Commission Implementing Regulation (EU) 2017/2327 of 14 December 2017 approving 2-methyl-1,2-benzisothiazol-3(2H)-one as an active substance for use in biocidal products of product-type 6 (Text with EEA relevance)

COMMISSION IMPLEMENTING REGULATION (EU) 2017/2327

of 14 December 2017

approving 2-methyl-1,2-benzisothiazol-3(2H)-one as an active substance for use in biocidal products of product-type 6

(Text with EEA relevance)

THE EUROPEAN COMMISSION,

Having regard to the Treaty on the Functioning of the European Union,

Having regard to Regulation (EU) No 528/2012 of the European Parliament and of the Council of 22 May 2012 concerning the making available on the market and use of biocidal products⁽¹⁾, and in particular Article 90(2) thereof,

Whereas:

- (1) Poland received on 26 November 2009 an application, in accordance with Article 11(1) of Directive 98/8/EC of the European Parliament and of the Council⁽²⁾, for the inclusion of the active substance 2-methyl-1,2-benzisothiazol-3(2H)-one in Annex I to that Directive for use in products of product-type 6, in-can preservatives, as described in Annex V to that Directive, which corresponds to product-type 6 as described in Annex V to Regulation (EU) No 528/2012.
- Poland submitted the assessment report together with its recommendations on 24 March 2016 in accordance with Article 90(2) of Regulation (EU) No 528/2012.
- (3) The opinion of the European Chemicals Agency was formulated on 27 June 2017 by the Biocidal Products Committee, having regard to the conclusions of the evaluating competent authority.
- (4) According to that opinion, biocidal products of product-type 6 containing 2-methyl-1,2-benzisothiazol-3(2H)-one may be expected to satisfy the criteria of Article 19(1)(b) of Regulation (EU) No 528/2012, provided that certain specifications and conditions concerning their use are complied with.
- (5) It is therefore appropriate to approve 2-methyl-1,2-benzisothiazol-3(2H)-one for use in biocidal products of product-type 6, subject to compliance with certain specifications and conditions.
- (6) Since 2-methyl-1,2-benzisothiazol-3(2H)-one meets the criteria for classification as a skin sensitiser sub-category 1A (strong sensitiser) as specified in point 3.4.2.2.1.2 of Annex I to Regulation (EC) No 1272/2008 of the European Parliament

- and of the Council⁽³⁾, treated articles treated with or incorporating 2-methyl-1,2-benzisothiazol-3(2H)-one should be labelled appropriately when placed on the market.
- (7) A reasonable period should be allowed to elapse before an active substance is approved in order to permit interested parties to take the preparatory measures necessary to meet the new requirements.
- (8) The measures provided for in this Regulation are in accordance with the opinion of the Standing Committee on Biocidal Products,

HAS ADOPTED THIS REGULATION:

Article 1

2-methyl-1,2-benzisothiazol-3(2H)-one is approved as an active substance for use in biocidal products of product-type 6, subject to the specifications and conditions set out in the Annex.

Article 2

This Regulation shall enter into force on the twentieth day following that of its publication in the *Official Journal of the European Union*.

This Regulation shall be binding in its entirety and directly applicable in all Member States.

Done at Brussels, 14 December 2017.

For the Commission

The President

Jean-Claude JUNCKER

ANNEX

Common Name	IUPAC NameIdent Numbers	Minimum ifilegien of purity of the active substance	Date of approval	Expiry date of approval	Product type	Specific conditions
2-methyl-1,2-benzisothiaz one (MBIT)	IUPAC Name: o2-3(2H)- methyl-1,2- benzisothiaz one EC No: Not available CAS No: 2527-66-4	≥ 997 g/kg ol-3(2H)-	1 July 2018	30 June 2028	6	The authorisations of biocidal products are subject to the following conditions: (1) the product assessment shall pay particular attention to the exposures, the risks and the efficacy linked to any use covered by an application for authorisation, but not addressed in the Union-level risk assessment

a The purity indicated in this column was the minimum degree of purity of the active substance evaluated. The active substance in the product placed on the market can be of equal or different purity if it has been proven to be technically equivalent to the evaluated active substance.

				of	
				the	
				active	
				substance	2;
			(2)	in	
				view	
				of	
				the	
				risks	
				identified	1
				for	
				the	
				uses	
				assessed,	
				the	
				product	
				assessme	nt
				shall	
				pay	
				particular	r
				attention	
				to:	
				(a)	professional
					users;
				(b)	surface
					water
					and
					groundwater
					for
					the
					outdoor
					use
					of
					preserved
					paints
					and
					plasters
					and
					the
					use
					of
					preserved
					fluids
					used
					in
					paper,
					textile
					or
					leather
					production.

The purity indicated in this column was the minimum degree of purity of the active substance evaluated. The active substance in the product placed on the market can be of equal or different purity if it has been proven to be technically equivalent to the evaluated active substance.

1	1		1	
				The placing
				on the
				market
				of treated
				articles
				is subject
				to the
				following
				condition:
				The person
				responsible
				for the
				placing on
				the market
				of a treated
				article
				treated
				with or
				incorporating
				2-
				methyl-1,2-
				benzisothiazol-3(2H)-
				one shall
				ensure that
				the label of
				that treated
				article
				provides
				the
				information
				listed in
				the second
				subparagraph
				of Article
				58(3) of
				Regulation
				(EU) No
				528/2012.
1	1	1		

a The purity indicated in this column was the minimum degree of purity of the active substance evaluated. The active substance in the product placed on the market can be of equal or different purity if it has been proven to be technically equivalent to the evaluated active substance.

- **(1)** OJ L 167, 27.6.2012, p. 1.
- (2) Directive 98/8/EC of the European Parliament and of the Council of 16 February 1998 concerning the placing of biocidal products on the market (OJ L 123, 24.4.1998, p. 1).
- (3) Regulation (EC) No 1272/2008 of the European Parliament and of the Council of 16 December 2008 on classification, labelling and packaging of substances and mixtures, amending and repealing Directives 67/548/EEC and 1999/45/EC, and amending Regulation (EC) No 1907/2006 (OJ L 353, 31.12.2008, p. 1).