Commission Directive 2008/44/EC of 4 April 2008 amending Council Directive 91/414/EEC to include benthiavalicarb, boscalid, carvone, fluoxastrobin, Paecilomyces lilacinus and prothioconazole as active substances (Text with EEA relevance)

COMMISSION DIRECTIVE 2008/44/EC

of 4 April 2008

amending Council Directive 91/414/EEC to include benthiavalicarb, boscalid, carvone, fluoxastrobin, *Paecilomyces lilacinus* and prothioconazole as active substances

(Text with EEA relevance)

THE COMMISSION OF THE EUROPEAN COMMUNITIES,

Having regard to the Treaty establishing the European Community,

Having regard to Council Directive 91/414/EEC of 15 July 1991 concerning the placing of plant protection products on the market⁽¹⁾, and in particular Article 6(1) thereof,

Whereas:

- (1) In accordance with Article 6(2) of Directive 91/414/EEC Belgium received on 19 April 2002 an application from Kumiai Chemicals Industry Co. Ltd for the inclusion of the active substance benthiavalicarb in Annex I to Directive 91/414/EEC. Commission Decision 2003/35/EC⁽²⁾ confirmed that the dossier was 'complete' in the sense that it could be considered as satisfying, in principle, the data and information requirements of Annexes II and III to Directive 91/414/EEC.
- (2) In accordance with Article 6(2) of Directive 91/414/EEC Germany received on 26 April 2001 an application from BASF AG for the inclusion of the active substance boscalid in Annex I to Directive 91/414/EEC. Commission Decision 2002/268/EC⁽³⁾ confirmed that the dossier was 'complete' in the sense that it could be considered as satisfying, in principle, the data and information requirements of Annexes II and III to Directive 91/414/EEC.
- (3) In accordance with Article 6(2) of Directive 91/414/EEC the Netherlands received on 26 March 1997 an application from Luxan B.V. for the inclusion of the active substance carvone in Annex I to Directive 91/414/EEC. Commission Decision 1999/610/EC⁽⁴⁾ confirmed that the dossier was 'complete' in the sense that it could be considered as satisfying, in principle, the data and information requirements of Annexes II and III to Directive 91/414/EEC.
- (4) In accordance with Article 6(2) of Directive 91/414/EEC United Kingdom received on 25 March 2002 an application from Bayer AG for the inclusion of the active substance fluoxastrobin in Annex I to Directive 91/414/EEC. Decision 2003/35/EC confirmed that the dossier was 'complete' in the sense that it could be considered as satisfying,

- in principle, the data and information requirements of Annexes II and III to Directive 91/414/EEC.
- (5) In accordance with Article 6(2) of Directive 91/414/EEC Belgium received on 15 September 2002 an application from Prophyta for the inclusion of the active substance *Paecilomyces lilacinus* strain 251 (hereafter *Paecilomyces lilacinus*) in Annex I to Directive 91/414/EEC. Commission Decision 2003/305/EC⁽⁵⁾ confirmed that the dossier was 'complete' in the sense that it could be considered as satisfying, in principle, the data and information requirements of Annexes II and III to Directive 91/414/EEC.
- (6) In accordance with Article 6(2) of Directive 91/414/EEC United Kingdom received on 25 March 2002 an application from Bayer CropScience for the inclusion of the active substance prothioconazole in Annex I to Directive 91/414/EEC. Decision 2003/35/EC confirmed that the dossier was 'complete' in the sense that it could be considered as satisfying, in principle, the data and information requirements of Annexes II and III to Directive 91/414/EEC.
- (7) For those active substances, the effects on human health and the environment have been assessed, in accordance with the provisions of Article 6(2) and (4) of Directive 91/414/EEC, for the uses proposed by the applicants. The designated rapporteur Member States submitted draft assessment report on 13 April 2004 (benthiavalicarb), 22 November 2002 (boscalid), 16 October 2000 (carvone), 2 September 2003 (fluoxastrobin), 3 November 2004 (*Paecilomyces lilacinus*) and 18 October 2004 (prothioconazole).
- (8) The assessment reports were peer reviewed by the Member States and the EFSA within its Working Group Evaluation and presented to the Commission in the format of the EFSA Scientific Reports on 15 June 2007 for fluoxastrobin⁽⁶⁾ and *Paecilomyces lilacinus*⁽⁷⁾ and on 12 July for benthiavalicarb⁽⁸⁾ and prothioconazole⁽⁹⁾. These reports and the draft assessment reports for boscalid and carvone were reviewed by the Member States and the Commission within the Standing Committee on the Food Chain and Animal Health and the review was finalised on 22 January 2008 in the format of the Commission review reports for benthiavalicarb, boscalid, carvone, fluoxastrobin, *Paecilomyces lilacinus* and prothioconazole.
- (9) It has appeared from the various examinations made that plant protection products containing the active substances concerned may be expected to satisfy, in general, the requirements laid down in Article 5(1) (a) and (b) and Article 5(3) of Directive 91/414/EEC, in particular with regard to the uses which were examined and detailed in the Commission review report. It is therefore appropriate to include benthiavalicarb, boscalid, carvone, fluoxastrobin, *Paecilomyces lilacinus* and prothioconazole in Annex I to that Directive, in order to ensure that in all Member States the authorisations of plant protection products containing these active substances may be granted in accordance with the provisions of that Directive.
- (10) Without prejudice to the above conclusion, for fluoxastrobin and prothioconazole it is appropriate to obtain further information on certain specific points. Article 6(1) of Directive 91/414/EEC provides that inclusion of a substance in Annex I may be subject to conditions. Therefore, it is appropriate to require that fluoxastrobin should be subjected to further testing for confirmation of the risk assessment for surface water and

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for non-rat metabolites and that prothioconazole should be subjected to further testing for confirmation of the risk assessment as regards the triazole metabolite derivatives and the risk to granivorous birds and mammals and that such studies should be presented by the notifiers.

- (11) Without prejudice to the obligations defined by Directive 91/414/EEC as a consequence of including an active substance in Annex I, Member States should be allowed a period of six months after inclusion to review existing provisional authorisations of plant protection products containing benthiavalicarb, boscalid, carvone, fluoxastrobin, *Paecilomyces lilacinus* or prothioconazole to ensure that the requirements laid down by Directive 91/414/EEC, in particular in its Article 13 and the relevant conditions set out in Annex I, are satisfied. Member States should transform existing provisional authorisations into full authorisations, amend them or withdraw them in accordance with the provisions of Directive 91/414/EEC. By derogation from the above deadline, a longer period should be provided for the submission and assessment of the complete Annex III dossier of each plant protection product for each intended use in accordance with the uniform principles laid down in Directive 91/414/EEC.
- (12) It is therefore appropriate to amend Directive 91/414/EEC accordingly.
- (13) The measures provided for in this Directive are in accordance with the opinion of the Standing Committee on the Food Chain and Animal Health,

HAS ADOPTED THIS DIRECTIVE:

Article 1

Annex I to Directive 91/414/EEC is amended as set out in the Annex to this Directive.

Article 2

1 Member States shall adopt and publish by 31 January 2009 at the latest the laws, regulations and administrative provisions necessary to comply with this Directive. They shall forthwith communicate to the Commission the text of those provisions and a correlation table between those provisions and this Directive.

They shall apply those provisions from 1 February 2009.

When Member States adopt those provisions, they shall contain a reference to this Directive or shall be accompanied by such a reference on the occasion of their official publication. Member States shall determine how such reference is to be made.

2 Member States shall communicate to the Commission the text of the main provisions of national law which they adopt in the field covered by this Directive.

Article 3

Member States shall in accordance with Directive 91/414/EEC, where necessary, amend or withdraw existing authorisations for plant protection products containing benthiavalicarb, boscalid, carvone, fluoxastrobin, *Paecilomyces lilacinus* or prothioconazole as active substance by 31 January 2009. By that date, they shall in particular verify that the conditions in Annex I to that Directive relating to benthiavalicarb, boscalid, carvone, fluoxastrobin, *Paecilomyces lilacinus* or prothioconazole, respectively, are met, with the exception of those identified in part B of the entry concerning the active substance, and that the

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holder of the authorisation has, or has access to, a dossier satisfying the requirements of Annex II to that Directive in accordance with the conditions of Article 13(2) of that Directive.

By way of derogation from paragraph 1, for each authorised plant protection product containing benthiavalicarb, boscalid, carvone, fluoxastrobin, Paecilomyces lilacinus or prothioconazole as either the only active substance or as one of several active substances all of which were listed in Annex I to Directive 91/414/EEC by 31 July 2008 at the latest, Member States shall re-evaluate the product in accordance with the uniform principles provided for in Annex VI to Directive 91/414/EEC, on the basis of a dossier satisfying the requirements of Annex III to that Directive and taking into account part B of the entry in Annex I to that Directive concerning benthiavalicarb, boscalid, carvone, fluoxastrobin, Paecilomyces lilacinus or prothioconazole. On the basis of that evaluation, they shall determine whether the product satisfies the conditions set out in Article 4(1)(b), (c), (d) and (e) of Directive 91/414/EEC.

Following that determination Member States shall:

- in the case of a product containing benthiavalicarb, boscalid, carvone, fluoxastrobin, Paecilomyces lilacinus or prothioconazole as the only active substance, where necessary, amend or withdraw the authorisation by 31 January 2010 at the latest; or
- in the case of a product containing benthiavalicarb, boscalid, carvone, fluoxastrobin, Paecilomyces lilacinus or prothioconazole as one of several active substances, where necessary, amend or withdraw the authorisation by 31 January 2010 or by the date fixed for such an amendment or withdrawal in the respective Directive or Directives which added the relevant substance or substances to Annex I to Directive 91/414/EEC, whichever is the latest.

Article 4

This Directive shall enter into force on 1 August 2008.

Article 5

This Directive is addressed to the Member States.

Done at Brussels, 4 April 2008.

For the Commission Androulla VASSILIOU Member of the Commission

ANNEX

In Annex I to Directive 91/414/EEC, the following rows are added at the end of the table:

No	Common name, identification	IUPAC name on	Purity ^a	Entry into force	Expiration of inclusion	Specific provisions
'169	Benthiavalic CAS No 413615-35-7 CIPAC No 744	{[(R)-1-(6- fluoro-1,3-	lacerbamic toxicological concern and each of them must not exceed a certain amount in the technical material: 6,6 diff dib < 3 mg kg bis am fluctors	/- luoro-2,2'- enzothiazole: ,5 / (2- ino-5- orophenyl) ulfide: 4	31 July 2018	Part A Only uses as fungicide may be authorised. Part B For the implementation of the uniform principles of Annex VI, the conclusions of the review report on benthiavalicart and in particular Appendices I and II thereof, as finalised in the Standing Committee on the Food

a Further details on identity and specification of active substances are provided in the review report.

			Chain
			and
			Animal
			Health
			on 22
			January
			2008
			shall
			be
			taken
			into
			account.
			In this
			overall
			assessment
			Member
			States
			must pay
			particular
			attention to:
			— the
			operator
			safety,
			— the
			protection
			of
			non-
			target
			arthropods.
			Conditions
			of use shall
			include
			adequate
			risk
			mitigation
			measures, where
			appropriate. In assessing
			applications
			to authorise
			plant
			protection
			products
			containing
			benthiavalicarb
			for uses
			other
			than in
			glasshouses,
			Member

a Further details on identity and specification of active substances are provided in the review report.

						States shall pay particula attention the criter in Article 4(1)(b), a shall ensith that any necessar data and informat is provid before such an authorisa is grante. The Member States shinform the Commissin accordar with Art 13(5) on the specificate of the technical material commerce manuface.	r to to ria e and ure y ion led ation d. all ne sion ace icle tion
170	Boscalid CAS No 188425-85-6 CIPAC No 673	2- Chloro- <i>N</i> - (4'- chlorobipher yl)nicotinam	≥ 960 g/kg nyl-2- ide	1 August 2008	31 July 2018	Part A Part B	Only uses as fungicide may be authorised. For the implementation of the uniform principles of Annex VI,

a Further details on identity and specification of active substances are provided in the review report.

				the
				conclusions
				of
				the
				review
				report
				on
				boscalid,
				and
				in
				particular
				Appendices
				I
				and II
				thereof,
				as
				finalised
				in
				the
				Standing
				Committee
				on
				the
				Food
				Chain
				and
				Animal Health
				on
				22
				January
				2008
				shall
				be
				taken
				into
				account.
			In this	
			overall	
			assessme Member	ent
			States	
			must pay	•
			particular	r
			attention	
			-	to
				the
				operator
				safety,
				to
				the

 $^{{\}bf a} \qquad \text{Further details on identity and specification of active substances are provided in the review report.}$

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						Condition of use shinclude adequate risk mitigation measures where appropri	all on S,
171	Carvone CAS No 99-49-0 (d/ 1 mixture) CIPAC No 602	5- isopropenyl- methylcyclol en-1-one	≥ 930 g/kg 2with a d/l heatið-of at least 100:1	1 August 2008	31 July 2018	Part A Part B	Only uses as plant growth regulator may be authorised.
				.,			the

Further details on identity and specification of active substances are provided in the review report.

			implementation
			of
			the
			uniform
			principles
			of
			Annex
			VI,
			the
			conclusions
			of
			the
			review
			report
			on
			carvone,
			and
			in
			particular
			Appendices
			I
			and
			II c
			thereof,
			as finalized
			finalised
			in the
			Standing
			Committee
			on
			the
			Food
			Chain
			and
			Animal
			Health
			on
			22
			January
			2008
			shall
			be
			taken
			into
			account.
			In this
			overall
			assessment
			Member States
			States
			must pay

a Further details on identity and specification of active substances are provided in the review report.

						particular attention the open safety. Condition of use sinclude mitigati measure where appropri	n to rator ons hall risk on es,
172	Fluoxastrobi CAS No 361377-29-9 CIPAC No 746	n(E)-{2- [6-(2-) chloropheno fluoropyrimi yloxy]pheny (5,6- dihydro-1,4, dioxazin-3- yl)methanon O- methyloxime	din-4- 1} 2- e	1 August 2008	31 July 2018	Part B	Only uses as fungicide may be authorised. For the implementation of the uniform principles of Annex VI, the conclusions of the review report on fluoxastrobin, and in particular Appendices I and II thereof, as finalised in the Standing Committee

a Further details on identity and specification of active substances are provided in the review report.

				on
				the
				Food
				Chain
				and
				Animal
				Health
				on
				22
				January
				2008
				shall
				be
				taken
				into
			In this	account.
			In this overall	
			assessme	nt
			Member	111
			States	
			must pay	
			particular	•
			attention	to:
				the
				operator
				safety,
				in
				particular
				when
				handling
				the
				undiluted
				concentrate. Conditions
				of
				use
				shall
				include
				adequate
				protective
				measures,
				such
				as
				wearing
				a
				face
				shield,
				the .
				protection
				of
 	 			aquatic

a Further details on identity and specification of active substances are provided in the review report.

			organisms. Risk
			mitigation
			measures
			such
			as
			buffer
			zones
			shall
			be
			applied, where
			appropriate,
			 the
			levels
			of
			residues
			of
			the
			metabolites
			of
			fluoxastrobin, when
			straw
			from
			treated
			areas
			is
			used
			as
			animal
			feeding
			stuff. Conditions
			of
			use
			shall
			include
			restrictions
			for
			feeding
			to
			animals, where
			appropriate,
			 the
			risk
			of
			accumulation
			in
			the
			soil

a Further details on identity and specification of active substances are provided in the review report.

			surface,
			if
			the
			substance
			is
			used
			in
			perennial
			crops
			or
			in
			succeeding
			crops
			in
			crop
			rotation.
			Conditions
			of use shall
			include risk
			mitigation
			measures,
			where
			appropriate.
			The
			concerned
			Member
			States shall
			request the
			submission
			of:
			— data
			to
			allow
			a
			comprehensive
			aquatic
			risk
			assessment
			to
			be
			made
			taking
			into
			account
			spray
			drift,
			run-
			off,
			draine as
			drainage
			and
			the
			effectiveness

a Further details on identity and specification of active substances are provided in the review report.

					They sha ensure the notificat whose request fluoxastr has been included this Anniprovide such studies to the Commission within two year from the entry into force of Directive	atier robin in ex sion s the
173	Paecilomyce	sNot	1 August	31 July		e of
-10	lilacinus (Thom) Samson 1974 strain 251 (AGAL:	applicable	2008	2018		uses as nematicide may be authorised.

a Further details on identity and specification of active substances are provided in the review report.

Lar	T	ı		D . D	Б.
No				Part B	For
89/030550)					the
CIPAC No					implementation
753					of
					the
					uniform
					principles
					of
					Annex
					VI, the
					conclusions
					of
					the
					review
					report
					on
					Paecilomyces
					lilacinus,
					and
					in
					particular
					Appendices
					I
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					II
					thereof,
					finalised
					in
					the
					Standing
					Committee
					on
					the
					Food
					Chain
					and
					Animal
					Health
					on 22
					January 2008
					shall
					be
					taken
					into
					account.
				In this	
				overall	
				assessme	ent

a Further details on identity and specification of active substances are provided in the review report.

174	Prothioconaz	:(oReS)-2-	≥ 970 g/kg	1 August		Member States must pay particula attention — Condition of use shinclude r mitigatic measures where appropri	r to: the operator safety (although there was no need to set an AOEL, as a general rule, microorganisms should be considered as potential sensitisers), the protection of leaf dwelling non- target arthropods. ns all risk on ss,
1/4	CAS No	[2-(1- chlorocyclop (2- chloropheny hydroxyprop	The ntopld)wing manufacturin Dinapurities	2008 ng	2018	гап А	only uses as fungicide may be authorised.

a Further details on identity and specification of active substances are provided in the review report.

triazole-3-	and each of			Part B	For
thione	them must				the
	not exceed				implementation
	a certain				of
	amount				the
	in the				uniform
	technical				principles
	material:				of
		uene:			Annex
	< 5				VI,
	kg	8			the
	Pro	thioconazole-			conclusions
		thio			of
	(2-				the
	(1-				review
		orocycloprop	v1)1_		report
	(2-		y1)1-		on
		orophenyl)-3-			prothioconazole,
			•		and
	(1,	2,4- zol-1-			
					in portioular
	yl)				particular
		pan-2-			Appendices I
	ol)				-
		,5 g/			and
	kg)D)			11
	(LC	DD)			thereof,
					as
					finalised
					in
					the
					Standing
					Committee
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					Animal
					Health
					on
					22
					January
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					shall
					be
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					into
					account.
				In this	
				overall	
				assessme	nt
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			States	
			must pay	
			particula	r
			particula	<u>.</u>
			attention	
				the
				operator
				safety
				in
				spray
				applications.
				Conditions
				Conditions
				of
				use
				shall
				include
				adequate
				protective
				measures,
				the
				protection
				of
				aquatic
				organisms.
				Risk
				mitigation
				measures
				such
				as
				buffer
				zones
				shall
				be
				applied,
				where
				appropriate,
				the
				protection
				of
				of binds
				birds
				and
				small
				mammals.
				Risk
				mitigation
				measures
				shall
				be
				applied,
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				appropriate.
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 	 	 	of use sh	all_

a Further details on identity and specification of active substances are provided in the review report.

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			mitigatio	
			measures	
			where	,
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			The	
			concerne	d
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			request tl	
			submissi	on
			of:	
				information
				to
				allow
				the
				assessment
				of
				consumer
				exposure
				to
				triazole
				metabolite
				derivatives
				in
				primary
				crops,
				rotational
				crops,
				and
				products
				of
				animal
				origin,
				a ammariaan
				comparison of
				the
				mode
				of
				action
				of
				prothioconazole
				and
				the
				triazole
				metabolite
				derivatives
				to
				allow
				the
				assessment

a Further details on identity and specification of active substances are provided in the review report.

			of
			the
			toxicity
			resulting
			from
			the
			combined
			exposure
			to
			these
			compounds,
			— information
			to
			further
			address
			the
			long-
			term
			risk
			to
			granivorous
			birds
			and
			mammals
			arising
			from
			the
			use
			of
			prothioconazole
			as
			a 1
			seed
			treatment.
			They shall
			ensure that
			the notifier
			at whose
			request
			prothioconazala
			prothioconazole
			has been
			included in
			this Annex
			provide
			such
			studies
			to the
			Commission
			within
			two years
			from the
 	 	 	entry into

 $^{{\}bf a} \qquad \text{Further details on identity and specification of active substances are provided in the review report.}$

						force of the Directive of inclusion.'
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a Further details on identity and specification of active substances are provided in the review report.

- OJ L 230, 19.8.1991, p. 1. Directive as last amended by Commission Directive 2008/41/EC (OJ L 89, 1.4.2008, p. 12).
- (2) OJ L 11, 16.1.2003, p. 52.
- (**3**) OJ L 92, 9.4.2002, p. 34.
- (4) OJ L 242, 14.9.1999, p. 29.
- (**5**) OJ L 112, 6.5.2003, p. 10.
- (6) EFSA Scientific Report (2007) 102, 1-84, Conclusion regarding the peer review of the pesticide risk assessment of the active substance fluoxastrobin (finalised: 13 June 2007).
- (7) EFSA Scientific Report (2007) 103, 1-35, Conclusion regarding the peer review of the pesticide risk assessment of the active substance Paecilomyces lilacinus strain 251 (finalised: 13 June 2007).
- (8) EFSA Scientific Report (2007) 107, 1-81, Conclusion regarding the peer review of the pesticide risk assessment of the active substance benthiavalicarb (finalised: 12 July 2007).
- (9) EFSA Scientific Report (2007) 106, 1-98, Conclusion regarding the peer review of the pesticide risk assessment of the active substance prothioconazole (finalised: 12 July 2007).