liable to result in adverse effects after prolonged exposure 8.9.2. Sub-chronic repeated dose toxicity study 90 days) preferred species is rat The sub-chronic toxicity study (90 days), preferred species is rat The information provided should be for the purified active substance of stated specification. The information provided should be for the purified active substance of stated specification. The information provided should be for the purified active substance of stated specification. The information provided should be for the purified active substance of stated specification. The information provided should be for the purified active substance of stated specification. The information provided should be for the purified active substance of stated specification. The information provided should be for the purified active substance of stated specification.			1	I	
8.9.2. Sub-chronic repeated dose toxicity study (90 days), preferred species is rat 8.9.2. Sub-chronic repeated dose toxicity study (90 days), preferred species is rat 8.9.2. The sub-chronic toxicity study (90 days), preferred species is rat 8.9.2. The sub-chronic toxicity study (90 days), preferred species is rat 8.9.2. The sub-chronic toxicity study (90 days) does not need to be conducted iff. — a reliable short-term toxicity study (28 days) is available showing severe toxicity effects according to the criteria for classifying the substance as a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different. b The information provided should be for the purified active substance of stated specification. c OJL 20, 26.1.1980, p. 43.					
8.9.2. Sub-chronic repeated dose toxicity study (90 days) prefered species is rat 8.9.2. Sub-chronic repeated dose toxicity study (90 days), preferred species is rat 8.9.2. The sub-chronic toxicity study (90 days), preferred species is rat 8.9.2. The sub-chronic toxicity study (90 days) does not need to be conducted if: — a reliable short-term toxicity study (28 days) is available showing severe toxicity effects according to the criteria for classifying to the criteria for classifying the substance as manufactured, if different. b The information provided should be for the purified active substance of stated specification. c OJ L 20, 26.1.1980, p. 43.					
8.9.2. Sub-chronic repeated dose toxicoly study (90 days) does not need to be conducted if: — a reliable short-term toxicity study (90 days), preferred species is rat 8.9.2. The sub-chronic repeated dose toxicity study (90 days) does not need to be conducted if: — a reliable short-term toxicity study (28 days) is available showing severe toxicity effects according to the criteria for classifying the substance as a The information provided should be for the purified active substance of stated specification. b The information provided should be for the purified active substance of stated specification. c OJL 20, 26.1.1980, p. 43.					
8.9.2. Sub-chronic repeated dose toxicity study (90 days), preferred species is rat 8.9.2. Sub-chronic repeated dose toxicity study (90 days), preferred species is rat The sub-chronic toxicity study (90 days), preferred species is rat The sub-chronic toxicity study (90 days) preferred species is rat The sub-chronic toxicity study (90 days) is available showing severe toxicity study (28 days) is available showing severe toxicity effects according to the criteria for classifying the substance as manufactured, if different. The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.					
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study (90 days) does not need to be conducted if: ———————————————————————————————————				The sub-chronic	toxicity
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			for the purified active substance of state	ed specification.	
	c OJ L 20, 26.	1.1980, p. 43.			

8.9.3. Long-term rep dose toxicity (months)		H372 and H373 (Regulation (EC) No 1272/2008), for which the observed NOAEL-28 days, with the application of an appropriate uncertainty factor allows the extrapolation towards the NOAEL-90 days for the same route of exposure, and — a reliable chronic toxicity study is available, provided that an appropriate species and route of administration were used, or — the substance is unreactive, insoluble, not bioaccumulative and not inhalable and there is no evidence of absorption and no evidence of toxicity in a 28-day 'limit test', particularly if such a pattern is coupled with limited human exposure The long-term toxicity study (≥ 12 months) does not need to be conducted if: — Long-term exposure can be excluded and no effects have been seen at the limit dose in the 90-day study or
a The information provided a	should be for the purified active subs	tance of stated specification or for the active substance as
a The information provided s manufactured, if different.		
manufactured, if different.	should be for the purified active subs	tance of stated specification.
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OJ L 20, 26.1.1980, p. 43.

OJ L 372, 27.12.2006, p. 19. OJ L 348, 24.12.2008, p. 84.

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. (See end of Document for details) View outstanding changes

		a combined long- term repeated dose/ carcinogenicity study (8.11.1) is undertaken
8.9.4. Further repeat dose	ADS	
studies		
Further repeat dose studies		
including testing on a		
second species (non-rodent),		
studies of longer duration		
or through a different route		
of administration shall be		
undertaken in case of:		
no other information on		
toxicity for a second		
non-rodent species		
is provided for, or		
— failure to identify		
a no observed		
adverse effect level		
(NOAEL) in the 28-		
or the 90-day study,		
unless the reason is that no effects have		
been observed at the		
limit dose, or		
substances bearing		
positive structural		
alerts for effects		
for which the rat		
or mouse is an		
inappropriate or		
insensitive model,		
or toxicity of		
particular concern		
(e.g. serious/severe		
effects), or		
indications of an		
effect for which		
the available data		
is inadequate		
a The information provided should be manufactured, if different.	for the purified active substance of stated s	pecification or for the active substance as

The information provided should be for the purified active substance of stated specification.

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for toxicological and/or risk characterisation. In such cases it may also be more appropriate to perform specific toxicological studies that are designed to investigate these effects (e.g. immunotoxicity, neurotoxicity, hormonal activity), concern regarding local effects for which a risk characterisation cannot be performed by route-to route extrapolation, or particular concern regarding exposure (e.g. use in biocidal products leading to exposure levels which are close to the toxicologically relevant dose levels), or effects shown in substances with a clear relationship in molecular structure with the substance being studied were not detected in the 28- or the 90-day study, or the route of administration used in the initial repeated dose study

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- **c** OJ L 20, 26.1.1980, p. 43.
- **d** OJ L 372, 27.12.2006, p. 19.
- e OJ L 348, 24.12.2008, p. 84.

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was inappropriate in relation to the expected route of human exposure and route-to-route extrapolation cannot be made.	
8.10. Reproductive toxicity For evaluation of consumer safety of active substances that may end up in food or feed, it is necessary to conduct toxicity studies by the oral route	The studies need not be conducted if: — the substance is known to be a genotoxic carcinogen and appropriate risk management measures are implemented including measures related to reproductive toxicity, or — the substance is known to be a germ cell mutagen and appropriate risk management measures are implemented including measures related to reproductive toxicity, or — the substance is of low toxicological activity (no evidence of toxicity seen in any of the tests available provided that the dataset is sufficiently comprehensive and informative), it can be proven from toxicokinetic data

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b The information provided should be for the purified active substance of stated specification.

c OJ L 20, 26.1.1980, p. 43.

d OJ L 372, 27.12.2006, p. 19.

e OJ L 348, 24.12.2008, p. 84.

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that no systemic absorption occurs via relevant routes of exposure (e.g. plasma/blood concentrations below detection limit using a sensitive method and absence of the substance and of metabolites of the substance in urine, bile or exhaled air) and the pattern of use indicates there is no or no significant human exposure If a substance is known to have an adverse effect on fertility, meeting the criteria for classification as Reproductive toxicity Cat 1A or 1B: May damage fertility (H360F), and the available data are adequate to support a robust risk assessment, then no further testing for fertility will be necessary. However, testing for developmental toxicity must be considered If a substance is known to cause developmental toxicity, meeting the criteria for classification as

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- e OJ L 348, 24.12.2008, p. 84.

	Reproductive toxicity Cat 1A or 1B: May damage the unborn child (H360D), and the available data are adequate to support a robust risk assessment, then no further testing for developmental toxicity will be necessary. However, testing for effects on fertility must be considered
8.10.1. Pre-natal developmental toxicity study, preferred species is rabbit; oral route of administration is the preferred route. The study shall be initially performed on one species	
8.10.2. Two-generation reproductive toxicity study, rat, oral route of administration is the preferred route. If another reproductive toxicity test is used justification shall be provided. The extended one-generation reproductive toxicity study adopted at OECD level shall be considered as an alternative approach to the multigeneration study	

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- **e** OJ L 348, 24.12.2008, p. 84.

8.10.3. Further pre-natal developmental toxicity study. A decision on the need to perform additional studies on a second species or mechanistic studies should be based on the outcome of the first test (8.10.1) and all other relevant available data (in particular rodent reprotox studies). Preferred species is rat, oral route of administration	ADS	
8.11. Carcinogenicity See 8.11.1 for new study requirements		A carcinogenicity study does not need to be conducted if: — the substance is classified as mutagen category 1A or 1B. The default presumption would be that a genotoxic mechanism for carcinogenicity is likely. In these cases, a carcinogenicity test will normally not be required
8.11.1. Combined carcinogenicity study and long- term repeated dose toxicity Rat, oral route of administration is the preferred route. If an	for the purified active substance of stated s	

- a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.
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- **e** OJ L 348, 24.12.2008, p. 84.

a justific provided For eval safety of that may or feed,	uation of consumer f active substances end up in food it is necessary to toxicity studies by		
8.11.2.	Carcinogenicity testing in a second species A second carcinogenicity study should normally be conducted using the mouse as test species For evaluation of consumer safety of active substances that may end up in food or feed, it is necessary to conduct toxicity studies by the oral route		
8.12. Relevant health data, observations and treatments			
	tion should be I if data is not e		
8.12.1.	Medical surveillance data on manufacturing plant personnel		
8.12.2.	Direct observation, e.g. clinical cases, poisoning incidents		
a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.			
b The in	nformation provided should be	for the purified active substance of stated sp	pecification.
c OJ L 20, 26.1.1980, p. 43.			
d OJ L 372, 27.12.2006, p. 19.			
e OJ L	348, 24.12.2008, p. 84.		

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8.12.3.	Health records, both from industry and any other available sources		
8.12.4.	Epidemiological studies on the general population		
8.12.5.	Diagnosis of poisoning including specific signs of poisoning and clinical tests		
8.12.6.	Sensitisation/ allergenicity observations		
8.12.7.	Specific treatment in case of an accident or poisoning: first aid measures, antidotes and medical treatment, if known		
8.12.8.	Prognosis following poisoning		
be requi- characte use of th Other av Availabl methods includin based ris vitro and proteom	Additional studies hal data which may red depending on the ristics and intended he active substance railable data: he data from emerging and models, go toxicity pathwaysk assessment, in d'omic' (genomic, ic, metabolomic,	ADS	position for the setting substance
a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.			

The information provided should be for the purified active substance of stated specification.

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OJ L 20, 26.1.1980, p. 43. OJ L 372, 27.12.2006, p. 19. OJ L 348, 24.12.2008, p. 84.

etc.) studies, systems biology, computational toxicology, bioinformatics, and high- throughput screening shall be submitted in parallel		
8.13.1. Phototoxicity	ADS	
8.13.2. Neurotoxicity including developmental neurotoxicity	ADS	
 The preferred test species is the rat unless another test species is justified to be more 		
appropriate — For delayed neurotoxicity tests the preferred species will be the		
adult hen If anticholinesterase activity is detected a test for response to reactivating		
agents should be considered		
If the active substance is an organophosphorus compound or if there is any evidence		
e.g. knowledge of the mechanism of action or from		
repeat dose studies that the active substance may have		
neurotoxic or developmental		
neurotoxic properties then		
additional information or specific studies will be		
required.		
For evaluation of consumer		
safety of active substances		
that may end up in food		
or feed, it is necessary to	for the purified active substance of stated si	

- a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.
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- **c** OJ L 20, 26.1.1980, p. 43.
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- e OJ L 348, 24.12.2008, p. 84.

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conduct toxicity studies by		
the oral route		
8.13.3. Endocrine disruption If there is any evidence from in vitro, repeat dose or reproduction toxicity studies, that the active substance may have endocrine disrupting properties then additional information or specific studies shall be required to: elucidate the mode/ mechanism of action provide sufficient evidence for relevant adverse effects For evaluation of consumer safety of active substances that may end up in food or feed, it is necessary to conduct toxicity studies by the oral route	ADS	
8.13.4. Immunotoxicity including developmental immunotoxicity If there is any evidence, from skin sensitisation, repeat dose or reproduction toxicity studies, that the active substance may have immunotoxic properties then additional information or specific studies shall be required to: — elucidate the mode/ mechanism of action — provide sufficient evidence for	ADS	position for the setting substance.
a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.		
b The information provided should be for the purified active substance of stated specification.		

relevant adverse effects in huma. For evaluation of consursafety of active substance that may end up in food or feed, it is necessary to conduct toxicity studies the oral route.	ns ner es
8.13.5. Mechanistic data — any stu necessary to cl effects reported toxicity studies	arify I in
8.14. Studies related to the exposure of humans to the active substance.	ne e
8.15. Toxic effects o livestock and p	
stuffs st including for producing an and their pro (milk, eggs honey) Additional information	imals
related to the exposure of humans to the active substance contained in biocidal products	
8.16.1. Proposed accepresidue levels i maximum residuimits (MRL) a the justification their acceptabi	e. lue nd of

a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.

 $[\]label{eq:barrier} \textbf{b} \qquad \text{The information provided should be for the purified active substance of stated specification.}$

c OJ L 20, 26.1.1980, p. 43.

d OJ L 372, 27.12.2006, p. 19.

e OJ L 348, 24.12.2008, p. 84.

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OJ L 20, 26.1.1980, p. 43.

OJ L 372, 27.12.2006, p. 19. OJ L 348, 24.12.2008, p. 84.

8.16.2. Behaviour of the residue of the active substance on the treated or contaminated food or feeding stuffs including the kinetics of disappearance Residue definitions should be provided where relevant. It is also important to compare residues found in toxicity studies with residues formed in food-producing animals and their products, as well as food and feed	ADS	
8.16.3. Overall material balance for the active substance Sufficient residue data from supervised trials on foodproducing animals and their products, as well as food and feed, to demonstrate that residues likely to arise from the proposed use would not be of concern for human or animal health	ADS	
8.16.4. Estimation of potential or actual exposure of humans to the active substance and residues through diet and other means	ADS	
8.16.5. If residues of the active substance occur in or on feeding stuffs	ADS	
	for the purified active substance of stated s	pecification or for the active substance as
b The information provided should be	for the purified active substance of stated s	pecification.

OJ L 348, 24.12.2008, p. 84.

for a significant period of time or are found in food of animal origin after treatment on or around food-producing animals (e.g. direct treatment on animals or indirect treatment of animal houses or surroundings) then feeding and metabolism studies in livestock shall be required to permit evaluation of residues in food of animal origin		
8.16.6. Effects of industrial processing and/ or domestic preparation on the nature and magnitude of residues of the active substance	ADS	
8.16.7. Any other available information that is relevant It may be appropriate to include information on migration into food, especially in the case of treatment of food contact materials		
8.16.8. Summary and evaluation of data submitted under 8.16.1 to 8.16.8	ADS	
a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.		
b The information provided should be	for the purified active substance of stated specification.	
c OJ L 20, 26.1.1980, p. 43.		
d OJ L 372, 27.12.2006, p. 19.		

wheth found or plan as thos studies risk as	inportant to establish er the metabolites in food (from animals ints) are the same se tested in toxicity s. Otherwise values for issessment (e.g. ADI) t valid for the residues		
8.17.	If the active substance is to be used in products for action against plants including algae then tests shall be required to assess toxic effects of metabolites from treated plants, if any, where different from those identified in animals	ADS	
and co to all t and ar	Summary of mammalian toxicology de overall evaluation onclusion with regard toxicological data my other information rating the active ences including NOAEL		
9.1. T	TOXICOLOGICAL DIES Toxicity to Aquatic nisms	-	
9.1.1.	Short-term toxicity testing on fish short-term fish toxicity		The study does not need to be conducted if: — a valid long-term
	required the threshold		aquatic toxicity
a Th			
b Th	The information provided should be for the purified active substance of stated specification.		
c OJ	OJ L 20, 26.1.1980, p. 43.		
d OJ	OJ L 372, 27.12.2006, p. 19.		
e OJ	OJ L 348, 24.12.2008, p. 84.		

approach (tiered strategy) should be applied		study on fish is available	
9.1.2. Short-term toxicity testing on aquatic invertebrates			
9.1.2.1. Daphnia magna			
9.1.2.2. Other species	ADS		
9.1.3. Growth inhibition study on algae			
9.1.3.1. Effects on growth rate of green algae			
9.1.3.2. Effects on growth rate of cyanobacteria or diatoms			
9.1.4. Bioconcentration		The experimental determination may not need	
9.1.4.1. Estimation methods 9.1.4.2. Experimental determination		to be carried out if: — it can be demonstrated on the basis of physico- chemical properties (e.g. log Kow < 3) or other evidence that the substance has a low potential for bioconcentration	
9.1.5. Inhibition of microbial activity The study may be replaced by a nitrification inhibition test if available data show that the substance is likely to be an inhibitor of microbial growth or function, in particular nitrifying bacteria			
a The information provided should be manufactured, if different.	for the purified active substance of stated s	specification or for the active substance as	
b The information provided should be	The information provided should be for the purified active substance of stated specification.		
c OJ L 20, 26.1.1980, p. 43.			
d OJ L 372, 27.12.2006, p. 19.	OJ L 372, 27.12.2006, p. 19.		
e OJ L 348, 24.12.2008, p. 84.			

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		ADS	
9.1.6.	Further Toxicity Studies on Aquatic Organisms		
If the res	sults of the		
	ological studies,		
	on fate and behaviour ne intended use(s)		
	etive substance		
	a risk for the aquatic		
	nent, or if long- posure is expected,		
	or more of the tests		
	d in this Section shall		
be condu	ucted		
9.1.6.1.		ADS	
(a)	testing on Fish Fish Early Life		
, ,	Stage (FELS) Test		
(b)	Fish short term toxicity test on		
	embryo and sack fry		
	stages		
(c)	Fish juvenile growth test		
(d)	Fish full life cycle		
	test		
9.1.6.2.	Long term toxicity	ADS	
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	testing on		
(2)	invertebrates		
(a)	Daphnia growth and reproduction study		
(b)	Other species		
	reproduction and		
(c)	growth (e.g. Mysid) Other species		
(-)	development and		
	emergence (e.g.		
	Chironomus)	A D G	
9.1.7.	Bioaccumulation	ADS	
	in an appropriate		
	aquatic species		
	nformation provided should be factured, if different.	for the purified active substance of stated s	pecification or for the active substance as
	eformation provided about he		

The information provided should be for the purified active substance of stated specification.

OJ L 20, 26.1.1980, p. 43.

OJ L 372, 27.12.2006, p. 19.

OJ L 348, 24.12.2008, p. 84.

c d

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9.1.8.	Effects on any other specific, non-target organisms (flora and fauna) believed to be at risk	ADS	
9.1.9.	Studies on sediment- dwelling organisms	ADS	
9.1.10.	Effects on aquatic macrophytes	ADS	
9.2.	Terrestrial toxicity, initial tests	ADS	
9.2.1.	Effects on soil micro-organisms		
9.2.2.	Effects on earthworms or other soil- dwelling non- target invertebrates		
9.2.3.	Acute toxicity to plants		
9.3.	Terrestrial tests, long term	ADS	
9.3.1.	Reproduction study with earthworms or other soil- dwelling non-target invertebrates		
9.4.	Effects on birds	ADS	For endpoint 9.4.3 the study does not need to be
9.4.1.	Acute oral toxicity		conducted if: — the dietary toxicity
9.4.2.	Short-term toxicity — eight-		study shows that the LC ₅₀ is above 2 000 mg/kg
	nformation provided should be factured, if different.	for the purified active substance of stated s	pecification or for the active substance as
b The is	nformation provided should be	for the purified active substance of stated s	pecification.
c OJ L	20, 26.1.1980, p. 43.		
d OJ L	372, 27.12.2006, p. 19.		
e OJ L	348, 24.12.2008, p. 84.		

	day dietary study in at least one species (other than chickens, ducks and geese)		
9.4.3.	Effects on reproduction		
9.5.	Effects on arthropods	ADS	
9.5.1.	Effects on honeybees		
9.5.2.	Other non- target terrestrial arthropods, e.g. predators		
9.6.	Bioconcentration, terrestrial	ADS	
9.7.	Bioaccumulation, terrestrial	ADS	
9.8.	Effects on other non-target, non-aquatic organisms	ADS	
9.9.	Effects on mammals	ADS	Data are derived from the mammalian toxicological
9.9.1.	Acute oral toxicity		assessment. The most sensitive relevant mammalian
9.9.2.	Short term toxicity		long-term toxicological endpoint (NOAEL) expressed
9.9.3.	Long term toxicity		as mg test compound/kg bw/ day shall be reported
9.9.4.	Effects on reproduction		
9.10.	Identification of endocrine activity	ADS	
a The	information provided should be	for the purified active substance of stated s	pecification or for the active substance as

a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.

b The information provided should be for the purified active substance of stated specification.

c OJ L 20, 26.1.1980, p. 43.

d OJ L 372, 27.12.2006, p. 19.

e OJ L 348, 24.12.2008, p. 84.

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10. ENVIRONMENTAL FATE AND BEHAVIOUR

10.1. Fate and behaviour in water and sediment

10.1.1. Degradation, initial studies

If the assessment performed indicates the need to investigate further the degradation of the substance and its degradation products or the active substance has an overall low or absent abiotic degradation, then the tests described in 10.1.3 and 10.3.2 and where appropriate — in 10.4 shall be required. The choice of the appropriate test(s) depends on the results of the initial assessment performed	
10.1.1.1. Abiotic	
 (a) Hydrolysis as a function of pH and identification of breakdown products The identification of breakdown products is 	
required when the breakdown products at any sampling time are present at \geq 10 %	
(b) Phototransformation in water, including identification of transformation products	
10.1.1.2. Biotic	

- **a** The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.
- **b** The information provided should be for the purified active substance of stated specification.
- **c** OJ L 20, 26.1.1980, p. 43.
- **d** OJ L 372, 27.12.2006, p. 19.
- e OJ L 348, 24.12.2008, p. 84.

(a)	Ready biodegradability		
(b)	Inherent biodegradability (where appropriate)		
10.1.2	Adsorption/ desorption		
route inclu- of me	3. Rate and confidence of degradation ding identification etabolites and adation products		
10.1.3	.1. Biological sewage treatment		
(a)	Aerobic biodegradation	ADS	
(b)	Anaerobic biodegradation	ADS	
(c)	STP simulation test	ADS	
10.1.3	.2. Biodegradation in freshwater		
(a)	Aerobic aquatic degradation study	ADS	
(b)	Water/sediment degradation test	ADS	
10.1.3	.3. Biodegradation in sea water	ADS	
10.1.3	.4. Biodegradation during manure storage	ADS	
	e information provided should be unufactured, if different.	for the purified active substance of stated s	pecification or for the active substance as
b Th	e information provided should be	for the purified active substance of stated s	pecification.
c OJ	L 20, 26.1.1980, p. 43.		
d OJ	L 372, 27.12.2006, p. 19.		
e OJ	L 348, 24.12.2008, p. 84.	-	

10.1.4.	Adsorption and desorption in water/ aquatic sediment systems and, where relevant, adsorption and desorption of metabolites and degradation products	ADS	
10.1.5.	Field study on accumulation in sediment	ADS	
10.1.6.	Inorganic substances: information on fate and behaviour in water	ADS	
10.2.	Fate and behaviour in soil	ADS	
of degra	Laboratory study on rate and route of degradation including identification of the processes involved and identification of any metabolites and degradation products in one soil type (unless pH dependent route) under appropriate conditions by studies on rate dation in three al soil types	ADS	
10.2.2.	Field studies, two soil types	ADS	
a The information provided should be for the purified active substance of stated specification or for the active substance as			

a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.

b The information provided should be for the purified active substance of stated specification.

c OJ L 20, 26.1.1980, p. 43.

d OJ L 372, 27.12.2006, p. 19.

e OJ L 348, 24.12.2008, p. 84.

10.2.3.	Soil accumulation studies	ADS	
10.2.4.	Adsorption and desorption in at least three soil types and, where relevant, adsorption and desorption of metabolites and degradation products	ADS	
10.2.5.	Further studies on sorption		
10.2.6.	Mobility in at least three soil types and where relevant mobility of metabolites and degradation products	ADS	
10.2.6.1.	Column leaching studies		
10.2.6.2.	Lysimeter studies		
10.2.6.3	Field leaching studies		
character residues	Extent and nature of bound residues rmination and ristics of bound is recommended mbined with a soil on study	ADS	
10.2.8.	Other soil degradation studies	ADS	
a The information provided should be for the purified active substance of stated specification or for the active substance as			

a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.

b The information provided should be for the purified active substance of stated specification.

c OJ L 20, 26.1.1980, p. 43.

d OJ L 372, 27.12.2006, p. 19.

e OJ L 348, 24.12.2008, p. 84.

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OJ L 372, 27.12.2006, p. 19.
OJ L 348, 24.12.2008, p. 84.

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10.2.9.	Inorganic substances: information on fate and behaviour in soil		
10.3. Fain air	ate and behaviour		
10.3.1. Identific transform	Phototransformation in air (estimation method) ation of mation products		
10.3.2.	Fate and behaviour in air, further studies	ADS	
10.4.	Additional studies on fate and behaviour in the environment	ADS	
10.5.	Definition of the residue	ADS	
10.5.1.	Definition of the residue for risk assessment		
10.5.2.	Definition of the residue for monitoring		
10.6.	Monitoring data	ADS	
10.6.1.	Identification of all degradation products (> 10 %) must be included in the studies on degradation in soil, water and sediments		
	nformation provided should be factured, if different.	for the purified active substance of stated sp	pecification or for the active substance as
		for the purified active substance of stated sp	pecification.
c OJ L 20, 26.1.1980, p. 43.			

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11. MEASURES
NECESSARY TO
PROTECT HUMANS,
ANIMALS AND THE
ENVIRONMENT

manufactured, if different.

OJ L 20, 26.1.1980, p. 43.

OJ L 372, 27.12.2006, p. 19.

OJ L 348, 24.12.2008, p. 84.

b

c

d

11.1.	Recommended methods and precautions concerning handling, use, storage, transport or fire		
11.2.	In case of fire, nature of reaction products, combustion gases etc.		
11.3.	Emergency measures in case of accident		
(a) (b) (c)	Possibility of destruction or decontamination following release in or on the following: air water, including drinking water soil		
11.5.	Procedures for waste management of the active substance for industry or professional users		
11.6.	Possibility of reuse or recycling		
a The in	a The information provided should be for the purified active substance of stated specification or for the active substance as		

The information provided should be for the purified active substance of stated specification.

11.7. Possibility of neutralisation of effects 11.8. Conditions for controlled discharge including leachate qualities on disposal 11.9. Conditions for controlled incineration 11.10. Identification of any substances falling within the scope of List I or List II of the Annex to Council Directive 80/68/EEC of 17 December 1979 on the protection of groundwater against pollution caused by certain dangerous substances', of Annexes I and II to Directive 2006/118/ EC of the European Parliament and of the Council of 12 December 2006 on the protection of groundwater against pollution and deterioration', of Annex I to Directive 2008/105/ EC of the European Parliament and of the Council of 16 Directive 2008/105/ EC of the European Parliament and of the Council of 16 December 2008 on environmental and of the Council of 16 December 2008 on environmental and the council of 12 Dece			
controlled discharge including leachate qualities on disposal 11.9. Conditions for controlled incineration 11.10. Identification of any substances falling within the scope of List I or List II of the Annex to Council Directive 80/68/EEC of 17 December 1979 on the protection of groundwater against pollution caused by certain dangerous substances ⁴ , of Annexes I and II to Directive 2006/118/EC of the European Parliament and of the Council of 12 December 2006 on the protection of groundwater against pollution and deterioration ⁴ , of Annex I to Directive 2008/105/EC of the European Parliament and of the Council of 16 Directive 2008/105/EC of the European Parliament and of the Council of 16 December 2008 on environmental	11.7.	neutralisation of	
for controlled incineration 11.10. Identification of any substances falling within the scope of List I or List II of the Annex to Council Directive 80/68/EEC of 17 December 1979 on the protection of groundwater against pollution caused by certain dangerous substances ^c , of Annexes I and II to Directive 2006/118/EC of the European Parliament and of the Council of 12 December 2006 on the protection of groundwater against pollution and deterioration ^d , of Annex I to Directive 2008/105/EC of the European Parliament and of the Council of 16 December 2008 on environmental	11.8.	controlled discharge including leachate	
substances falling within the scope of List I or List II of the Annex to Council Directive 80/68/EEC of 17 December 1979 on the protection of groundwater against pollution caused by certain dangerous substances', of Annexes I and II to Directive 2006/118/ EC of the European Parliament and of the Council of 12 December 2006 on the protection of groundwater against pollution and deterioration ⁴ , of Annex I to Directive 2008/105/ EC of the European Parliament and of the Council of 16 December 2008 on environmental	11.9.	for controlled	
quanty standards in	11.10.	substances falling within the scope of List I or List II of the Annex to Council Directive 80/68/EEC of 17 December 1979 on the protection of groundwater against pollution caused by certain dangerous substances ^c , of Annexes I and II to Directive 2006/118/EC of the European Parliament and of the Council of 12 December 2006 on the protection of groundwater against pollution and deterioration ^d , of Annex I to Directive 2008/105/EC of the European Parliament and of the Council of 16 December 2008	

a The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.

b The information provided should be for the purified active substance of stated specification.

c OJ L 20, 26.1.1980, p. 43.

d OJ L 372, 27.12.2006, p. 19.

e OJ L 348, 24.12.2008, p. 84.

	the field of water policy ^e , of Part B of Annex I to Directive 98/83/EC or Annexes VIII and X to Directive 2000/60/EC		
LABEI	ASSIFICATION, LLING AND AGING		
12.1.	State any existing classification and labelling		
classific substar from th	he hazard cation of the ace resulting a application alation (EC) No		
In addition, for each entry, the reasons why no classification is given for an endpoint should be provided			
12.2.1.	Hazard classification		
12.2.2.	Hazard pictogram		
12.2.3.	Signal word		
12.2.4.	Hazard statements		
12.2.5.	Precautionary statements including prevention, response, storage and disposal		
12.3.	Specific concentration		
	nformation provided should be factured, if different.	for the purified active substance of s	tated specification or for the active substance as
b The in	nformation provided should be	for the purified active substance of s	tated specification.
c OJ L	20, 26.1.1980, p. 43.		
	372, 27.12.2006, p. 19.		
e OJ L	348, 24.12.2008, p. 84.		

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limits, where applicable, resulting from the application of Regulation (EC) No 1272/2008			
13. SUMMARY AND EVALUATION The key information identified from the endpoints in each subsection (2-12) is summarised, evaluated and a draft risk assessment is performed			
The information provided should be for the purified active substance of stated specification or for the active substance as manufactured, if different.			
1 TI ' C (' '11 1 111			

- **b** The information provided should be for the purified active substance of stated specification.
- **c** OJ L 20, 26.1.1980, p. 43.
- **d** OJ L 372, 27.12.2006, p. 19.
- **e** OJ L 348, 24.12.2008, p. 84.

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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

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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)
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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)

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    Annex 6 para. 66 words substituted by S.I. 2019/720 Sch. 2 para. 143(11)
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- 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)(o) 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)