Status: There are multiple versions of this provision on screen. These apply to different geographical extents. Skip to: E+W+S - England, Wales and Scotland extentN.I. - Northern Ireland extent Changes to legislation: There are currently no known outstanding effects for the The Veterinary Medicines Regulations 2013, Paragraph 23. (See end of Document for details)

#### SCHEDULE 1

# Marketing authorisations [F1in Great Britain][F2in Northern Ireland]

#### **Textual Amendments**

- F1 Words in Sch. 1 heading inserted (E.W.S.) (31.12.2020) by The Veterinary Medicines and Residues (Amendment) (EU Exit) Regulations 2020 (S.I. 2020/1461), regs. 1(2)(b), 4(7)(a)
- **F2** Words in Sch. 1 heading inserted (N.I.) (31.12.2020) by The Animals (Health, Identification, Trade and Veterinary Medicines) (Amendment) (EU Exit) Regulations (Northern Ireland) 2020 (S.R. 2020/353), regs. 1(3), 10(13)(a)

# PART 3

## Grant of a marketing authorisation

### Marketing authorisations for food-producing species E+W+S

- **23.**—(1) The Secretary of State must not grant a marketing authorisation for a veterinary medicinal product for food-producing species unless [F3 maximum residue limits have been established under Regulation (EC) No 470/2009 of the European Parliament and of the Council in respect of [all its pharmacologically active substances F4....
- (2) This does not apply in the case of a marketing authorisation for a veterinary medicinal product for administration to a horse that has been declared on its horse passport as not intended for slaughter for human consumption; but in this case the product must not include an active substance that [F5has been classified under Article 14 of Regulation (EC) No 470/2009 of the European Parliament and of the Council as prohibited for use in food producing animals.]

### **Extent Information**

E1 This version of this provision extends to England and Wales and Scotland only; a separate version has been created for Northern Ireland only

#### **Textual Amendments**

- F3 Words in Sch. 1 para. 23(1) inserted (E.W.S.) (31.12.2020) by The Veterinary Medicines and Animals and Animal Products (Examination of Residues and Maximum Residue Limits) (Amendment etc.) (EU Exit) Regulations 2019 (S.I. 2019/676), regs. 1(2)(b), 3(20)(a)(i) (as amended by S.I. 2020/1461, regs. 1(2)(a), 3(2)(b)); 2020 c. 1, Sch. 5 para. 1(1)
- F4 Words in Sch. 1 para. 23(1) omitted (E.W.S.) (31.12.2020) by virtue of The Veterinary Medicines and Animals and Animal Products (Examination of Residues and Maximum Residue Limits) (Amendment etc.) (EU Exit) Regulations 2019 (S.I. 2019/676), regs. 1(2)(b), 3(20)(a)(ii) (as amended by S.I. 2020/1461, regs. 1(2)(a), 3(2)(b)); 2020 c. 1, Sch. 5 para. 1(1)
- Words in Sch. 1 para. 23(2) substituted (E.W.S.) (31.12.2020) by The Veterinary Medicines and Animals and Animal Products (Examination of Residues and Maximum Residue Limits) (Amendment etc.) (EU Exit) Regulations 2019 (S.I. 2019/676), regs. 1(2)(b), 3(20)(b) (as amended by S.I. 2020/1461, regs. 1(2) (a), 3(2)(b)); 2020 c. 1, Sch. 5 para. 1(1)

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Status: There are multiple versions of this provision on screen. These apply to different geographical extents. Skip to: E+W+S - England, Wales and Scotland extentN.I. - Northern Ireland extent Changes to legislation: There are currently no known outstanding effects for the The Veterinary Medicines Regulations 2013, Paragraph 23. (See end of Document for details)

# Marketing authorisations for food-producing species N.I.

- **23.**—(1) The Secretary of State must not grant a marketing authorisation for a veterinary medicinal product for food-producing species unless all its pharmacologically active substances appear in Table 1 in the Annex to Commission Regulation (EU) No 37/2010.
- (2) This does not apply in the case of a marketing authorisation for a veterinary medicinal product for administration to a horse that has been declared on its horse passport as not intended for slaughter for human consumption; but in this case the product must not include an active substance that appears in Table 2 in the Annex to Commission Regulation (EU) No 37/2010 and must not be intended for the treatment of a condition for which a veterinary medicinal product is already authorised for horses.

#### **Extent Information**

E2 This version of this provision extends to Northern Ireland only; a separate version has been created for England and Wales and Scotland only

### **Status:**

There are multiple versions of this provision on screen. These apply to different geographical extents.

# Skip to:

- E+W+S England, Wales and Scotland extent
- N.I. Northern Ireland extent

# **Changes to legislation:**

There are currently no known outstanding effects for the The Veterinary Medicines Regulations 2013, Paragraph 23.