Commission Regulation (EC) No 2380/2001 of 5 December 2001 concerning the 10 year authorisation of an additive in feedingstuffs (Text with EEA relevance)

## COMMISSION REGULATION (EC) No 2380/2001

of 5 December 2001

## concerning the 10 year authorisation of an additive in feedingstuffs

(Text with EEA relevance)

## THE COMMISSION OF THE EUROPEAN COMMUNITIES,

Having regard to the Treaty establishing the European Community,

Having regard to Council Directive 70/524/EEC of 23 November 1970 concerning additives in feedingstuffs<sup>(1)</sup>, as last amended by Directive 2001/46/EC of the European Parliament and of the Council<sup>(2)</sup>, and in particular Article 4 thereof,

Whereas:

- (1) Article 2(aaa) of Directive 70/524/EEC requires authorisations for coccidiostats to be linked to the person responsible for putting them into circulation.
- (2) Article 9 of Directive 70/524/EEC provides that a substance may be authorised if all conditions laid down in Article 3a of that Directive are met.
- (3) The assessment of the dossier submitted shows that the coccidiostat described in the Annex satisfies all the requirements of Article 3a of Directive 70/524/EEC, when used for the animal category and under the conditions described in the Annex to this Regulation: the substance should therefore be authorised under those conditions.
- (4) Article 9b of Directive 70/524/EEC provides that the authorisations of such substances shall be given for a period of 10 years from the date on which the final authorisation takes effect.
- (5) The assessment of the dossier shows that certain procedures may be required to protect workers from exposure to the additives. Such protection should however be assured by the application of Council Directive 89/391/EEC of 12 June 1989 on the introduction of measures to encourage improvements in the safety and health of workers at work<sup>(3)</sup>.
- (6) The Scientific Committee for Animal Nutrition has delivered a favourable opinion with regard to the safety and with regard to the favourable effect on animal production of the coccidiostat under the conditions described in the said Annex.
- (7) The measures provided for in this Regulation are in accordance with the opinion of the Standing Committee for Feedingstuffs,

HAS ADOPTED THIS REGULATION:

Status: Point in time view as at 31/12/2020. Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EC) No 2380/2001. (See end of Document for details)

#### Article 1

The additive belonging to the 'Coccidiostats and other medicinal substances' listed in the Annex to the present Regulation is authorised for use as additive in animal nutrition under the conditions laid down in that Annex.

Article 2

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

It shall apply from 15 December 2001.

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

Status: Point in time view as at 31/12/2020. Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EC) No 2380/2001. (See end of Document for details)

# [<sup>F1</sup>ANNEX

#### **Textual Amendments**

**F1** Substituted by Commission Implementing Regulation (EU) No 406/2011 of 27 April 2011 amending Regulation (EC) No 2380/2001 as regards the composition of the feed additive maduramicin ammonium alpha (Text with EEA relevance).

additive of a	f the older f uthori	name) sation	chemica formula descrip analytic method	alor a, categor ti <b>of</b> i, ca <b>l</b> nimal	age		nce/kg plete gstuff re		End period of authorisation
Coccidiosta E 770 [ <sup>F2</sup> Be SA	Zoetis elgium	Madurar ammoniu alpha 1 g/100 g (Cygro 1 %)	nicin	Autoritieiyse composit Maduran ammoniu alpha: 1 g/100 g	tivoneeks nicin um methylce e: e: e: nicin um 17N 1-5,	5 Ilulose	5	1. 'This feedings: contains an ionophor simultan use with certain medicina substanc (e.g. tiamulin)	re: eous Il es

#### Status: Point in time view as at 31/12/2020. Changes to legislation: There are currently no known outstanding effects for the Commission Regulation (EC) No 2380/2001. (See end of Document for details)

polyether	can be
monocarboxylic	contraindicated'
acid	
produced	
by	
a	
fermentation	
process	
by	
the	
strain	
Actinomadura	
yumaensis	
(ATCC	
31585)	
(NRRL	
12515)	
Related	
impurities:	
Maduramicin	
ammonium	
β:	
<	
10	
%	

#### **Textual Amendments**

F2 Substituted by Commission Implementing Regulation (EU) No 1014/2013 of 22 October 2013 amending Regulations (EC) No 2380/2001, (EC) No 1289/2004, (EC) No 1455/2004, (EC) No 1800/2004, (EC) No 600/2005, (EU) No 874/2010, Implementing Regulations (EU) No 388/2011, (EU) No 532/2011 and (EU) No 900/2011 as regards the name of the holder of the authorisation of certain additives in animal feed (Text with EEA relevance).

- (1) OJ L 270, 14.12.1970, p. 1.
- (2) OJ L 234, 1.9.2001, p. 55.
- (**3**) OJ L 183, 29.6.1989, p. 1.

## Status:

Point in time view as at 31/12/2020.

### Changes to legislation:

There are currently no known outstanding effects for the Commission Regulation (EC) No 2380/2001.