

Medicines Act 1968

1968 CHAPTER 67

PART II

LICENCES AND CERTIFICATES RELATING TO MEDICINAL PRODUCTS

General provisions and exemptions

6 The licensing authority.

- (1) For the purposes of this Part of this Act the authority responsible for the grant, renewal, variation, suspension and revocation of licences and certificates shall be [^{F1}a body consisting of the Ministers].
- (2) Any function conferred on the licensing authority by or under this Act may be performed [^{F2}by either of the Ministers acting alone or both of them acting jointly].
- (3) In accordance with the preceding provisions of this section, in this Act "the licensing authority" means [^{F3}either or both of the] Ministers, and, in the case of anything falling to be done by the licensing authority, means [^{F3}either or both of the] Ministers acting as mentioned in subsection (2) of this section.

Textual Amendments

- **F1** Words in s. 6(1) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 6(a) (with regs. 2(4), 3)
- F2 Words in s. 6(2) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 6(b) (with regs. 2(4), 3)
- **F3** Words in s. 6(3) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 6(c) (with regs. 2(4), 3)

Modifications etc. (not altering text)

- C1 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C2 S. 6 applied (with modifications) (3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.
 S. 6 extended (with modifications) (14.2.1994) by S.I. 1994/105, reg. 19, Sch.4

S. 6 amendment to earlier affecting provision SI 1994/105 Sch. 4 (30.10.2005) by Medicines **C3** (Homoeopathic Medicinal Products for Human Use) Amendment Regulations 2005 (S.I. 2005/2753), regs. 1(1), 19(2)

7 General provisions as to dealing with medicinal products.

(1) The following provisions of this section shall have effect subject to—

- any exemption conferred by or under this Part of this Act; (a)
- ^{F4}(b)
 - the provisions of section 48 of this Act. (c)
- (2) Except in accordance with a licence granted for the purposes of this section (in this Act referred to as a "product licence") no person shall, in the course of a business carried on by him, and in circumstances to which this subsection applies,-
 - (a) sell, supply or export any medicinal product, or
 - (b) procure the sale, supply or exportation of any medicinal product, or
 - procure the manufacture or assembly of any medicinal product for sale, supply (c) or exportation.

- (3) No person shall import any medicinal product except in accordance with a product licence.
- $[^{F6}(3A)$ The restrictions imposed by subsections (2) and (3) of this section shall not apply where the medicinal product concerned is an investigational medicinal product within the meaning of the Clinical Trials Regulations.]
- $[^{F7}(3B)$ The restrictions imposed by subsections (2) and (3) of this section shall not apply where the medicinal product concerned is a homoeopathic medicinal product to which the 2001 Directive applies and which fulfils the conditions laid down in Article 14(1)of that Directive.]
 - (4) In relation to an imported medicinal product, subsection (2) of this section applies to circumstances in which the person selling, supplying or exporting the medicinal product in question, or procuring the sale, supply or exportation or the manufacture or assembly for sale, supply or exportation of that product, has himself imported the product or procured its importation.
 - (5) In relation to any medicinal product which has not been imported, subsection (2) of this section applies to any circumstances in which the person selling, supplying or exporting the medicinal product in question, or procuring the sale, supply or exportation or the manufacture or assembly for sale, supply or exportation of that product,
 - [^{F8}(a) is responsible for the composition of the product, or
 - if that product is a proprietary medicinal product $[^{F10F11}...$ or an industrially produced medicinal product $^{F11}...]$, is responsible for the placing of the product ^{F9}(b) on the market in the United Kingdom.]]
 - (6) For the purposes of subsection (5) of this section a person shall be taken to be responsible for the composition of a medicinal product if (but only if) in the course of a business carried on by him-

Changes to legislation: Medicines Act 1968, Part II is up to date with all changes known to be in force on or before 27 June 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details)

- (a) he procures the manufacture of the product to his order by another person, where the order specifies, or incorporates by reference to some other document, particulars of the composition of the product ordered, whether those particulars amount to a complete specification or not, or
- (b) he manufactures the product otherwise than in pursuance of an order which fulfils the conditions specified in the preceding paragraph.

[^{F12}(6A) ^{F13}... subsection (5)(b) of this section shall not apply if the product is—

- ^{F14}(a)
 - (b) a radiopharmaceutical in which the radionuclide is in the form of a sealed source, F15 ...

^{F15}(c) ...

 $[^{F17}(7)]_{F18}$ In this section—

F19[F20 ...

"homoeopathic medicinal product" means any medicinal product (which may contain a number of principles) prepared from F21 ..., substances F21 ... called homoeopathic stocks in accordance with a homoeopathic manufacturing procedure described by the European Pharmacopoeia or, in the absence thereof, by any pharmacopoeia used officially in a member State;]

"proprietary medicinal product" means a ready-prepared medicinal product placed on the market in the United Kingdom under a special name and in a special pack;

"radiopharmaceutical" means a medicinal product which, when ready for use, contains one or more radionuclides included for a medicinal purpose; ^{F22}...] ^{F23}...^{F23}...^{F24F23}...]

Textual Amendments

- **F4** S. 7(1)(b) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 7(a) (with regs. 2(4), 3)
- F5 S. 7(2A)(2B) repealed (30.10.2005) by Medicines (Homoeopathic Medicinal Products for Human Use) Amendment Regulations 2005 (S.I. 2005/2753), reg. 1(1), Sch. para. 1(2)
- **F6** S. 7(3A) inserted (1.5.2004) by Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, **Sch. 10 para. 3**
- F7 S. 7(3B) inserted (30.10.2005) by Medicines (Homoeopathic Medicinal Products for Human Use) Amendment Regulations 2005 (S.I. 2005/2753), reg. 1(1), Sch. para. 1(3)
- F8 Words substituted by (E.W.)(S.) S.I. 1977/1050, art. 2(2) and (N.I.) S.R. 1977 No. 170, reg. 3
- **F9** S. 7(5)(*b*) substituted by S.I. 1983/1724, art. 2(2)
- F10 Words in S. 7(5)(b) substituted (3.4.1992) by S.I. 1992/604, regs. 2(2), 4
- **F11** Words in s. 7(5)(b) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 7(b) (with regs. 2(4), 3)
- F12 S. 7(6A)(6B) inserted (3.4.1992) by S.I. 1992/604, regs. 2(3), 4
- **F13** Words in s. 7(6A) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 7(c) (with regs. 2(4), 3)
- F14 S. 7(6A)(a) omitted (8.11.2005) by virtue of The Blood Safety and Quality Regulations 2005 (S.I. 2005/50), regs. 1(2), 25(1)(a) (with reg. 2(2)-(4))
- F15 In s. 7(6A) para.(c) and word

"or" omitted (13.2.1994) by S.I.1994/276, reg.3(3)(b) (which S.I. revoked and replaced defective S.I. 1994/101 before the latter S.I. came into force) F16 S. 7(6B) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 7(d) (with regs. 2(4), 3) F17 S. 7(7) substituted by S.I. 1983/1724, art. 2(3) Words in s. 7(7) substituted (3.4.1992) by S.I. 1992/604, regs. 2(4), 4 F18 F19 Words in s. 7 repealed (30.10.2005) by Medicines (Homoeopathic Medicinal Products for Human Use) Amendment Regulations 2005 (S.I. 2005/2753), reg. 1(1), Sch. para. 1(4)(a) F20 Definitions in s. 7(7) inserted (13.2.1994) by S.I. 1994/276, reg. 3(4) (which S.I. revoked and replaced defective S.I. 1994/101 before the latter S.I. came into force) F21 Words in s. 7 repealed (30.10.2005) by Medicines (Homoeopathic Medicinal Products for Human Use) Amendment Regulations 2005 (S.I. 2005/2753), reg. 1(1), Sch. para. 1(4)(b) F22 Word in s. 7(7) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 7(e)(i) (with regs. 2(4), 3) F23 Words in s. 7(7) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 7(e)(ii) (with regs. 2(4), 3) Words in s. 7(7) repealed (3.4.1992) by S.I. 1992/604, regs. 2(5), 4 F24 Modifications etc. (not altering text) C4 Pt. II(ss. 6-50) extended with modifications by S.I. 1985/1403, art. 3(1) C5 S. 7 excluded by S.I. 1989/2325, art. 2(1) **C6** S. 7 excluded (11.12.1992) by S.I. 1992/2844, art. 2 S. 7 excluded (31.12.1994) by S.I. 1994/2986, reg.3(1) S. 7 excluded (1.1.1995) by S.I. 1994/3142, reg. 18(1) S. 7 excluded (1.1.1995) by S.I. 1994/3144, reg.9(2) **C7** S.7 excluded by S.I. 1981/164, art. 3 S. 7 excluded (30.10.2005) by Medicines (Traditional Herbal Medicinal Products for Human Use) **C**8 Regulations 2005 (S.I. 2005/2750), regs. 1(a), 10(1) (with Sch. 6) **C9** S. 7 amendment to earlier affecting provision SI 1994/3144 reg. 9 (30.10.2005) by Medicines (Marketing Authorisations Etc.) Amendment Regulations 2005 (S.I. 2005/2759), regs. 1(a), 2(12) C10 S. 7 excluded (19.8.2010) by The Medicines for Human Use (Advanced Therapy Medicinal Products and Miscellaneous Amendments) Regulations 2010 (S.I. 2010/1882), regs. 1(1), 2 C11 S. 7(1)(a)(2)(4)(5)(6) applied (with modifications) (3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.

8 Provisions as to manufacture and wholesale dealing.

- (1) The following provisions of this section shall have effect without prejudice to the operation of section 7 of this Act, but subject to the exemptions and provisions referred to in paragraphs [^{F25}(a) and (c)] of subsection (1) of that section.
- (2) [^{F26}Subject to [^{F27}subsections (2A) and (2C)] of this section]No person shall, in the course of a business carried on by him, [^{F28}manufacture, assemble or import from a third country] any medicinal product except in accordance with a licence granted for the purposes of this subsection (in this Act referred to as a "manufacturer's licence").
- [^{F29}(2A) In the case of a medicinal product that is an investigational medicinal product, the restrictions imposed by subsection (2) of this section only apply—
 - (a) if the product has a product licence or marketing authorization, and
 - (b) to the extent that the manufacture or assembly of the product is in accordance with the terms and conditions of that licence or authorization.]

[^{F29}(2B) In subsection (2A) of this section—

"investigational medicinal product" has the meaning given by the Clinical Trials Regulations; and

"marketing authorization" means—

- (a) a marketing authorization issued by a competent authority in accordance with $[{}^{F30}\mbox{the 2001 Directive}]$, or
- (b) a marketing authorization granted by the European Commission under Council Regulation (EEC) 2309/93 [^{F31}or under Regulation (EC) No 726/2004 of the European Parliament and of the Council laying down Community procedures for the authorisation and supervision of medicinal products for human and veterinary use and establishing a European Medicines Agency].]
- [^{F32}(2C) The prohibition in subsection (2) does not apply to a person who, in connection with the importation of a medicinal product from a third country—
 - (a) provides facilities solely for transporting the product, or
 - (b) in the course of a business carried on by him as an import agent, imports the medicinal product solely to the order of another person who holds a manufacturer's licence authorising the importation of the product.]
- [^{F32}(2D) The Ministers may prescribe requirements (either generally or in relation to a prescribed class of medicinal product or activity)—
 - (a) with which the holder of a manufacturer's licence must comply, and
 - (b) which are to have effect as if they were provisions of the licence.]
- [^{F33}(3) [^{F34}Subject to [^{F35}subsections (3C) and (3D)] of this section,] no person shall, in the course of a business carried on by him—
 - (a) sell, or offer for sale, any medicinal product by way of wholesale dealing, or
 - (b) distribute, otherwise than by way of sale, any proprietary medicinal product ^{F36}[^{F37}... or industrially produced medicinal product ^{F36}...] which has been imported, but was not consigned from a member State,

except in accordance with a [^{F38}wholesale dealer's licence].]

- [^{F39}(3A) Without prejudice to the generality of subsection (3) of this section but subject to [^{F40}subsections (3C) and (3D)], no person shall, in the course of a business carried on by him, distribute by way of wholesale dealing a product to which [^{F41}the 2001 Directive applies] apply except in accordance with a wholesale dealer's licence.
 - (3B) Distribution of such a product by way of wholesale dealing shall not be taken to be in accordance with a wholesale dealer's licence unless, in particular, it occurs in the course of a business which is carried on at a place or places specified in the licence.
 - (3C) The restrictions imposed by subsections (3) and (3A) of this section do not apply to anything done in relation to a product to which [^{F41}the 2001 Directive applies] apply by the holder of a manufacturer's licence in respect of it.]
- [^{F42}(3D) The restrictions imposed by subsections (3) and (3A) of this section do not apply where the product concerned is an investigational medicinal product within the meaning given by the Clinical Trials Regulations.]
- [^{F43}(3E) The Ministers may prescribe requirements (either generally or in relation to a prescribed class of medicinal product or activity)—
 - (a) with which the holder of a wholesale dealer's licence must comply, and
 - (b) which are to have effect as if they were provisions of the licence.]

[^{F44}(4) ^{F45}... subsection (3)(b) of this section shall not apply if the product is—

- $F^{46}(a)$
 - (b) a radiopharmaceutical in which the radionuclide is in the form of a sealed source, $[F^{47}...]$
- ^{F47}(c) ...
- $F^{48}(5)$
 - (6) In this section, [^{F49}[^{F50}"proprietary medicinal product" and "radiopharmaceutical"] have the same meanings as in section 7 of this Act.]

[In this section any reference to distribution of a product by way of wholesale dealing $^{F51}(7)$ is a reference to—

- (a) selling or supplying it, or
- (b) procuring, holding or exporting it for the purposes of sale or supply,
- to a person who receives it for the purposes of-
 - (i) selling or supplying it, or

(ii) administering it or causing it to be administered to one or more human beings, in the course of a business carried on by that person.

(8) In this Act any reference to a wholesale dealer's licence is a reference to a licence granted for the purposes of subsection (3) or (3A) of this section.]]

Textual Amendments

- **F25** Words in s. 8(1) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 8(a) (with regs. 2(4), 3)
- F26 Words in s. 8(2) inserted (1.5.2004) by Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 4(2)
- F27 Words in s. 8(2) substituted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 1(2)(a) (with Sch. 6)
- F28 Words in s. 8(2) substituted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 1(2)(b) (with Sch. 6)
- **F29** S. 8(2A)(2B) inserted (1.5.2004) by Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 4(3)
- **F30** Words in s. 8(2B) substituted (29.12.2008) by The Medicines for Human Use (Marketing Authorisations Etc.) Amendment Regulations 2008 (S.I. 2008/3097), regs. 1(1), 4(a)(i)
- **F31** Words in s. 8(2B) inserted (29.12.2008) by The Medicines for Human Use (Marketing Authorisations Etc.) Amendment Regulations 2008 (S.I. 2008/3097), regs. 1(1), 4(a)(ii)
- F32 S. 8(2C)(2D) inserted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 1(3) (with Sch. 6)
- **F33** S. 8(3)(4) substituted for s. 8(3) by (E.W.)(S.) S.I. 1977/1050, art. 3(2) and (N.I.) S.R. 1977 No. 170, reg. 4
- F34 Words in s. 8(3) inserted (14.4.1993) by S.I. 1993/834, reg. 2(2)
- **F35** Words in s. 8(3) substituted (1.5.2004) by Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 4(4)
- **F36** Words in s. 8(3)(b) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 8(b) (with regs. 2(4), 3)
- F37 Words in s. 8(3)(b) substituted (3.4.1992) by virtue of S.I.1992/604, regs. 3(2), 4

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- **F38** Words in s. 8(3) substituted (14.4.1993) by S.I. 1993/834, reg. 2(3)
- **F39** S. 8(3A)-(3C) inserted (14.4.1993) by S.I. 1993/834, reg. 2(4)
- F40 Words in s. 8(3A) substituted (1.5.2004) by Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 4(4)
- F41 Words in s. 8(3A)(3C) substituted (28.2.2002) by S.I. 2002/236, reg. 2(a)(i)
- F42 S. 8(3D) inserted (1.5.2004) by Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 4(5)
- **F43** S. 8(3E) inserted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 1(4) (with Sch. 6)
- F44 S. 8(4)(5)(6) substituted (3.4.1992) for s. 8(4) by virtue of S.I. 1992/604, regs. 3(3), 4
- **F45** Words in s. 8(4) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 8(c) (with regs. 2(4), 3)
- **F46** S. 8(4)(a) omitted (8.11.2005) by virtue of The Blood Safety and Quality Regulations 2005 (S.I. 2005/50), regs. 1(2), **25(1)(b)** (with reg. 2(2)-(4))
- **F47** In S. 8(4) paragraph (c) and words immediately preceeding it omitted (13.2.1994) by virtue of S.I. 1994/276, reg. 4(2)(b) (which S.I. revoked and replaced defective S.I. 1994/101 before the latter S.I. came into force)
- **F48** S. 8(5) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 8(d) (with regs. 2(4), 3)
- F49 Words in s.8(6) inserted (13.2.1994) by S.I. 1994/276, reg. 4(3) (which S.I. revoked and replaced defective S.I. 1994/101 before the latter S.I. came into force)
- **F50** Words in s. 8(6) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 8(e) (with regs. 2(4), 3)
- F51 S. 8(7)(8) added (14.4.1993) by S.I. 1993/834, reg. 2(5)

Modifications etc. (not altering text)

- C12 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C13 S. 8 excluded by S.I. 1989/2325, art. 2(3)
- C14 S. 8(2) excluded by S.I. 1979/1114, arts. 2, 4 and by S.I. 1979/1585, arts. 2, 3
- C15 S. 8(3) excluded by S.I. 1989/2322, art. 2(1)
- C16 S. 8(3) excluded by S.I. 1990/566, art. 2(1)
- C17 S. 8(3)(*b*) excluded by S.I. 1989/2322, art. 2(3)
- C18 S. 8(3)(b) excluded by S.I. 1990/566, art. 2(3)

9 [^{F52}Exemptions for doctors and dentists]

- (1) The restrictions imposed by sections 7 and 8 of this Act do not apply to anything done by a doctor or dentist which—
 - (a) relates to a medicinal product specially prepared, or specially imported by him or to his order, for administration to a particular patient of his, and consists of manufacturing or assembling, or procuring the manufacture or assembly of, the product, or of selling or supplying, or procuring the sale or supply of, the product to that patient or to a person under whose care that patient is, or
 - (b) relates to a medicinal product specially prepared at the request of another doctor or dentist, or specially imported by him or to his order at the request of another doctor or dentist, for administration to a particular patient of that other doctor or dentist, and consists of manufacturing or assembling, or procuring the manufacture or assembly of, the product, or of selling or supplying, or procuring the sale or supply of, the product to that other doctor or dentist or to that patient or to a person under whose care that patient is.

Textual Amendments

- **F52** Words in s. 9 heading substituted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 9
- **F53** S. 9(2)(3) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 9(a) (with regs. 2(4), 3)

Modifications etc. (not altering text)

- C19 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C20 S. 9(2) restricted by S.I. 1987/2217, art. 3

10 Exemptions for pharmacists.

- (1) ^{F54}... the restrictions imposed by sections 7 and 8 of this Act do not apply to anything which is done in a registered pharmacy, a hospital [^{F55}, a care home service] or a health centre and is done there by or under the supervision of a pharmacist and consists of—
 - (a) preparing or dispensing a medicinal product in accordance with a prescription given by a practitioner, or
 - (b) assembling a medicinal product [^{F56}provided that where the assembling takes place in a registered pharmacy—
 - (i) it shall be in a registered pharmacy at which the business in medicinal products carried on is restricted to retail sale or to supply in circumstances corresponding to retail sale and the assembling is done with a view to such sale or supply either at that registered pharmacy or at any other such registered pharmacy forming part of the same retail pharmacy business, and
 - (ii) the medicinal product has not been the subject of an advertisement]; and those restrictions do not apply to anything done by or under the supervision of a pharmacist which consists of procuring the preparation or dispensing of a medicinal product in accordance with a prescription given by a practitioner, or of procuring the assembly of a medicinal product.
- - (3) Those restrictions do not apply to the preparation or dispensing in a registered pharmacy of a medicinal product by or under the supervision of a pharmacist in accordance with a specification furnished by the person to whom the product is or is to be sold or supplied, where—
 - (a) the product is prepared or dispensed for administration to that person or to a person under his care, ^{F58}...
 - ^{F58}(b)
 - (4) Without prejudice to the preceding subsections, the restrictions imposed by sections 7 and 8 of this Act do not apply to anything which is done in a registered pharmacy by or under the supervision of a pharmacist and consists of—

- (a) preparing or dispensing a medicinal product for administration to a person where the pharmacist is requested by or on behalf of that person to do so in accordance with the pharmacist's own judgment as to the treatment required, and that person is present in the pharmacy at the time of the request in pursuance of which that product is prepared or dispensed, or
- (b) preparing a stock of medicinal products with a view to dispensing them as mentioned in subsection (1)(a) or subsection (3) of this section or in paragraph (a) of this subsection [^{F59}provided that such stock is prepared with a view to retail sale or to supply in circumstances corresponding to retail sale and the preparation is done with a view to such sale or supply either at that registered pharmacy or at any other registered pharmacy forming part of the same retail pharmacy business];

and those restrictions do not apply to anything which is done in a hospital or a health centre by or under the supervision of a pharmacist and consists of preparing a stock of medicinal products with a view to dispensing them as mentioned in subsection (1) (a) of this section.

- [^{F60}(5) Without prejudice to the preceding subsections, the restrictions imposed by section 7 of this Act do not apply to the preparation or dispensing in a registered pharmacy of a medicinal product by or under the supervision of a pharmacist where—
 - (a) the medicinal product is prepared or dispensed otherwise than in pursuance of an order from any other person, and
 - (b) the medicinal product is prepared with a view to retail sale or supply in circumstances corresponding to retail sale at the registered pharmacy at which it is prepared, and
 - (c) the medicinal product has not been the subject of an advertisement.
 - (6) Without prejudice to the preceding subsections, the restrictions imposed by section 8(2) of this Act do not apply to anything which is done in a registered pharmacy by or under the supervision of a pharmacist and consists of preparing a medicinal product with a view to retail sale or to supply in circumstances corresponding to retail sale at that registered pharmacy.

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^{F62}(6A)]

- (7) Without prejudice to the preceding subsections, the restrictions imposed by section 8(3) [^{F63}or (3A)] of this Act do not apply to anything which is done in a registered pharmacy by or under the supervision of a pharmacist and amounts to wholesale dealing, where such dealing constitutes no more than an inconsiderable part of the business carried on by the pharmacist at that pharmacy.
- [The ^{F65}... Ministers may make regulations prescribing conditions which must be ^{F64}(7A) complied with if a thing is to be considered for the purposes of this section as done under the supervision of a pharmacist.
 - (7B) Conditions prescribed under subsection (7A) may relate to supervision in the case where the pharmacist is not at the place where the thing is being done, and in that case the thing is not to be so considered if no such conditions are prescribed.
 - (7C) In any case, compliance with any applicable conditions is sufficient for the thing to be so considered.]

Changes to legislation: Medicines Act 1968, Part II is up to date with all changes known to be in force on or before 27 June 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details)

- (8) For the purposes of this section "advertisement" shall have the meaning assigned to it by section 92 of this Act, except that it shall not include words inscribed on the medicinal product, or on its container or package.]
- [^{F66}(9) In subsection (1) of this section, "care home service" has the meaning given by [^{F67}paragraph 2 of schedule 12 to the Public Services Reform (Scotland) Act 2010 (asp 8)].]

Textual Amendments

- **F54** Words in s. 10(1) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 10(a) (with regs. 2(4), 3)
- F55 Words in s. 10(1) inserted (S.) (1.4.2002) by 2001 asp 8, s. 79, Sch. 3 para. 5(a); S.S.I. 2002/162, art. 2(h) (subject to arts. 3-13)
- F56 Words added by S.I. 1971/1445, art. 3(a)
- **F57** S. 10(2) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 10(b) (with regs. 2(4), 3)
- **F58** S. 10(3)(b) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 10(c) (with regs. 2(4), 3)
- **F59** Words added by S.I. 1971/1445, art. 3(b)
- **F60** S. 10(5)–(8) added by S.I. 1971/1445, art. 3(c)
- **F61** S. 10(6A) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 10(d) (with regs. 2(4), 3)
- F62 S. 10(6A) inserted (31.12.1994) by S.I. 1994/2987, reg. 11(2)
- F63 Words in s. 10(7) inserted (14.4.1993) by S.I. 1993/834, reg. 3
- F64 S. 10(7A)-(7C) inserted (19.7.2006 for specified purposes) by Health Act 2006 (c. 28), ss. 26(1), 83(1) (e)
- **F65** Word in s. 10(7A) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 10(e) (with regs. 2(4), 3)
- F66 S. 10(9) added (S.) (1.4.2002) by 2001 asp 8, ss. 79, Sch. 3 para. 5(b); S.S.I. 2002/162, art. 2(h) (subject to arts. 3-13)
- F67 Words in s. 10(9) substituted (28.10.2011) by The Public Services Reform (Scotland) Act 2010 (Consequential Modifications of Enactments) Order 2011 (S.I. 2011/2581), art. 1(2)(b), Sch. 2 para. 1

Modifications etc. (not altering text)

- C21 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C22 S. 10 amended (E.W.S.) (*prosp*) by 1954 c. 61, s. 13I(1)(b) (as inserted (*prosp.*) by 1997 c. 19, ss. 1, 2(1), Sch. para. 2)

11 Exemption for nurses and midwives.

- (1) The restrictions imposed by section 8 of this Act do not apply to the assembly of any medicinal products by a person in the course of that person's profession as [^{F68}a registered ^{F69}... nurse or a registered midwife] ...

Textual Amendments

F68 Words substituted by Nurses, Midwives and Health Visitors Act 1979 (c. 36, SIF 83:1), s. 24(2), Sch. 7 para. 14(a)

- **F69** Words in s. 11(1) omitted (1.8.2004) by virtue of The Health Act 1999 (Consequential Amendments) (Nursing and Midwifery) Order 2004 (S.I. 2004/1771), art. 1(1), **Sch. para. 10(a)**
- F70 Words repealed by Nurses, Midwives and Health Visitors Act 1979 (c. 36, SIF 83:1), s. 24(2), Sch. 8
- F71 S. 11(2) repealed by Nurses, Midwives and Health Visitors Act 1979 (c. 36, SIF 83:1), s. 24(2), Sch. 8

Modifications etc. (not altering text)

C23 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

12 Exemptions in respect of herbal remedies.

- (1) The restrictions imposed by sections 7 and 8 of this Act do not apply to the sale, supply, manufacture or assembly of any herbal remedy in the course of a business where—
 - (a) the remedy is manufactured or assembled on premises of which the person carrying on the business is the occupier and which he is able to close so as to exclude the public, and
 - (b) the person carrying on the business sells or supplies the remedy for administration to a particular person after being requested by or on behalf of that person and in that person's presence to use his own judgment as to the treatment required.

 $F^{72}(2)$

Textual Amendments

F72 S. 12(2) repealed (30.4.2011) by The Medicinal Products (Herbal Remedies) (Amendment) Regulations 2011 (S.I. 2011/915), regs. 1, **2**

Modifications etc. (not altering text)

C24 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

13 Exemptions for imports.

- (1) The restriction imposed by section 7(3) of this Act does not apply to the importation of a medicinal product by any person for administration to himself or to any person or persons who are members of his household, and does not apply to the importation of a medicinal product where it is specially imported by or to the order of a doctor or dentist for administration to a particular patient of his.
- (2) Without prejudice to the preceding subsection, the restriction imposed by section 7(3) of this Act shall not apply to the importation of a medicinal product in such circumstances as may be specified in an order made by the Ministers for the purposes of this section.
- (3) Any exemption conferred by an order under this section may be conferred either in relation to medicinal products generally or in relation to a class of medicinal products specified in the order, and (in either case) may be so conferred subject to such conditions or limitations as may be so specified.

Modifications etc. (not altering text)

C25 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

Changes to legislation: Medicines Act 1968, Part II is up to date with all changes known to be in force on or before 27 June 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details)

14 Exemption for re-exports.

- [^{F73}(1)] [^{F74}Subject to subsection (2) of this section,] the restrictions imposed by sections 7 and 8 of this Act do not apply to the exportation, or the sale or offer for sale for the purposes of exportation, of any imported medicinal product if it is, or is to be, exported—
 - (a) in the form in which it was imported, and
 - (b) without being assembled in a way different from the way in which it was assembled on being imported.
- ^{F75}[(2) Section 8(3A) of this Act applies to the exportation, or the sale for exportation, of any product to which [^{F76}the 2001 Directive applies] apply if it is, or is to be exported to [^{F77}an EEA State.]]

Textual Amendments

- F73 S. 14 renumbered as s. 14(1) (14.4.1993) by S.I. 1993/834, reg. 4(a)
- **F74** Words in s. 14(1) inserted (14.4.1993) by S.I. 1993/834, reg. 4(a)
- **F75** S. 14(2) inserted (14.4.1993) by S.I. 1993/834, reg. 4(b)
- F76 Words in s. 14(2) substituted (28.2.2002) by S.I. 2002/236, reg. 2(a)(ii)
- F77 Words in s. 14(2) substituted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 2 (with Sch. 6)

Modifications etc. (not altering text)

- C26 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C27 S. 14 applied (with modifications)(3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.

15 Provision for extending or modifying exemptions.

- (1) The ^{F78}... Ministers may by order provide that sections 7 and 8 of this Act shall have effect subject to such exemptions (other than those for the time being having effect by virtue of sections 9 to 14 of this Act) as may be specified in the order.
- (2) Any exemption conferred by an order under the preceding subsection may be conferred subject to such conditions or limitations as may be specified in the order.
- (3) The ^{F79}... Ministers may by order provide that any of the provisions of sections 9 to 14 of this Act specified in the order shall cease to have effect, or shall have effect subject to such exceptions or modifications as may be so specified.
- (4) No order shall be made under subsection (3) of this section unless a draft of the order has been laid before Parliament and approved by a resolution of each House of Parliament.

Textual Amendments

- **F78** Word in s. 15(1) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 11(a) (with regs. 2(4), 3)
- **F79** Word in s. 15(3) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 11(b) (with regs. 2(4), 3)

Changes to legislation: Medicines Act 1968, Part II is up to date with all changes known to be in force on or before 27 June 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details)

Modifications etc. (not altering text)

C28 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

16 Transitional exemptions.

- (1) The restrictions imposed by sections 7 and 8 of this Act do not apply to anything done before such day as the Ministers may by order appoint for the purposes of this subsection (in this Act referred to as "the first appointed day"); and, except as otherwise provided by any order made under section 17 of this Act, the following provisions of this section shall have effect in relation to things done on or after that day.
- (2) Section 7(2) of this Act shall not have effect in relation to a person in respect of his selling or supplying, or procuring the sale, supply, manufacture or assembly of, medicinal products of any description if, in the course of a business carried on by him, any medicinal products of that description were sold or supplied, or procured to be sold, supplied, manufactured or assembled, at any time before the first appointed day and medicinal products of that description were effectively on the market in the United Kingdom immediately before the first appointed day, and either—
 - (a) information with regard to the composition of medicinal products of that description, and as to their being available for sale or supply in the United Kingdom, had before that day been made known generally to doctors, or to any particular class of doctors, or to dentists or pharmacists, or to veterinary surgeons and veterinary practitioners, in the United Kingdom, or
 - (b) information that the products were available for sale or supply in the United Kingdom had before that day been made known generally to the public in the United Kingdom.
- (3) Section 7(3) of this Act shall not have effect in relation to a person in respect of his importing medicinal products of any description in the course of a business carried on by him if, in the course of that business, medicinal products of that description were imported within the period of twenty-four months ending with the first appointed day.
- (4) Section 8(2) of this Act shall not have effect in relation to a person in respect of his manufacturing or assembling medicinal products of any description in the course of a business carred on by him if in the course of that business—
 - (a) medicinal products of that description were manufactured or assembled within the period of twelve months ending with the first appointed day, or
 - (b) medicinal products of that description were manufactured or assembled before the beginning of that period and further supplies of such products could, if required, have been manufactured or assembled within that period:

Provided that this subsection shall not have effect in relation to any particular operations carried out in the course of a business on or after the first appointed day unless the manufacture or assembly of the products as mentioned in paragraph (a) or paragraph (b) of this subsection, as the case may be, included those operations.

(5) Section 8(3) of this Act shall not have effect in relation to a person in respect of his selling or offering for sale any medicinal products by way of wholesale dealing in the course of a business carried on by him if, in the course of that business, medicinal products were being sold or offered for sale by way of wholesale dealing within the period of twelve months ending with the first appointed day.

Modifications etc. (not altering text)

- C29 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C30 S. 16(1) modified by S.I. 1985/1403, art. 3(2) and S.I. 1985/1539, art. 1
- C31 S. 16(2)–(5) excluded by S.I. 1981/1690, art. 2

17 Termination of transitional exemptions.

For the purposes of subsections (2) to (5) of the last preceding section, the Ministers may by one or more orders under this section appoint one or more days, subsequent to the first appointed day, and may by any such order provide that such one or more of those subsections as may be specified in that order shall cease to have effect either—

- (a) generally in relation to anything done on or after the day appointed by that order, or
- (b) in relation to anything done on or after that day in so far as it consists of operations or activities, or relates to medicinal products of any such class, as may be so specified.

Modifications etc. (not altering text) C32 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

Applications for, and grant and renewal of, licences

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Modifications etc. (not altering text)
C33 Definitions in ss. 18-22 applied (N.I.) (1.4.1992) by S.I. 1991/194 (N.I. 1), art. 7(2), Sch. 2 para. 5(4);
S.R. 1991/131, art. 2(e), Sch. Pt.III
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18 Application for licence.

- (1) Any application for the grant of a licence under this Part of this Act shall be made to the licensing authority and shall be made in such form and manner, and shall contain, or be accompanied by, such information, documents, samples and other material, as may be prescribed.
- (2) Any such application shall indicate the descriptions of medicinal products in respect of which the licence is required, either by specifying the descriptions of medicinal products in question or by way of an appropriate general classification.

^{F80}(3)....

Textual Amendments

F80 S. 18(3) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 12 (with regs. 2(4), 3)

Modifications etc. (not altering text)

C34 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

- C35 S. 18 applied (with modifications)(11.3.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.
- C36 Definitions in ss. 18-22 applied (N.I)(1.4.1992) by S.I. 1991/194 (N.I. 1), art. 7(2), Sch. 2 para. 5(4); S.R. 1991/131, art. 2(e), Sch. Pt. III

19 Factors relevant to determination of application for licence.

- (1) Subject to the following provisions of this Part of this Act, in dealing with an application for a product licence the licensing authority shall in particular take into consideration—
 - (a) the safety of medicinal products of each description to which the application relates;
 - (b) the efficacy of medicinal products of each such description for the purposes for which the products are proposed to be administered; and
 - (c) the quality of medicinal products of each such description, according to the specification and the method or proposed method of manufacture of the products, and the provisions proposed for securing that the products as sold or supplied will be of that quality.
- (2) In taking into consideration the efficacy for a particular purpose of medicinal products of a description to which such an application relates, the licensing authority shall leave out of account any question whether medicinal products of another description would or might be equally or more efficacious for that purpose:

Provided that nothing in this subsection shall be construed as requiring the licensing authority, in considering the safety of medicinal products of a particular description, in relation to a purpose for which they are proposed to be administered, to leave out of account any question whether medicinal products of another description, being equally or more efficacious for that purpose, would or might be safer in relation to that purpose.

- (3) Where any such application indicates that the purposes for which the licence is required relate (wholly or partly) to medicinal products which have been or are to be imported, then in dealing with the application, in so far as it relates to such products, the licensing authority shall also take into consideration in particular the methods, standards and conditions of manufacture of those products and may, if they think fit, require the production by the applicant of any one or more of the following, that is to say—
 - (a) an undertaking, given by the manufacturer of any such products, to permit the premises where they are or are to be manufactured, and the operations carried on or to be carried on in the course of manufacturing them, to be inspected by or on behalf of the licensing authority;
 - (b) an undertaking, given by or on behalf of the manufacturer of any such products, to comply with any prescribed conditions or any conditions attached to the licence by the licensing authority;
 - (c) a declaration, given by or on behalf of the manufacturer of any such products, that, in relation to the manufacture of those products, any requirements imposed by or under the law of the country in which they are or are to be manufactured have been or will be complied with.
- (4) Where any such application indicates that the purposes for which the license is required relate exclusively to the exportation of medicinal products, the licensing authority shall leave out of account considerations of safety and efficacy (as mentioned in paragraphs (a) and (b) of subsection (1) of this section) if satisfied that in the circumstances it is reasonable to do so.

- (5) In dealing with an application for a manufacturer's licence the licensing authority shall in particular take into consideration—
 - (a) the operations proposed to be carried out in pursuance of the licence:
 - (b) the premises in which those operations are to be carried out;
 - (c) the equipment which is or will be available on those premises for carrying out those operations;
 - (d) the qualifications of the persons under whose supervision those operations will be carried out; and
 - (e) the arrangements made or to be made for securing the safekeeping of, and the maintenance of adequate records in respect of, medicinal products manufactured or assembled in pursuance of the licence.
- (6) In dealing with an application for a wholesale dealer's licence the licensing authority shall in particular take into consideration—
 - (a) the premises on which medicinal products of the descriptions to which the application relates will be stored;
 - (b) the equipment which is or will be available for storing medicinal products on those premises;
 - (c) the equipment and facilities which are or will be available for distributing medicinal products from those premises; and
 - (d) the arrangements made or to be made for securing the safekeeping of, and the maintenance of adequate records in respect of, medicinal products stored on or distributed from those premises.
- (7) The preceding provisions of this section shall have effect subject to the provisions of this Part of this Act relating to licences of right.

Modifications etc. (not altering text)

- C37 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C38 Definitions in ss. 18-22 applied (N.I) (1.4.1992) by S.I. 1991/194, (N.I. 1), art. 7(2), Sch. 2 para. 5(4); S.R. 1991/194, art. 2(e), Sch. Pt. III
- C39 S. 19 (1)(2)(3) applied (with modifications) by S.I. 1992/605, reg. 2(1)(2), Sch.

20 Grant or refusal of licence.

- (1) Subject to [^{F81}sections 8(2E) and (3E) and 19,], and to the following provisions of this Act, on any application to the licensing authority for a licence under this Part of this Act the licensing authority—
 - (a) may grant a licence containing such provisions as they consider appropriate, or
 - (b) if, having regard to the provisions of this Act [^{F82}and any [^{F83}EU] obligation], they consider it necessary or expedient to do so, may refuse to grant a licence.
- [^{F84}(1A) The licensing authority must either grant or refuse any application for a licence under this Part, before the end of a period of 90 days from the date upon which they receive the application.]
- [^{F84}(2B) If there are requirements in force under section 18 that apply to the application, subsection (1A) applies only if the requirements have been met]

- [^{F84}(2C) If a notice under section 44 requires the applicant to provide the licensing authority with information, the period specified in subsection (1) stops running when the notice is given, and does not start running again until—
 - (a) the licensing authority receives the information; or
 - (b) the applicant has shown to the reasonable satisfaction of the licensing authority why he is unable to provide it.]
 - (2) The licensing authority shall not refuse to grant such a licence on any grounds relating to the price of any product, and shall not insert in any such licence any provisions as to the price at which any product may be sold, supplied, imported or exported.
 - (3) The licensing authority shall not refuse to grant such a licence on any grounds relating to the safety, quality or efficacy of medicinal products of any description, except after consultation with the appropriate committee ^{F85}....
 - - (5) Where on an application for a licence under this Part of this Act—
 - (a) the licensing authority refuse to grant a licence, or
 - (b) the licensing authority grant a licence otherwise than in accordance with the application, and the applicant requests the licensing authority to state their reasons,

the licensing authority shall serve on the applicant a notice stating the reasons for their decision.

Textual Amendments

- F81 Words in s. 20(1) substituted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 3(a) (with Sch. 6)
- F82 Words inserted by (E.W.) S.I. 1977/1050, art. 4(3) and (N.I.) S.R. 1977 No. 170, reg. 5(3)
- **F83** Words in Act substituted (22.4.2011) by The Treaty of Lisbon (Changes in Terminology) Order 2011 (S.I. 2011/1043), arts. 2, **3**, 6 (with art. 3(2)(3)4(2)6(4)6(5))
- F84 S. 20(1A)(2B)(2C) inserted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 3(b) (with Sch. 6)
- **F85** Words in s. 20(3) omitted (30.10.2005) by virtue of Medicines (Advisory Bodies) Regulations 2005 (S.I. 2005/1094), reg. 1(1), Sch. 1 para. 1(a)
- **F86** S. 20(4) omitted (30.10.2005) by virtue of Medicines (Advisory Bodies) Regulations 2005 (S.I. 2005/1094), reg. 1(1), Sch. 1 para. 1(b)

Modifications etc. (not altering text)

- C40 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C41 Ss. 20-22 applied (with modifications) (3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.
- C42 Definitions in ss. 18-22 applied (N.I.) (1.4.1992) by S.I. 1991/194, (N.I. 1), art. 7(2), Sch. 2 para. 5(4); S.R. 1991/131, art. 2(e), Sch. Pt.III

[^{F87}21 Procedure on reference to appropriate committee

- (1) Where the appropriate committee are consulted under section 20(3) of this Act and are of the provisional opinion that, on grounds relating to safety, quality or efficacy, they—
 - (a) may be unable to advise the licensing authority to grant the licence; or
 - (b) may be unable to advise the licensing authority to grant it unless it contains provisions otherwise than in accordance with the application,

they shall notify the applicant accordingly.

- (2) A person who has been so notified may, within the time allowed, give notice of his wish to make written or oral representations to the appropriate committee.
- (3) The appropriate committee shall give the applicant an opportunity to make such representations in accordance with subsections (4) to (7) of this section.
- (4) Subject to subsection (5) of this section, the applicant shall provide the appropriate committee with—
 - (a) his written representations or a written summary of the oral representations he intends to make; and
 - (b) any documents on which he wishes to rely in support of those representations,

before the end of the period of six months beginning with the date of the notice referred to in subsection (2) of this section, or within such shorter period as the appropriate committee may specify in the notification under subsection (1).

- (5) If the applicant so requests, the appropriate committee may extend the time limit referred to in subsection (4) of this section, up to a maximum period of twelve months beginning with the date of the notice referred to in subsection (2) of this section.
- (6) The applicant may not submit any additional written representations or documents once the time limit referred to in subsections (4) and (5) of this section has expired, except with the permission of the appropriate committee.
- (7) If the applicant gave notice of his wish to make oral representations, the appropriate committee shall, after receiving a written summary and any other documents in accordance with subsection (4) of this section, arrange for the applicant to make such representations at a hearing before the committee.
- (8) The appropriate committee shall—
 - (a) take into account such representations as are made in accordance with this section; and
 - (b) report their findings and advice to the licensing authority, together with the reasons for their advice.
- (9) After receiving the report of the appropriate committee, the licensing authority shall—
 - (a) decide whether to grant or refuse the application, or to grant it otherwise than in accordance with the application; and
 - (b) take the report into account when making their decision.
- (10) The licensing authority shall notify the applicant of-
 - (a) the decision made pursuant to subsection (9) of this section; and
 - (b) the advice given to them by the appropriate committee and the reasons for that advice.

(11) If—

- (a) the applicant has made representations in accordance with this section and the licensing authority have notified the applicant of the authority's decision to refuse to grant the licence, or to grant it otherwise than in accordance with the application; or
- (b) the applicant has not made representations in accordance with this section and the licensing authority have notified the applicant of the authority's decision to refuse to grant the licence, or to grant it otherwise than in accordance with the application, on grounds which differ from those relied on in the advice of the appropriate committee,

the applicant may, within the time allowed, notify the licensing authority that he wishes to appear before and be heard by a person appointed by the licensing authority with respect to the decision.

(12) In this Part of the Act, "the time allowed" means the period of twenty-eight days beginning with the date of the relevant notification, or such longer period as the licensing authority may allow in any particular case.]

Textual Amendments

F87 S. 21 substituted (30.10.2005) by Medicines (Advisory Bodies) Regulations 2005 (S.I. 2005/1094), reg. 1(1), Sch. 1 para. 2

Modifications etc. (not altering text)

- C43 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C44 Ss. 20-22 applied (with modifications) (3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.
- C45 Definitions in ss. 18-22 applied (N.I.) (1.4.1992) by S.I. 1991/194, (N.I. 1), art. 7(2), Sch. 2 para. 5(4); S.R. 1991/131, art. 2(e), Sch. Pt.III
- C46 S. 21 extended (with modifications) (14.2.1994) by S.I. 1994/105, reg. 19, Sch.4
- C47 S. 21 amendment to earlier affecting provision SI 1994/105 Sch. 4 (30.10.2005) by Medicines (Homoeopathic Medicinal Products for Human Use) Amendment Regulations 2005 (S.I. 2005/2753), regs. 1(1), 19(2)
- C48 S. 21 applied (with modifications) (30.10.2005) by Medicines (Advisory Bodies) (No.2) Regulations 2005 (S.I. 2005/2754), reg. 1(2)(b), Sch. 5 para. 11(3)
- C49 S. 21 applied (with modifications) (30.10.2005) by Medicines (Advisory Bodies) (No.2) Regulations 2005 (S.I. 2005/2754), reg. 1(2)(b), Sch. 5 para. 6(3)
- C50 S. 21(11)(a)(b) applied (with modifications) (30.10.2005) by Medicines (Advisory Bodies) (No.2) Regulations 2005 (S.I. 2005/2754), reg. 1(2)(b), Sch. 5 para. 2(4)

[^{F88} 22 Procedure in other cases.

(1) This section applies when-

- (a) an application is made for the grant of a licence under this Part of this Act; and
- (b) the appropriate committee—
 - (i) is not consulted under subsection (3) of section 20, or
 - (ii) is consulted under that subsection but does not give a provisional opinion in accordance with section 21(1).
- (2) If the licensing authority propose—
 - (a) to refuse to grant the licence, or

- (b) to grant it otherwise than in accordance with the application,
- they shall notify the applicant of their proposals and the reasons for them.
- (3) If the applicant is so notified, he may, within the time allowed—
 - (a) notify the licensing authority of his wish to appear before and be heard by a person appointed by the licensing authority with respect to the proposal; or
 - (b) make representations in writing to the licensing authority with respect to the proposal referred to in the notification.
- (4) If the applicant makes written representations in accordance with subsection (3)(b) of this section, the licensing authority shall take those representations into account before determining the application.

Textual Amendments

F88 Ss. 22, 22A substituted for s. 22 (30.10.2005) by Medicines (Advisory Bodies) Regulations 2005 (S.I. 2005/1094), reg. 1(1), **Sch. 1 para. 3**

Modifications etc. (not altering text)

- C51 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C52 Ss. 20-22 applied (with modifications) (3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.
- C53 Definitions in ss. 18-22 applied (N.I.) (1.4.1992) by S.I. 1991/194 (N.I. 1), art. 7(2), Sch. 2 para. 5(4); S.R. 1991/131, art. 2(e), Sch. Pt.III
- C54 s. 22 extended (with modifications) (14.2.1994) by S.I. 1994/105, reg. 19, Sch.4
- C55 S. 22 amendment to earlier affecting provision SI 1994/105 Sch. 4 (30.10.2005) by Medicines (Homoeopathic Medicinal Products for Human Use) Amendment Regulations 2005 (S.I. 2005/2753), regs. 1(1), **19(2)**

22A. Hearing before person appointed

- (1) If the applicant gives notice under section 21(11) or section 22(3) of his wish to appear before and be heard by a person appointed by the licensing authority, the authority shall—
 - (a) make that appointment; and
 - (b) arrange for the applicant to have an opportunity of appearing before that person.

(2) The person appointed—

- (a) shall not be, or at any time have been, a member of—
 - (i) the Commission on Human Medicines or any of its Expert Advisory Groups,
 - (ii) the Medicines Commission formerly established under section 2 of this Act or any of its committees, or
 - (iii) a committee established under section 4 of this Act, or any subcommittee of such a committee; and
- (b) shall not be an officer or servant of any Minister of the Crown.
- (3) Subject to subsection (4) of this section, the applicant shall provide the person appointed with—
 - (a) a written summary of the oral representations he intends to make; and
 - (b) any documents on which he wishes to rely in support of those representations,

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before the end of the period of three months beginning with the date of the notice referred to in subsection (1) of this section.

- (4) If the applicant so requests, the person appointed may, after consulting the licensing authority, extend the time limit referred to in subsection (3) of this section, up to a maximum period of six months beginning with the date of the notice referred to in subsection (1) of this section.
- (5) If the applicant fails to comply with the time limit in subsection (3) of this section, or, where he has been granted an extended time limit under subsection (4) of this section, that time limit—
 - (a) he may not appear before or be heard by the person appointed, and
 - (b) the licensing authority shall decide whether to grant or refuse the licence, or to grant it otherwise than in accordance with the application, and notify the applicant accordingly.
- (6) The applicant may not submit any additional written representations or documents once the time limit has expired, except with the permission of the person appointed.
- (7) At the hearing before the person appointed, both the applicant and the licensing authority may make representations.
- (8) If the applicant so requests the hearing shall be in public.
- (9) After the hearing—
 - (a) the person appointed shall provide a report to the licensing authority; and
 - (b) the licensing authority shall take the report into account and decide whether to grant or refuse the licence, or to grant it otherwise than in accordance with the application, or to confirm or alter their decision, as the case may be.
- (10) The licensing authority shall then—
 - (a) notify the applicant of their decision;
 - (b) if the applicant so requests, provide the applicant with a copy of the report of the person appointed.]

Textual Amendments

F88 Ss. 22, 22A substituted for s. 22 (30.10.2005) by Medicines (Advisory Bodies) Regulations 2005 (S.I. 2005/1094), reg. 1(1), Sch. 1 para. 3

Modifications etc. (not altering text)

- C56 S. 22A applied by SI 1992/605 Sch. (as amended) (30.10.2005) by Medicines (Advisory Bodies) (No.2) Regulations 2005 (S.I. 2005/2754), reg. 1(2)(b), Sch. 4 para. 6(a)
- C57 S. 22A excluded (30.10.2005) by Medicines (Advisory Bodies) (No.2) Regulations 2005 (S.I. 2005/2754), reg. 1(2)(b), Sch. 5 para. 19(3)(a)(i)
- C58 S. 22A(10)(b) applied by SI 1978/1006 reg. 7(3) (as substituted (30.10.2005) by Medicines (Advisory Bodies) (No.2) Regulations 2005 (S.I. 2005/2754), reg. 1(2)(b), Sch. 4 para. 3(b))
- C59 Ss. 22A(2)-(9) applied by SI 1978/1006 reg. 7(3) (as substituted (30.10.2005) by Medicines (Advisory Bodies) (No.2) Regulations 2005 (S.I. 2005/2754), reg. 1(2)(b), Sch. 4 para. 3(b))

23 Special provisions as to effect of manufacturer's licence.

- (1) Subject to ^{F89}...^{F90F89}... the following provisions of this section, a manufacturer's licence shall not have effect so as to authorise the manufacture or assembly of medicinal products of any description for sale or supply to any other person, or for exportation, unless either
 - the holder of the licence is also the holder of a product licence which is (a) applicable to medicinal products of that description, or
 - $[^{F91}(b)]$ the products are manufactured or assembled to the order of-
 - (i) a person who is the holder of such a product licence, or
 - (ii) if the products are to be used for the purposes of a clinical trial, the sponsor of that trial,]

and (in either case) the products are manufactured or assembled in accordance with that product licence.

- (2) ^{F92}... the preceding subsection shall not have effect in relation to the manufacture or assembly of any medicinal product to the order of a practitioner, where the practitioner-
 - (a) being a doctor or dentist, states that the product is required for administration to a patient of his or is required, at the request of another doctor or dentist, for administration to a patient of that other doctor or dentist, ^{F93}...
 - ^{F93}(b)

and shall not have effect in relation to the manufacture or assembly of any medicinal product to the order of a pharmacist in accordance with a prescription given by a practitioner.

- - (4) If by virtue of an order made under section 15 of this Act an exemption is conferred in respect of the restrictions imposed by section 7 of this Act, but no corresponding exemption is conferred in respect of the restrictions imposed by section 8(2) of this Act, the order may provide that subsection (1) of this section shall have effect subject to such exceptions or modifications as the Ministers consider appropriate in the circumstances.
 - (5) Where subsection (1) of this section has effect in relation to medicinal products of any description, and the conditions specified in that subsection are not fulfilled, the manufacture or assembly of medicinal products of that description for sale or supply to another person, or for exportation, notwithstanding that it complies with the provisions contained in the manufacturer's licence, shall for the purposes of this Act be deemed to be not in accordance with that licence.
- [^{F95}(6) In this section, "clinical trial" and "sponsor", in relation to a clinical trial, have the meaning given by Clinical Trials Regulations.]

Textual Amendments

Words in s. 23(1) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. F89 2006/2407), reg. 1, Sch. 8 para. 13(a) (with regs. 2(4), 3)

Words in s. 23(1) omitted (1.5.2004) by virtue of Medicines for Human Use (Clinical Trials) F90 Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 5(2)(a)

F91 S. 23(1)(b) substituted (1.5.2004) by Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 5(2)(b)

- **F92** Words in s. 23(2) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 13(b)(i) (with regs. 2(4), 3)
- **F93** S. 23(2)(b) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 13(b)(ii) (with regs. 2(4), 3)
- **F94** S. 23(3) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 13(c) (with regs. 2(4), 3)
- **F95** S. 23(6) inserted (1.5.2004) by Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 5(3)

Modifications etc. (not altering text)

- C60 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C61 S.23 extended (with modifications) (14.2.1994) by S.I. 1994/105, reg. 19, Sch.4
 - S. 23 modified (1.1.1995) by S.I. 1994/3144, reg.9(3)
 - S. 23 applied (1.1.1995) by S. I. 1994/3142, reg. 18(2)
 - S. 23 amended (E.W.S.) (*prosp.*) by 1954 c. 61, s. 13I para. 1(b) (as inserted (*prosp.*) by 1997 c. 19, ss. 1,2, Sch. para.2)
- C62 S. 23 modified (30.10.2005) by Medicines (Traditional Herbal Medicinal Products for Human Use) Regulations 2005 (S.I. 2005/2750), regs. 1(a), **10(2)** (with Sch. 6)
- C63 S. 23 amendment to earlier affecting provision SI 1994/3144 reg. 9 (30.10.2005) by Medicines (Marketing Authorisations Etc.) Amendment Regulations 2005 (S.I. 2005/2759), regs. 1(a), 2(12)

24 Duration and renewal of licence.

- [^{F96}(1) A licence granted under this Part expires—
 - (a) in accordance with the provisions of the licence, or
 - (b) if there is no such provision, at the end of the period of five years beginning with the date on which the licence was granted, or if it has been renewed the date on which it was last renewed.
- (1AA) But so far as the licence relates to a medicinal product to which the 2001 Directive applies, it remains in force until—
 - (a) revoked by the licensing authority; or
 - (b) surrendered by the holder.]
 - (2) Any [^{F97}licence granted under this Part of this Act], if it has not been revoked, may, on the application of the holder of the licence, be renewed by the licensing authority for a further period of five years from the date on which it would otherwise expire or such shorter period from that date as the licensing authority may determine.
- [^{F98}(2A) Subsection (2) does not apply to a licence insofar as it relates to a medicinal product to which the 2001 Directive applies.]
 - (3) On an application to the licensing authority for the renewal of a licence under this Part of this Act, the licensing authority—
 - (a) may renew the licence, with or without modifications, for such a further period as is mentioned in subsection (2) of this section, or
 - (b) may grant to the applicant a new licence containing such provisions as the licensing authority consider appropriate, or
 - (c) if, having regard to the provisions of this Act [^{F99} and any [^{F83}EU] obligation under [^{F100}the 2001 Directive other than Titles VI, VII and VIII of that Directive]], they consider it necessary or expedient to do so, may refuse to renew the licence or to grant a new licence.

- [^{F101}(3A) References to a licence in subsection (3) are to be read as references to a licence only insofar as that licence relates to a medicinal product to which the 2001 Directive does not apply.]
 - (4) In relation to any such application the provisions of sections 18 and 19, subsections (2) to (5) of section 20 and sections [^{F102}21 to 22A] of this Act shall have effect as if in those provisions any reference to refusing a licence included a reference to refusing to renew a licence and any reference to granting a licence included a reference to renewing it.
 - (5) Every application for the grant or renewal of a licence under this Part of this Act shall, unless it otherwise expressly provides, be taken to be an application for the grant or renewal of the licence for the full period of five years mentioned in subsection (1) or subsection (2) of this section, as the case may be; and in this Part of this Act any reference (including a reference implied by virtue of the last preceding subsection) to the grant or renewal of a licence otherwise than in accordance with the application shall be construed accordingly.
- [^{F103}(5A) Subsection (5) does not apply to a licence insofar as it relates to a medicinal product to which the 2001 Directive applies.]
 - (6) Where an application for the renewal of a licence under this Act has been duly made—
 - (a) the licence shall not cease to be in force by virtue of the preceding provisions of this section before the licensing authority have determined the application, and
 - (b) if by an interim order made under section 107(3)(a) of this Act the operation of the decision of the licensing authority on the application is suspended, the licence shall not cease to be in force by virtue of those provisions so long as the operation of the decision continues to be suspended by the order.

^{F104}(7)...

Textual Amendments

- **F83** Words in Act substituted (22.4.2011) by The Treaty of Lisbon (Changes in Terminology) Order 2011 (S.I. 2011/1043), arts. 2, 3, 6 (with art. 3(2)(3)4(2)6(4)6(5))
- F96 S. 24(1)(1AA) substituted for s. 24(1) (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 4(2) (with Sch. 6)
- F97 Words substituted by (E.W.)(S.) S.I. 1977/1050, art. 4(4) and (N.I.) S.R. 1977 No. 170, reg. 5(4)
- **F98** S. 24(2A) inserted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 4(3) (with Sch. 6)
- **F99** Words in s. 24(3)(c) inserted (13.2.1994) by S.I. 1994/276, **reg. 5(b)** (which S.I. revoked and replaced defective S.I. 1994/101 before the latter S.I. came into force)
- F100 Words in "s. 24(3)(C)" substituted (28.2.2002) by virtue of S.I. 2002/236, reg. 2(c)(ii)
- **F101** S. 24(3A) inserted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 4(4) (with Sch. 6)
- F102 Words in s. 24(4) substituted (30.10.2005) by Medicines (Advisory Bodies) Regulations 2005 (S.I. 2005/1094), reg. 1(1), Sch. 1 para. 4

Changes to legislation: Medicines Act 1968, Part II is up to date with all changes known to be in force on or before 27 June 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details)

F103 S. 24(5A) inserted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 4(5) (with Sch. 6)

F104 S. 24(7) deleted (28.2.2002) by virtue of S.I. 2002/236, reg. 2(c)(ii)

Modifications etc. (not altering text)

C64 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

C65 S. 24 applied (with modifications) (3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.

Licences of right

25 Entitlement to licence of right.

(4) In this Act "licence of right" means a licence to which a person is entitled by virtue of this section, including such a licence which has been renewed (with or without modifications) but not a licence granted instead of the renewal of such a licence.

Textual Amendments

F105 S. 25(1)-(3) repealed (22.7.2004) by Statute Law (Repeals) Act 2004 (c. 14), Sch. 1 Pt. 17 Group 7

Modifications etc. (not altering text)

C66 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

^{F106}26 Scope of licence of right in different cases.

Textual Amendments

F106 S. 26 repealed (22.7.2004) by Statute Law (Repeals) Act 2004 (c. 14), Sch. 1 Pt. 17 Group 7

Modifications etc. (not altering text)

C67 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

27 **Proceedings on application for licence of right.**

- (1) Sections 19 to [^{F107}22A] of this Act shall not have effect in relation to any application for a licence of right.
- (2) If on any such application the licensing authority—
 - (a) propose to refuse to grant a licence on that application, on the grounds that none of the provisions of subsections (2) to (5) of section 16 of this Act has been proved to have effect in relation to the applicant, or

(b) propose to grant a licence which will not extend to some of the matters specified in the application.

the licensing authority shall, before the end of the period of three months from the date on which the application is received by them, serve on the applicant a notice stating their proposals and the reasons for them and, in a case falling within paragraph (b) of this subsection, the matters specified in the application to which it is proposed that the licence should not extend.

- (3) If, within the time allowed after the service of a notice under subsection (2) of this section, the applicant gives notice to the licensing authority of his desire to be heard under this subsection or makes representations in writing to the licensing authority with respect to their proposals, then, before determining the application, the licensing authority shall afford to him an opportunity of appearing before, and being heard by, a person appointed for the purpose by the licensing authority, or shall take those representations into account, as the case may be.
- (4) Where the applicant avails himself of the opportunity afforded to him in pursuance of subsection (3) of this section or makes representations in writing as mentioned in that subsection, then if—
 - (a) the licensing authority refuse to grant a licence on the application, or
 - (b) grant a licence otherwise than in accordance with the application, and the applicant requests the licensing authority to state their reasons.

the licensing authority shall serve on the applicant a notice stating the reasons for their decision.

- (5) If, in a case where the licensing authority have served a notice under subsection (2) of this section, the application is not finally disposed of before the date which, in relation to any matters specified in the application, is the relevant date, then on and after that date, and until the application has been finally disposed of, the provisions of this Act shall have effect in relation to those matters as if the licensing authority had granted a licence of right in accordance with the application.
- (6) Where, on an application for a licence of right, the licensing authority do not serve a notice under subsection (2) of this section before the end of the period mentioned in that subsection, the licensing authority shall be required to grant a licence in accordance with sections 25 and 26 of this Act as if all the matters specified in the application had been proved; and if such a licence has not been granted before the date which, in relation to any of those matters, is the relevant date, the provisions of this Act shall have effect on and after that date in relation to those matters as if the licensing authority had granted a licence of right in accordance with the application.
- (7) For the purposes of this section the relevant date, in relation to any matters specified in an application, is the date on which, in accordance with one or more orders made under section 17 of this Act, that subsection of section 16 of this Act which has effect in relation to those matters ceases to have effect in relation to them; and an application shall for the purposes of this section be taken to be finally disposed of on (but not before) the occurrence of whichever of the following events last occurs, that is to say—
 - (a) the licensing authority make a decision determining the application;
 - (b) the time within which an application under section 107 of this Act with respect to that decision can be made expires without its having been made;
 - (c) if such an application under section 107 of this Act is made, the proceedings on the application under that section are finally determined or abandoned or otherwise disposed of;

- (d) if there is an appeal against the decision in any such proceedings as are mentioned in paragraph (c) of this subsection, or an appeal against the decision on such an appeal, the proceedings on that appeal are finally determined or abandoned or otherwise disposed of;
- (e) the time for bringing any such appeal as is mentioned in paragraph (d) of this subsection expires without its having been brought.

[^{F108}(8) Subsections (2), (8) and (10)(b) of section 22A of this Act shall have effect in relation to a person appointed under subsection (3) of this section and to proceedings before him and his report as they have effect for the purposes of that section.]

Textual Amendments

F107 Word in s. 27(1) substituted (30.10.2005) by Medicines (Advisory Bodies) Regulations 2005 (S.I. 2005/1094), reg. 1(1), Sch. 1 para. 5(a)

F108 S. 27(8) substituted (30.10.2005) by Medicines (Advisory Bodies) Regulations 2005 (S.I. 2005/1094), reg. 1(1), Sch. 1 para. 5(b)

Modifications etc. (not altering text)

- C68 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C69 Power to exclude s. 27(5)(6) conferred by Medicines Act 1971 (c. 69), s. 1(2)(a)

Suspension, revocation and variation of licences

28 General power to suspend, revoke or vary licences.

- (1) Subject to the following provisions of this Part of this Act, the licensing authority may suspend a licence under this Part of this Act for such period as the authority may determine, or may revoke, or vary the provisions of, any such licence.
- (2) The suspension or revocation of a licence under this section may be total or may be limited to medicinal products of one or more descriptions or to medicinal products manufactured, assembled or stored on any particular premises or in a particular part of any premises.
- (3) [^{F109}Subject to subsection (3A) of this section]the powers conferred by this section shall not be exercisable by the licensing authority in relation to a product licence except on one or more of the following grounds, that is to say—
 - (a) that the matters stated in the application on which the licence was granted were false or incomplete in a material particular;
 - (b) that any of the provisions of the licence has to a material extent been contravened by the holder of the licence or by a person procured by him to manufacture or assemble medicinal products of a description to which the licence relates;
 - (c) that medicinal products of any such description, as sold, supplied, exported, imported, manufactured or assembled in pursuance of the licence, fail to a material extent to correspond to the characteristics by reference to which the licence was granted;
 - (d) that the holder of the licence has without reasonable excuse failed to comply with a requirement imposed on him under section 44(2) of this Act to furnish

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information to the licensing authority with respect to medicinal products of any such description;

- (e) that any premises on which, or in part of which, medicinal products of any such description are manufactured, assembled or stored by or on behalf of the holder of the licence are unsuitable;
- (f) in the case of a licence other than a licence of right, that the holder of the licence has not, within two years after the grant of the licence, notified to the licensing authority, in relation to each description of medicinal products to which the licence relates, a date on which medicinal products of that description were effectively on the market in the United Kingdom;
- (g) that medicinal products of any description to which the licence relates can no longer be regarded as products which can safely be administered for the purposes indicated in the licence, or can no longer be regarded as efficacious for those purposes;
- (h) that the specification and standards to which medicinal products of any such description are manufactured can no longer be regarded as satisfactory.

^{F110}(i) ...

- [^{F111}(j) that, in relation to medicinal products of any description to which the licence relates [^{F112}(other than products to which [^{F113}the 2001 Directive applies])] any of the provisions contained in regulations which—
 - (i) are made under section 85 of this Act (labelling and marking of containers and packages), and
 - (ii) impose requirements which give effect to [^{F83}EU] obligations,

has to a material extent been contravened by the holder of the licence or by a person procured by him to manufacture or assemble such medicinal products.]

- [^{F114}(3A) Where a product licence relates to a product to which [^{F113}the 2001 Directive applies], the power conferred by this section to suspend a licence shall be exercisable in relation to the licence on the ground that—
 - (a) any of the provisions contained in regulations made under section 85 (labelling and marking of containers and packages) or 86 (leaflets) of this Act, or
 - (b) section 86(4),

has to a material extent been contravened in relation to the product by the holder of the licence or by a person procured by him to manufacture or assemble the product.]

- (4) Subject to the following provisions of this section, the powers conferred by this section shall not be exercisable in relation to a manufacturer's licence or a wholesale dealer's licence except on one or more of the following grounds, that is to say—
 - (a) that the matters stated in the application on which the licence was granted were false or incomplete in a material particular;
 - (b) that a material change of circumstances has occurred in relation to any of those matters;
 - (c) that any of the provisions of the licence has to a material extent been contravened by the holder of the licence;
 - (d) that the holder of the licence has without reasonable excuse failed to comply with a requirement imposed on him under section 44(2) of this Act to furnish information to the licensing authority with respect to medicinal products of a description to which the licence relates.

- (5) In relation to a manufacturer's licence, the powers conferred by this section shall be exercisable on either of the following grounds, in addition to those specified in subsection (4) of this section, that is to say—
 - (a) that the holder of the manufacturer's licence has carried out processes of manufacture or assembly to the order of another person who is the holder of a product licence, and has habitually failed to comply with the provisions of that product licence;
 - (b) that the holder of the manufacturer's licence does not have the requisite facilities for carrying out properly processes of manufacture or assembly authorised by the licence.
- (6) In relation to a wholesale dealer's licence, the powers conferred by this section shall be exercisable on the following grounds, in addition to those specified in subsection (4) of this section, that is to say, that the equipment and facilities for storing or distributing medicinal products which are available to the holder of the licence are inadequate to maintain the quality of medicinal products of one or more descriptions to which the application for the licence related.
- (7) The preceding provisions of this section shall have effect subject to the next following section.

Textual Amendments

- **F83** Words in Act substituted (22.4.2011) by The Treaty of Lisbon (Changes in Terminology) Order 2011 (S.I. 2011/1043), arts. 2, 3, 6 (with art. 3(2)(3)4(2)6(4)6(5))
- **F109** Words in s. 28(3) inserted (13.2.1994) by S.I. 1994/276, reg. 6(2)(a) (with reg. 6(4)) (which S.I. revoked and replaced defective S.I. 1994/101 before the latter S.I. came into force)
- **F110** S. 28(3)(i) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 14 (with regs. 2(4), 3)
- F111 S. 28(3)(j) inserted by (E.W.)(S.) S.I. 1977/1050, art. 4(5) and (N.I.) S.R. 1977 No. 170, reg. 5(5)
- **F112** Words in s. 28(3)(j) inserted (13.2.1994) by S.I. 1994/276, reg. 6(2)(b) (with reg. 6(4))(which S.I. revoked and replaced defective S.I. 1994/101 before the latter S.I, came into force)
- F113 Words in s. 28(3)(j)(3A) substituted (28.2.2002) by S.I. 2002/236, reg. 2(a)(iii)
- F114 S. 28(3A) inserted (13.2.1994) by S.I. 1994/276, reg. 6(3) (with reg. 6(4)) (which S.I. revoked and replaced defective S.I. 1994/101 before the latter S.I. came into force)

Modifications etc. (not altering text)

- C70 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C71 S. 28 (1)(2)(3)(7) applied (with modifications)(3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.

29 Procedure where licensing authority propose to suspend, revoke or vary licence under s. 28.

- (1) The provisions of Schedule 2 to this Act shall have effect where the licensing authority propose to exercise any power conferred by section 28 of this Act.
- (2) Without prejudice to any requirement of that Schedule as to the service of notices, where in the exercise of any such power the licensing authority suspend, revoke or vary a licence, they shall serve on the holder of the licence a notice giving particulars of the suspension, revocation or variation and of the reasons for their decision to suspend, revoke or vary the licence.

Modifications etc. (not altering text)

- C72 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C73 S. 29 applied (with modifications) (3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch. s. 29 extended (with modifications) (14.2.1994) by S.I. 1994/105, reg. 19, Sch.4
- C74 S. 29 amendment to earlier affecting provision SI 1994/105 Sch. 4 (30.10.2005) by Medicines (Homoeopathic Medicinal Products for Human Use) Amendment Regulations 2005 (S.I. 2005/2753), regs. 1(1), 19(2)

[^{F115}30 Variation of licence on application of holder.

- (1) This section applies if the holder of a licence under this Part applies to the licensing authority for the licence to be varied.
- (2) The application must—
 - (a) be in writing,
 - (b) specify the required variation,
 - (c) be signed by or on behalf of the applicant,
 - (d) be accompanied by such information as is reasonably required to enable the licensing authority to consider the application, and
 - (e) if there is a requirement in force under section 1(1)(a) of the Medicines Act 1971 to pay a fee in respect of the application, be accompanied by the required fee.
- (3) The licensing authority must consider any application properly made under this section.
- (4) If subsection (5) applies, they must either vary the licence or refuse to vary it before the end of the period allowed for considering the application.
- (5) This subsection applies to a variation which would have the effect of altering—
 - (a) the types of medicinal product,
 - (b) any operation carried out under the licence,
 - (c) any premises, or
 - (d) any equipment or facilities,
 - in respect of which the licence was granted.
- (6) If the licensing authority considers that it is necessary for them to conduct an inspection of any premises to which the application relates, the period allowed is 90 days beginning with the date on which they receive the application.
- (7) Otherwise, the period allowed is 90 days beginning with that date.
- (8) The licensing authority may give the applicant written notice requiring him to give them such further information in connection with the application as they consider reasonable.
- (9) The period allowed for consideration stops running when a notice is given under paragraph (8) and does not start running again until—
 - (a) the licensing authority receives the information; or
 - (b) the applicant has shown to the reasonable satisfaction of the licensing authority why he is unable to provide it

Changes to legislation: Medicines Act 1968, Part II is up to date with all changes known to be in force on or before 27 June 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details)

(10) Nothing in this section affects the powers conferred by section 28.]

Textual Amendments

F115 S. 30 substituted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 5 (with Sch. 6)

Modifications etc. (not altering text)

- C75 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C76 S. 30 applied (with modifications) (3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.

Clinical trials and medicinal tests on animals

^{F116}31 Clinical trials.

Textual Amendments

F116 S. 31 omitted (1.5.2004) by virtue of Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 6

Modifications etc. (not altering text)

C77 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

^{F117}32 Medicinal tests on animals.

Textual Amendments

F117 Ss. 32-36 omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 15 (with regs. 2(4), 3)

Modifications etc. (not altering text)

- C78 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C79 Ss. 32-39 modified (1.1.1995) by S.I. 1994/3142, reg. 18(4)
- C80 S. 32 applied (with modifications) (2.8.1999) by S.I. 1999/1871, reg. 92(3)

F11733 Exemptions in respect of medicinal tests on animals.

Textual Amendments

F117 Ss. 32-36 omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 15 (with regs. 2(4), 3)

Modifications etc. (not altering text)

- C81 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- **C82** Ss. 32-39 modified (1.1.1995) by S.I. 1994/3142, reg. 18(4)
- C83 S. 33 amended (E.W.S.) (prosp.) by 1954 c. 61, s. 13I(1)(b) (as inserted (prosp.) by 1997 c. 19, ss. 1, 2, Sch. para.2)
- C84 S. 33 applied (with modifications) (2.8.1999) by S.I. 1999/1871, reg. 92(3)

F11734 Restrictions as to animals on which medicinal tests have been carried out.

Textual Amendments

F117 Ss. 32-36 omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 15 (with regs. 2(4), 3)

Modifications etc. (not altering text)

- C85 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C86 Ss. 32-39 modified (1.1.1995) by S.I. 1994/3142, reg. 18(4)
- C87 S. 34 applied (with modifications) (2.8.1999) by S.I. 1999/1871, reg. 92(3)

^{F117}35 Supplementary provisions as to clinical trials and medicinal tests on animals.

Textual Amendments

F117 Ss. 32-36 omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 15 (with regs. 2(4), 3)

Modifications etc. (not altering text)

C88 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

C89 S. 35 (other than subsection (8)(a)) applied (with modifications) (2.8.1999) by S.I. 1999/1871, reg. 92(3)

^{F117}36 Application for, and issue of, certificate.

Textual Amendments

F117 Ss. 32-36 omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 15 (with regs. 2(4), 3)

Modifications etc. (not altering text)

- C90 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- **C91** Ss. 32-39 modified (1.1.1995) by S.I. 1994/3142, reg. 18(4)
- C92 S. 36 applied (with modifications) (2.8.1999) by S.I. 1999/1871, reg. 92(3)

^{F118}37 Transitional provisions as to clinical trials and medicinal tests on animals.

Textual Amendments

F118 S. 37 repealed (22.7.2004) by Statute Law (Repeals) Act 2004 (c. 14), Sch. 1 Pt. 17 Group 7

Modifications etc. (not altering text)

- C93 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C94 Ss. 32-39 modified (1.1.1995) by S.I. 1994/3142, reg. 18(4)

^{F119}38 Duration and renewal of certificate.

Textual Amendments

F119 Ss. 38-40 omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 15 (with regs. 2(4), 3)

Modifications etc. (not altering text)

C95 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

- C96 Ss. 32-39 modified (1.1.1995) by S.I. 1994, reg. 18(4)
- C97 S. 38 applied (with modifications) (2.8.1999) by S.I. 1999/1871, art. 92(3)

^{F119}39 Suspension, revocation or variation of certificate.

Textual Amendments

F119 Ss. 38-40 omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 15 (with regs. 2(4), 3)

Modifications etc. (not altering text)

C98 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

C99 Ss. 32-39 modified (1.1.1995) by S.I. 1994/3142, reg. 18(4)

C100 S. 39 applied (with modifications) (2.8.1999) by S.I. 1999/1871, reg. 92(3)

Medicated animal feeding stuffs

^{F119}40 Medicated animal feeding stuffs.

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Textual Amendments

F119 Ss. 38-40 omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 15 (with regs. 2(4), 3)

Modifications etc. (not altering text)

C101 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1) C102 S. 40 applied (1.1.1995) by S.I. 1994/3142, reg. 18(2)

41–42^{F120}

Textual Amendments

F120 Ss. 41, 42 repealed by Animal Health and Welfare Act 1984 (c. 40, SIF 2:8), s. 16, Sch. 1 para. 3(3), Sch. 2

Supplementary provisions

43 Extension of s. 7 to certain special circumstances.

- (1) Where in the course of a business carried on by him a person sells, supplies or exports a substance or article for use wholly or mainly in either or both of the ways specified in section 130(1) of this Act, and the substance or article, not having been—
 - (a) manufactured or imported for such use, or
 - (b) previously sold or supplied for such use,

does not constitute a medicinal product before that person so sells, supplies or exports it, then (subject to subsection (2) of this section) subsection (2) of section 7 of this Act, if apart from this subsection it would not so have effect, shall have effect in relation to the sale, supply or exportation of the substance or article as if he were selling, supplying or exporting it in circumstances to which that subsection applies.

- (2) Subsection (1) of this section shall not have effect in relation to a transaction whereby a person, in the course of a business carried on by him, sells a substance or article by retail or supplies a substance or article in circumstances corresponding to retail sale unless in the course of that business the substance or article has been assembled for the purpose of being sold or supplied by him.
- (3) In any reference in this Part of this Act to the provisions of, or the restrictions imposed by, section 7 of this Act, the reference to that section shall be construed as including a reference to subsection (2) of that section as extended by the preceding subsections.
- (4) Where in the course of a business carried on by him a person proposes to sell, supply or export a substance or article for use as mentioned in subsection (1) of this section, where the substance or article will not constitute a medicinal product before he so sells, supplies or exports it and he will not be selling, supplying or exporting it in circumstances to which section 7(2) of this Act applies, he may, if he so desires, apply for a product licence in respect of that substance or article, and the licensing authority (subject to the provisions of sections 19 to [^{F121}22A] of this Act) may grant to him a product licence in respect of it, as if he were proposing to sell, supply or export it

Status: Point in time view as at 28/10/2011. Changes to legislation: Medicines Act 1968, Part II is up to date with all changes known to be in force on or

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in circumstances to which section 7(2) of this Act applies; and a product licence so granted may be renewed, suspended, revoked or varied accordingly.

(5) In subsection (2) of this section the reference to assembling a substance or article in the course of a business carried on by a person is a reference to doing in the course of that business anything which (in accordance with section 132(1) of this Act) would constitute assembling if it had been a medicinal product when sold or supplied to him.

Textual Amendments

F121 Word in s. 43(4) substituted (30.10.2005) by Medicines (Advisory Bodies) Regulations 2005 (S.I. 2005/1094), reg. 1(1), Sch. 1 para. 8

Modifications etc. (not altering text)

C103 Pt. II(ss. 6-50) extended with modifications by S.I. 1985/1403, art. 3(1)

44 **Provision of information to licensing authority.**

- (1) Where an application has been made to the licensing authority for a licence under this Part of this Act (including a licence of right) F122... the licensing authority, before determining the application, may request the applicant to furnish to the licensing authority such information relating to the application as the licensing authority may consider requisite; and, where any such request has been made, the licensing authority shall not be required to determine the application until either—
 - (a) the information requested has been furnished to them, or
 - (b) it has been shown to their reasonable satisfaction that the applicant is unable to furnish the information.
- (2) The licensing authority may serve on the holder of a licence under this Part of this Act^{F123}... a notice requiring him, within such time as may be specified in the notice, to furnish to the licensing authority information of any description specified in the notice in accordance with the following provisions of this section.
- (3) Except as provided by subsection (4) of this section, a notice under subsection (2) of this section shall not be served unless it appears to the licensing authority, or it is represented to them ^{F124}... by the appropriate committee, that circumstances exist by reason of which it is necessary to consider whether the licence ^{F125}... should be varied, suspended or revoked; and the information required by such a notice shall be such as appears to the licensing authority, or is represented to them ^{F124}... by the committee, to be requisite for considering that question.
- (4) Subsection (3) of this section shall not have effect in the case of a licence of right^{F126}... whether the licence ^{F127}... has been renewed or not; and, in the case of such a licence ^{F127}..., a notice under this section may be served at any time and may require any information which, in the opinion of the licensing authority, would be relevant if—
 - (a) [^{F128} section 25] of this Act had not been enacted, and
 - (b) the licensing authority were then dealing with an application, by the person who is the holder of the licence ^{F127}..., for the grant ^{F129}... of a licence ^{F127}... containing the same provisions as those contained in the licence ^{F127}... in question.
- (5) Before the end of the period of two years from the date on which a product licence, other than a licence of right, is granted, the holder of the licence shall, in respect of each

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description of medicinal products to which the licence relates which is effectively on the market in the United Kingdom within that period, notify to the licensing authority a date on which medicinal products of that description were effectively on that market.

Textual Amendments

- **F122** Words in s. 44(1) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 16(a) (with regs. 2(4), 3)
- **F123** Words in s. 44(2) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 16(b) (with regs. 2(4), 3)
- F124 Words in s. 44(3) omitted (30.10.2005) by virtue of Medicines (Advisory Bodies) Regulations 2005 (S.I. 2005/1094), reg. 1(1), Sch. 1 para. 9
- F125 Words in s. 44(3) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 16(c) (with regs. 2(4), 3)
- **F126** Words in s. 44(4) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 16(d)(i) (with regs. 2(4), 3)
- F127 Words in s. 44(4) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 16(d)(ii) (with regs. 2(4), 3)
- **F128** Words in s. 44(4)(a) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 16(d)(iii) (with regs. 2(4), 3)
- **F129** Words in s. 44(4)(b) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 16(d)(iv) (with regs. 2(4), 3)

Modifications etc. (not altering text)

C104 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

- C105 S. 44 extended (with modifications) (14.2.1994) by S.I. 1994/105, reg. 19, Sch.4
- C106 S. 44 amendment to earlier affecting provision SI 1994/105 Sch. 4 (30.10.2005) by Medicines (Homoeopathic Medicinal Products for Human Use) Amendment Regulations 2005 (S.I. 2005/2753), regs. 1(1), 19(3)
- C107 S. 44(1)(2)(3) applied (with modifications) (3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.

45 Offences under Part II.

- (1) Subject to the next following section, any person who contravenes any of the provisions of section 7 [^{F130}or section 8] of this Act, or who is in possession of any medicinal product ^{F131}... for the purpose of selling, supplying or exporting it in contravention of [^{F132}either of those sections], shall be guilty of an offence.
- (2) Where any medicinal product ^{F133}... is imported in contravention of section 7 ^{F134}... F¹³⁵... of this Act, any person who, otherwise than for the purpose of performing or exercising a duty or power imposed or conferred by or under this Act or any other enactment, is in possession of the product ^{F136}... knowing or having reasonable cause to suspect that it was so imported shall be guilty of an offence.
- (3) Any person who, being the holder of a product licence ^{F137}..., procures another person to carry out a process in the manufacture or assembly of medicinal products of a description to which the licence ^{F138}... relates, and—
 - (a) does not communicate to that person the provisions of the licence ^{F138}... which are applicable to medicinal products of that description, or
 - (b) in a case where any of those provisions has been varied by a decision of the licensing authority, does not communicate the variation to that person within fourteen days after notice of the decision has been served on him,

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shall be guilty of an offence.

- - (6) Any person who, in giving any information which he is required to give under section 44 of this Act, makes a statement which he knows to be false in a material particular shall be guilty of an offence.
 - (7) Any person who without reasonable excuse fails to comply with a requirement imposed on him by a notice under section 44(2) of this Act shall be guilty of an offence.
 - (8) Any person guilty of an offence under any of subsections (1) to (6) of this section shall be liable—
 - (a) on summary conviction, to a fine not exceeding £400;
 - (b) on conviction on indictment, to a fine or to imprisonment for a term not exceeding two years or to both.
 - (9) Any person guilty of an offence under subsection (7) of this section shall be liable on summary conviction to a fine not exceeding [^{F140}level 3 on the standard scale]

Textual Amendments

- **F130** Words in s. 45(1) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 17(a)(i) (with regs. 2(4), 3)
- **F131** Words in s. 45(1) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 17(a)(ii) (with regs. 2(4), 3)
- **F132** Words in s. 45(1) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 17(a)(iii) (with regs. 2(4), 3)
- **F133** Words in s. 45(2) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 17(b)(i) (with regs. 2(4), 3)
- **F134** Words in s. 45(2) omitted (1.5.2004) by virtue of Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 13(a)
- F135 Words in s. 45(2) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 17(b)(ii) (with regs. 2(4), 3)
- F136 Words in s. 45(2) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 17(b)(iii) (with regs. 2(4), 3)
- F137 Words in s. 45(3) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 17(c)(i) (with regs. 2(4), 3)
- **F138** Words in s. 45(3) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 17(c)(ii) (with regs. 2(4), 3)
- F139 S. 45(4)(5) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 17(d) (with regs. 2(4), 3)
- F140 Words substituted by virtue of (E.W.) Criminal Justice Act 1982 (c. 48, SIF 39:1), ss. 38, 46, (S.) Criminal Procedure (Scotland) Act 1975 (c.21, SIF 39:1), ss. 289F, 289G and (N.I.) S.I. 1984/703 (N.I. 3), arts. 5, 6

Modifications etc. (not altering text)

C108 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

- C109 s. 45 extended (with modifications) (14.2.1994) by S.I. 1994/105, reg. 19, Sch.4
- C110 S. 45 amendment to earlier affecting provision SI 1994/105 Sch. 4 (30.10.2005) by Medicines (Homoeopathic Medicinal Products for Human Use) Amendment Regulations 2005 (S.I. 2005/2753), regs. 1(1), 19(4)

C111 S. 45(1)(2)(6)(7)(8)(9) applied (with modifications) (3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.

46 Special defences under s. 45.

- (1) Where the holder of a product licence ^{F141}... is charged with an offence under the last preceding section in respect of any substance or article which has been manufactured (or, in the case of a medicinal product, manufactured or assembled) to his order by another person and has been so manufactured or assembled as not to comply with the provisions of that licence ^{F142}... which are applicable to it, it shall be a defence for him to prove—
 - (a) that he had communicated those provisions to that other person, and
 - (b) that he did not know, and could not by the exercise of reasonable care have discovered, that those provisions had not been complied with.
- (2) Where the holder of a manufacturer's licence is charged with an offence under the last preceding section in respect of any medicinal products which have been manufactured or assembled by him, in circumstances where he is not the holder of a product licence ^{F143}... which is applicable to those products, but the products were manufactured or assembled to the order of another person, it shall be a defence for him to prove that he believed, and had reasonable grounds for believing,—
 - (a) that the other person in question was the holder of a product licence applicable to those products, ^{F144}..., and
 - (b) that the products were manufactured or assembled in accordance with that product licence F145

Textual Amendments

- **F141** Words in s. 46(1) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 18(a)(i) (with regs. 2(4), 3)
- **F142** Words in s. 46(1) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 18(a)(ii) (with regs. 2(4), 3)
- **F143** Words in s. 46(2) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 18(b)(i) (with regs. 2(4), 3)
- **F144** Words in s. 46(2)(a) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 18(b)(ii) (with regs. 2(4), 3)
- F145 Words in s. 46(2)(b) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 18(b)(iii) (with regs. 2(4), 3)
- **F146** S. 46(3)(4) repealed by Animal Health and Welfare Act 1984 (c. 40, SIF 2:8), s. 16, Sch. 1 para. 3(3), Sch. 2

Modifications etc. (not altering text)

- C112 Pt. II(ss. 6-50) extended with modifications by S.I. 1985/1403, art. 3(1)
- C113 S. 46 extended (with modifications) (14.2.1994) by S.I. 1994/105, reg. 19, Sch.4
- C114 S. 46 amendment to earlier affecting provision SI 1994/105 Sch. 4 (30.10.2005) by Medicines (Homoeopathic Medicinal Products for Human Use) Amendment Regulations 2005 (S.I. 2005/2753), regs. 1(1), 19(2)
- C115 S. 46(1) applied (wth modifications)(3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.

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47 [^{F147}Standard provisions for licences]

- (1) The Ministers may by regulations prescribe standard provisions for the purposes of this Part of this Act, either generally or in relation to any class of medicinal products specified in the regulations.
- (2) Any standard provisions so prescribed may be incorporated by the licensing authority in any licence under this Part of this Act ^{F148}...^{F149F148}... granted ^{F148}... on or after the date on which the regulations come into operation, and may be so incorporated with or without modifications and either generally or in relation to medicinal products of any particular class.

(3) The following provisions of this section shall have effect where—

- (a) standard provisions are prescribed by regulations made under this section, or
- (b) after any such provisions have been so prescribed, they are amended by, or superseded by new standard provisions prescribed by, subsequent regulations so made;

and in the following provisions of this section, in a case falling within paragraph (a) but not within paragraph (b) of this subsection, "the operative standard provisions" means the standard provisions prescribed by the regulations and "the relevant regulations" means those regulations, and, in any other case, "the operative standard provisions" means the standard provisions as amended by the subsequent regulations or the new standard provisions prescribed by those regulations, as the case may be, and "the relevant regulations" means the subsequent regulations.

- (4) Subject to the following provisions of this section, as from the end of the period of three months from the date on which the relevant regulations come into operation, the operative standard provisions shall be deemed to be incorporated in any licence under this Part of this Act^{F150}...^{F151F150}... which is in force at the end of that period or, in the case of a suspended licence ^{F152}..., would then be in force if it were not suspended, in so far as, in accordance with the relevant regulations, the operative standard provisions are applicable to medicinal products of any description to which that licence ^{F152}... relates.
- (5) Notwithstanding anything in subsection (4) of this section, the operative standard provisions shall not by virtue of that subsection be deemed to be incorporated in any licence of right^{F153}... including any such licence ^{F154}... which has been renewed, except in circumstances where, immediately before the first appointed day, the manufacture or importation of substances or articles to which the licence ^{F154}... relates was authorised by a licence issued under Part I of the ^{M1}Therapeutic Substances Act 1956 or under Part II of the ^{M2}Diseases of Animals Act 1950, or of the ^{M3}Diseases of Animals Act (Northern Ireland) 1958, and, where those circumstances exist, shall be deemed to be so incorporated only in relation to substances or articles to which the licence so issued was applicable.
- (6) At any time after the relevant regulations are made and before the end of the period of three months from the date on which they come into operation, the holder of any licence ^{F155}... may apply to the licensing authority to direct—
 - (a) that the operative standard provisions shall not be deemed to be incorporated in that licence ^{F155}..., or
 - (b) that the operative standard provisions shall be deemed to be so incorporated subject to such exceptions or modifications as may be specified in the application;

and if, on any such application, the licensing authority direct that the operative standard provisions shall not be deemed to be so incorporated, or shall be deemed to be so incorporated subject to exceptions and modifications specified in the direction, with or without provision postponing the date as from which they are to be deemed to be so incorporated, that direction shall have effect notwithstanding anything in subsection (4) of this section.

- (7) Where an application is made to the licensing authority under subsection (6) of this section, then, if the licensing authority propose to refuse to give a direction in accordance with the application, the licensing authority, before determining the application, shall afford to the applicant an opportunity of appearing before, and being heard by, a person appointed for the purpose by the licensing authority, or of making representations in writing to the licensing authority with respect to that proposal; and, if the licensing authority then determine to refuse to give a direction in accordance with the application, they shall serve on the applicant a notice stating the reasons for their decision.
- (8) Without prejudice to any direction given under subsection (6) of this section, where such an application is made—
 - (a) the operative standard provisions shall not be deemed to be incorporated in the licence ^{F156}... to which the application relates before the licensing authority have made a decision on that application, and
 - (b) if an application under section 107 of this Act is made with respect to that decision, those provisions shall not be deemed to have been or to be so incorporated before the application under subsection (6) of this section has been finally disposed of;

and so much of subsection (7) of section 27 of this Act as relates to the time when an application is to be taken to be finally disposed of shall have effect for the purposes of this subsection as it has effect for the purposes of that section.

(9) The powers conferred on the licensing authority by the preceding provisions of this Part of this Act to vary the provisions of a licence ^{F157}... shall be exercisable with respect to any provisions which, in accordance with this section, are incorporated or deemed to be incorporated in a licence ^{F157}....

Textual Amendments

- F147 Words in s. 47 heading substituted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 19
- **F148** Words in s. 47(2) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 19(a) (with regs. 2(4), 3)
- F149 Words in s. 47(2) omitted (1.5.2004) by virtue of Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 15
- **F150** Words in s. 47(4) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 19(b)(i) (with regs. 2(4), 3)
- F151 Words in s. 47(4) omitted (1.5.2004) by virtue of Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 15
- **F152** Words in s. 47(4) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 19(b)(ii) (with regs. 2(4), 3)
- **F153** Words in s. 47(5) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 19(c)(i) (with regs. 2(4), 3)
- F154 Words in s. 47(5) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 19(c)(ii) (with regs. 2(4), 3)

Changes to legislation: Medicines Act 1968, Part II is up to date with all changes known to be in force on or before 27 June 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details)

- F155 Words in s. 47(6) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 19(d) (with regs. 2(4), 3)
- **F156** Words in s. 47(8) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 19(e) (with regs. 2(4), 3)
- F157 Words in s. 47(9) omitted (1.10.2006) by virtue of The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 19(f) (with regs. 2(4), 3)

Modifications etc. (not altering text)

C116 Pt. II(ss. 6-50) extended with modifications by S.I. 1985/1403, art. 3(1)

- C117 S. 47(1)(2)(3)(4)(6)(7) applied (with modifications) (3.4.1992) by S.I. 1992/605, reg. 2(1)(2), Sch.
- C118 S. 47(4) excluded (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 6 para. 1(2)(c) (i)Sch. 6 para. 3(2)(a)Sch. 6 para. 3(3)(a) (with Sch. 6)
- C119 S. 47(6) applied (with modifications) (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 6 para. 1Sch. 6 para. 3(2)Sch. 6 para. 3(3) (with Sch. 6)
- C120 S. 47(7) modified (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 6 para. 1Sch. 6 para. 3(2)Sch. 6 para. 3(3) (with Sch. 6)
- C121 S. 47(8) excluded (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 6 para. 1Sch. 6 para. 3(2)Sch. 6 para. 3(3) (with Sch. 6)

Marginal Citations

- M1 1956 c. 25.
- **M2** 1950 c. 36.
- M3 1958 c. 13 (N.I.)

48 **Postponement of restrictions in relation to exports.**

- (1) Notwithstanding anything in sections 7 to 47 of this Act but subject to [^{F158}[^{F159}section 49] of this Act,] in relation to anything done before such day (subsequent to the first appointed day) as the Ministers may by order appoint for the purposes of this subsection (in this section referred to as "the special appointed day") those sections shall have effect as if in them—
 - (a) every reference to exportation (in whatever form the reference occurs) were omitted;
 - (b) any reference to the sale or supply of a medicinal product did not include sale or supply which involves, or is for the purposes of, exporting the product; and
 - (c) any reference to offering a medicinal product for sale did not include an offer for sale where the prospective sale would involve, or would be for the purposes of, exporting the product.
- (2) The Ministers shall not make an order under the preceding subsection unless it appears to them to be necessary or expedient to do so for the purpose of giving effect to an agreement to which the United Kingdom or Her Majesty's Government in the United Kingdom is a party or will be a party on the day appointed by the order.
- (3) The following provisions of this section shall have effect where an order is made under subsection (1) of this section; and for the purposes of those provisions the relevant transitional conditions shall be taken to be fulfilled by a person in relation to medicinal products of any description if, in the course of a business carried on by him,—

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- (a) substantial quantities of medicinal products of that description (that is to say, quantities exceeding those required for distribution as samples) were exported or procured to be exported during the period of twenty-four months ending immediately before the special appointed day, and
- (b) during the whole of that period further substantial quantities of medicinal products of that description were available, or could within a reasonable time have been made available, to be so exported or procured to be exported if required.

(4) Unless the order expressly excludes the operation of this subsection,—

- (a) subject to any order made by virtue of paragraph (b) of this subsection, section 7(2) of this Act shall not have effect in relation to a person in respect of his exporting on or after the special appointed day, or procuring the exportation on or after that day of, medicinal products of any description in relation to which he fulfils the relevant transitional conditions;
- (b) section 17 of this Act shall have effect in relation to paragraph (a) of this subsection as it has effect in relation to the subsections of section 16 of this Act mentioned in that section.
- (5) Where a product licence which is in force on the special appointed day authorises the holder of the licence to sell medicinal products of any description, or to procure the sale, or procure the manufacture or assembly for sale, of medicinal products of any description, that licence shall have effect on and after that day as if—
 - (a) it also authorised him to export medicinal products of that description, or (as the case may be) to procure the exportation, or procure the manufacture or assembly for exportation, of medicinal products of that description, and
 - (b) it authorised him to do so subject to the like provisions as (apart from subsections (3) to (7) of section 47 of this Act) are specified in the licence in relation to selling or (as the case may be) procuring the sale, or procuring the manufacture or assembly for sale, of such products:

Provided that, if the operation of subsection (4) of this section is not excluded by the order, a product licence shall not have effect as mentioned in this subsection in relation to medicinal products of any description so long as paragraph (a) of that subsection has effect in relation to the holder of the licence in respect of his exporting, or procuring the exportation of, medicinal products of that description.

- (6) Where on an application for a product licence made before such date as may be appointed by the order for the purposes of this subsection, which states that it is an application made by virtue of this subsection, it is proved to the reasonable satisfaction of the licensing authority that the applicant fulfilled or will fulfil the relevant transitional conditions in relation to one or more descriptions of medicinal products, then (subject to the next following subsection) he shall be entitled to the grant of a product licence granted so as—
 - (a) to be limited to exportation, or procuring exportation, of medicinal products, and
 - (b) not to extend to medicinal products of any description other than those in respect of which it is so proved that the applicant fulfilled or will fulfil those conditions, and
 - (c) not to extend to medicinal products of any description in respect of which, at the time when the licence is granted, a product licence is already held by the applicant.

Changes to legislation: Medicines Act 1968, Part II is up to date with all changes known to be in force on or before 27 June 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations. (See end of Document for details)

- (7) If a person would, on making an application under subsection (6) of this section, be entitled to the grant of a product licence under that subsection in respect of medicinal products of a particular description, and he would at the same time, on making an application as mentioned in section 25(1) of this Act, be entitled to the grant of a licence of right in respect of medicinal products of the same description, he may apply to the licensing authority for a single product licence for both purposes, and he shall be entitled to the grant of a product licence having the same effect as the two licences, if granted separately, would together have had.
- (8) Subsection (6) of section 26 of this Act shall have effect for the purposes of subsections(6) and (7) of this section as it has effect for the purposes of that section.
- (9) An order made under subsection (1) of this section may contain such provisions relating to proceedings on an application made under subsection (6) or subsection (7) of this section (whether by way of applying with modifications any of the provisions of section 27 of this Act or otherwise) as the Ministers may consider appropriate.
- (10) No order shall be made under this section unless a draft of the order has been laid before Parliament and approved by a resolution of each House of Parliament.

Textual Amendments

- F158 Words in s. 48(1) substituted (14.4.1993) by S.I. 1993/834, reg. 5
- **F159** Words in s. 48(1) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 20 (with regs. 2(4), 3)

Modifications etc. (not altering text)

C122 Pt. II(ss. 6-50) extended with modifications by S.I. 1985/1403, art. 3(1)

49 Special provisions in respect of exporting certain products.

- (1) Nothing in subsection (1) of section 48 of this Act shall affect the operation of any of the provisions of sections 7 to 47 of this Act in relation to any medicinal product falling within a class specified in an order made under this section by [^{F160}the Ministers].
- (2) No class of medicinal products shall be specified in an order made by [^{F161}the Ministers] under this section unless it appears to [^{F162}them] to be requisite to do so for securing that any exemption conferred by section 48(1) of this Act does not apply to medicinal products consisting wholly or partly of substances the purity or potency of which cannot, in their opinion, be adequately tested by chemical means.
- (3) Subsections (3) to (7) of section 48 of this Act shall not have effect in relation to medicinal products of any description falling within a class specified in an order under this section which is in force immediately before the day appointed for the purposes of subsection (1) of that section.
- (4) Subject to the next following subsection, section 7(2) of this Act shall not have effect in relation to a person in respect of his exporting, or procuring the exportation of, medicinal products of any description falling within a class specified in an order under this section which is in force immediately before the first appointed day if, in the course of a business carried on by that person,—
 - (a) substantial quantities of medicinal products of that description (that is to say, quantities exceeding those required for distribution as samples) were exported

or procured to be exported during the period of twenty-four months ending with the first appointed day, and

- (b) during the whole of that period further substantial quantities of medicinal products of that description were available, or could within a reasonable time have been made available, to be so exported or procured to be exported if required.
- (5) Sections 17 and 25 of this Act shall have effect in relation to subsection (4) of this section as they have effect in relation to subsections (2) to (5) of section 16 of this Act.
- (6) Where a person is entitled to the grant of a licence of right by reason that subsection (4) of this section has effect in relation to him, he shall be entitled to the grant of a product licence; but, subject to the next following subsection, the licence shall be granted so as not to extend to medicinal products of any description other than those in respect of which the conditions specified in that subsection are proved to the reasonable satisfaction of the licensing authority to have been fulfilled, and shall be limited to exporting, or procuring the exportation of, medicinal products.
- (7) Subsection (5) of section 26 of this Act (with the omission of paragraph (b) of that subsection) and subsection (6) of that section shall have effect in relation to the grant of a licence of right in accordance with subsection (6) of this section as those subsections have effect in relation to the grant of such a licence in accordance with subsection (1) of that section.
- (8) In relation to any application for a licence of right which is made by virtue of section 25 of this Act as applied by subsection (5) of this section, the provisions of section 27 of this Act shall have effect subject to such modifications as may be specified by order made by the Ministers for the purposes of this subsection.

Textual	Amendments
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- **F160** Words in s. 49(1) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 21(a) (with regs. 2(4), 3)
- **F161** Words in s. 49(2) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 21(b)(i) (with regs. 2(4), 3)
- **F162** Word in s. 49(2) substituted (1.10.2006) by The Veterinary Medicines Regulations 2006 (S.I. 2006/2407), reg. 1, Sch. 8 para. 21(b)(ii) (with regs. 2(4), 3)

Modifications etc. (not altering text)

C123 Pt. II(ss. 6-50) extended with modifications by S.I. 1985/1403, art. 3(1)

F163 Special provisions in respect of exporting certain products to member States49A

Textual Amendments

F163 S. 49A repealed (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 6 (with Sch. 6)

[^{F164}49B.Special provisions in respect of exporting certain products to EEA States

Nothing in section 48 of this Act affects the operation of section 8(3A) of this Act in relation to the exportation of a product, or the sale or supply of a product which involves, or is for the purposes of, the exportation of the product if —

- (a) it is a product to which the 2001 Directive applies; and
- (b) the exportation is, or is to be, to an EEA State.]

Textual Amendments

F164 S. 49B inserted (30.10.2005) by The Medicines for Human Use (Manufacturing, Wholesale Dealing and Miscellaneous Amendments) Regulations 2005 (S.I. 2005/2789), reg. 1(1), Sch. 5 para. 7 (with Sch. 6)

50 Certificates for exporters of medicinal products.

On the application of any person who proposes to export medicinal products of any description, the licensing authority may issue to him a certificate containing any such statement relating to medicinal products of that description as the licensing authority may consider appropriate having regard—

- (a) to any requirements (whether having the force of law or not) which have effect in the country to which the products are to be exported, and
- (b) to the provisions of this Act and to any licence granted or other thing done by virtue of this Act.

[^{F165}, and

(c) to the provisions of the Clinical Trials Regulations and to any authorisation granted or other thing done by virtue of those regulations.]

Textual Amendments

F165 S. 50(c) and word inserted (1.5.2004) by Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 para. 16

Modifications etc. (not altering text)

C124 Pt. II(ss. 6–50) extended with modifications by S.I. 1985/1403, art. 3(1)

C125 S. 50 applied (with modifications)(3.4.1992) by S.I. 1992/605, reg. 2(1)(2),Sch.

Status:

Point in time view as at 28/10/2011.

Changes to legislation:

Medicines Act 1968, Part II is up to date with all changes known to be in force on or before 27 June 2024. There are changes that may be brought into force at a future date. Changes that have been made appear in the content and are referenced with annotations.