

# Official controls of veterinary drug residues in products of animal origin


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## Tables & Figures

Table 1 Group A: Prohibited or unauthorised pharmacologically active substances in food-producing animals	
Description	Substance
<b>A.1</b> Substances with hormonal and thyrostatic action and beta-agonists prohibited under Council Directive 96/22/EC	Stilbenes Antithyroid agents Steroids Resorcylic acid lactones, including zeranol Beta-agonists
<b>A.2</b> Prohibited substances listed in Regulation (EU) No 37/2010 (Annex, Table 2)	Chloramphenicol Nitrofurans Dimetridazole, metronidazole, ronidazole and other nitro-imidazoles "Other substances", i.e.: <ul style="list-style-type: none"> <li>• <i>Aristolochia</i> spp. and preparations thereof</li> <li>• Chloroform</li> <li>• Chlorpromazine</li> <li>• Colchicine</li> <li>• Dapsone</li> </ul>
<b>A.3</b> Pharmacologically active substances	Dyes Plant protection products as defined in Regulation (EU) No 1107/2009 and biocides as defined in Regulation (EU) No 528/2012 which may be used in animal husbandry of food-producing animals Antimicrobial substances Coccidiostats, histomonostats and other antiparasitic agents Protein and peptide hormones Anti-inflammatory substances, sedatives and any other pharmacologically active substances Antiviral substances

Source: Delegated Regulation (EU) [2022/1644](#), Annex I

Table 2 Group B: Pharmacologically active substances authorised for use in food-producing animals	
Until 14 December 2022 <sup>a</sup>	From 15 December 2022
Antibacterial substances including sulphonamides, quinolones	<b>B.1</b> Pharmacologically active substances listed in Regulation (EU) No 37/2010 (Annex, Table 1). MRLs are set for some of these. Antimicrobial substances Insecticides, fungicides, anthelmintics and other antiparasitic agents Sedatives Non-steroidal anti-inflammatory drugs, corticosteroids and glucocorticoids Other pharmacologically active substances
Other veterinary drugs Anthelmintics Anticoccidials, including nitroimidazoles Carbamates and pyrethroids Sedatives Non-steroidal anti-inflammatory drugs Other pharmacologically active substances	<b>B.2</b> Coccidiostats and histomonostats authorised with maximum levels and maximum residue limits
Other substances and environmental contaminants Organochlorine compounds including PCBs Organophosphorus compounds Chemical elements Mycotoxins Dyes	
a Directive 96/23/EC, Annex I.	
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Source: Delegated Regulation (EU) [2022/1644](#), Annex I


Table 3 Official controls of veterinary drug residues in products of animal origin	
New Regulation 2022/1644/EU	Old Directive 96/23/EC
<b>Group A:</b> prohibited and unauthorised pharmacologically active substances	<b>Group A:</b> prohibited substances
<b>Group B:</b> authorised pharmacologically active substances	<b>Group B:</b> veterinary drugs and contaminants
Specific plan for contaminants (2022/931 and 2022/932)	
Specific plan for plant protection products (2021/1355)	
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Table 4 Legislation based on the Official Controls Regulation <sup>1</sup>			
New rules apply from 15 Dec 2022 (1 Jan 2023 for contaminants) <sup>2</sup>			
New legislation	Control plans/programmes for exporting countries <sup>3</sup> are required for:		
	Pharmacologically active substances and their residues	Pesticide residues	Contaminants in food
	Risk-based, surveillance and import control	Coordinated multi-annual control programme	Food of animal origin and other foods
Delegated Act <sup>4</sup>	2022/1644	2021/2244	2022/931
Implementing Act <sup>5</sup>	2022/1646	2021/1355	2022/932

1 Official Controls Regulation (EU) 2017/625 (Art. 19).  
 2 New rules replace Directive 96/23/CE.  
 3 Control plans/programmes based on Regulation 2022/2292.  
 4 Delegated Acts supplement or amend non-essential elements of the basic law, and are adopted after consulting expert groups.  
 5 Implementing Acts aim to ensure uniform conditions for implementation, and are adopted with the agreement of Member States (a qualified majority) through a procedure called "comitology".

  
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