## STATUTORY INSTRUMENTS

# 1997 No. 1830

## **MEDICINES**

## The Prescription Only Medicines (Human Use) Order 1997

Made	25th July 1997
Laid before Parliament	28th July 1997
Coming into force	18th August 1997

The Secretary of State concerned with health in England, the Secretaries of State concerned with health and with agriculture in Wales and in Scotland respectively, the Minister of Agriculture, Fisheries and Food, the Department of Health and Social Services for Northern Ireland and the Department of Agriculture for Northern Ireland, acting jointly, in exercise of powers conferred on them by sections 58(1), (4) and (5), 59(1) and 129(4) of the Medicines Act 1968(1) or, as the case may be, those conferred by the said provisions and now vested in them(2), and of all other powers enabling them in that behalf, after consulting such organisations as appear to them to be representative of interests likely to be substantially affected by this Order, pursuant to section 129(6) of that Act, and after consulting and taking into account the advice of the Medicines Commission pursuant to sections 58(6) and 129(7) of that Act, hereby make the following Order:

#### Citation, commencement and interpretation

**1.**—(1) This Order may be cited as the Prescription Only Medicines (Human Use) Order 1997 and shall come into force on 18th August 1997.

(2) In this Order, unless the context otherwise requires-

"the Act" means the Medicines Act 1968;

[<sup>F1</sup>"additional supply optometrist" means a person who is registered as an optometrist, and against whose name particulars of the additional supply speciality have been entered in the relevant register;]

"aerosol" means a product which is dispersed from its container by a propellent gas or liquid;

<sup>(1) 1968</sup> c. 67. Section 58 has been amended by the Prescription by Nurses Etc. Act 1992 (c. 28), section 1. The expression "the appropriate Ministers" is defined in section 1(2) of the Medicines Act 1968.

<sup>(2)</sup> In the case of the Secretaries of State concerned with health in England and in Wales by virtue of article 2(2) of, and Schedule 1 to, the Transfer of Functions (Wales) Order 1969 (S.I. 1969/388); in the case of the Secretary of State concerned with Agriculture in Wales by virtue of article 2(3) of, and Schedule 1 to, the Transfer of Functions (Wales) (No. 1) Order 1978 (S.I. 1978/272) and in the case of the Northern Ireland Departments by virtue of section 40 of, and Schedule 5 to, the Northern Ireland Constitution Act 1973 (c. 36) and section 1(3) of, and paragraph 2(1)(b) of Schedule 1 to, the Northern Ireland Act 1974 (c. 28).

[<sup>F2</sup>"clinical management plan" means a written plan (which may be amended from time to time) relating to the treatment of an individual patient agreed by—

- (a) the patient to whom the plan relates,
- (b) the doctor or dentist who is a party to the plan, and
- (c) any supplementary prescriber who is to prescribe, give directions for administration or administer under the plan;]

[<sup>F3</sup>"clinical trial" has the meaning given by regulation 2(1) of the Medicines for Human Use (Clinical Trials) Regulations 2003;]

F4

[<sup>F5</sup>"Common Services Agency" means the Common Services Agency for the Scottish Health Service established under section 10 of the National Health Service (Scotland) Act 1978;]

[<sup>F5</sup>··Community marketing authorization" means a marketing authorization granted by the European Community under Council Regulation (EEC) No. 2309/93[<sup>F6</sup>or 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];]

[<sup>F7</sup>"community practitioner nurse prescriber" means a person—

- (a) who is a registered nurse or a registered midwife, and
- (b) against whose name is recorded in the professional register an annotation signifying that he is qualified to order drugs, medicines and appliances from the Nurse Prescribers' Formulary for Community Practitioners in the current edition of the British National Formulary;]

"controlled drug" has the meaning assigned to it by section 2 of the Misuse of Drugs Act 1971(3);

"cyanogenetic substances" means preparations which-

- (a) are presented for sale or supply under the name of, or as containing, amygdalin, laetrile or vitamin B17, or
- (b) contain more than 0.1 per cent by weight of any substance having the formula either  $\alpha$ -Cyanobenzyl-6-O- $\beta$ -d-glucopyranosyl- $\beta$ -d-glucopyranoside or  $\alpha$ -Cyanobenzyl- $\beta$ -d-glucopy ranosiduronic acid;

[<sup>F8</sup>"dental hygienist" means a person whose name is registered under the title of dental hygienist in the dental care professionals register;

"dental therapist" means a person whose name is registered under the title of dental therapist in the dental care professionals register;

"dental care professionals register" means the dental care professionals register established under section 36B of the Dentists Act 1984;]

F9...

"dosage unit" means-

- (a) where a medicinal product is in the form of a tablet or capsule or is an article in some other similar pharmaceutical form, that tablet, capsule or other article, or
- (b) where a medicinal product is not in any such form, the unit of measurement which is used as the unit by reference to which the dose of the medicinal product is measured;

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"external use" means application to the skin, hair, teeth, mucosa of the mouth, throat, nose, ear, eye, vagina or anal canal when a local action only is intended and extensive systemic absorption is unlikely to occur; and references to medicinal products for external use shall be read accordingly except that such references shall not include throat sprays, throat pastilles, throat lozenges, throat tablets, nasal drops, nasal sprays, nasal inhalations or teething preparations;

F12...

[<sup>F5</sup>"Health Authority"—

(a) <sup>F13</sup>...

- (b) in relation to Scotland, means a Health Board constituted under section 2 of the National Health Service (Scotland) Act 1978; and
- (c) in relation to Northern Ireland, means a Health and Social Services Board established under Article 16 of the Health and Personal Social Services (Northern Ireland) Order 1972;]

[<sup>F14</sup>"health care" means services for or in connection with the prevention, diagnosis or treatment of disease;]

"health prescription" means a prescription issued by a doctor, dentist [ $^{F15}$ , supplementary prescriber], [ $^{F16}$ a community practitioner nurse prescriber, a nurse independent prescriber or a pharmacist independent prescriber] under or by virtue of–

- (a) in England and Wales, the National Health Service Act 1977(4),
- (b) in Scotland, the National Health Service (Scotland) Act 1978(5), and
- (c) in Northern Ireland, the Health and Personal Social Services (Northern Ireland) Order 1972(6);

[<sup>F17</sup>"health record" has the meaning given by section 68(2) of the Data Protection Act 1998(6);]

[<sup>F5</sup> chomoeopathic certificate of registration" means a certificate of registration for the purposes of the Medicines (Homoeopathic Medicinal Products For Human Use) Regulations 1994;]

[<sup>F18</sup>"independent clinic"—

- (a) [<sup>F19</sup>in relation to Wales], has the meaning given by section 2(4) of the Care Standards Act 2000(**8**), and
- (b) in relation to Scotland, has the meaning given by [<sup>F20</sup>section 10F(2) of the National Health Service (Scotland) Act 1978][<sup>F21</sup>, and];
- (c) [<sup>F22</sup>in relation to Northern Ireland, has the meaning given by article 2(2) of the Health and Personal Social Services (Quality, Improvement and Regulation) (Northern Ireland) Order 2003;]]

[<sup>F18</sup>"independent hospital"—

(za) [<sup>F23</sup>in relation to England, means a hospital as defined by section 275 of the National Health Service Act 2006 that is not a health service hospital as defined by that section,

<sup>(</sup>**4**) 1977 c. 49.

<sup>(5) 1978</sup> c. 29.

<sup>(6)</sup> S.I. 1972/1265 (N.I. 14).

<sup>(6)</sup> S.I. 1972/1265 (N.I. 14).

<sup>(8)</sup> S.I. 1985/2066.

- (a) in relation to Wales, shall be construed in accordance with section 2(2), (3) and (6) of the Care Standards Act 2000,]
- (b) in relation to Scotland, means—
  - (i) an independent hospital, or
  - (ii) a private psychiatric hospital,

as defined by  $[^{F20}$ section 10F(2) of the National Health Service (Scotland) Act 1978] $[^{F24}$ , and]

(c) [<sup>F25</sup>in relation to Northern Ireland, has the meaning given by article 2(2) of the Health and Personal Social Services (Quality, Improvement and Regulation) (Northern Ireland) Order 2003;]]

[<sup>F18</sup>"independent medical agency"—

- (a) [<sup>F26</sup>in relation to England and Wales, means an undertaking (not being an independent hospital, or in Wales an independent clinic) which consists of or includes the provision of services by medical practitioners and the term "undertaking" in this definition is to be interpreted in accordance with paragraph (2A),]
- (b) in relation to Scotland, has the meaning given by [<sup>F20</sup>section 10F(2) of the National Health Service (Scotland) Act 1978][<sup>F27</sup>, and]
- (c) [<sup>F28</sup>in relation to Northern Ireland, has the meaning given by article 2(2) of the Health and Personal Social Services (Quality, Improvement and Regulation) (Northern Ireland) Order 2003;]]

"inhaler" does not include an aerosol;

[<sup>F29</sup>"IRME practitioner" means, in relation to a medical exposure, a practitioner for the purposes of the Ionising Radiation (Medical Exposure) Regulations 2000(4);]

[<sup>F30</sup>"Local Health Board" has the same meaning as in the National Health Service Act 1977;]

[<sup>F5</sup>"marketing authorization" includes a reference both to a United Kingdom marketing authorization and to a Community marketing authorization;]

"master" has the same meaning as in section 313(1) of the Merchant Shipping Act 1995(7);

"maximum daily dose" or "MDD" means the maximum quantity of a substance contained in the amount of a medicinal product which it is recommended should be taken or administered in a period of 24 hours;

"maximum dose" or "MD" means the maximum quantity of a substance contained in the amount of a medicinal product which it is recommended should be taken or administered at any one time;

"maximum strength" means-

- (a) the maximum quantity of a substance by weight or volume contained in a dosage unit of a medicinal product;
- (b) the maximum percentage of a substance contained in a medicinal product calculated in any of the following ways-
  - (i) weight in weight,
  - (ii) weight in volume,
  - (iii) volume in weight, or

<sup>(</sup>**4**) 1977 c. 49.

<sup>(7) 1995</sup> c. 21.

(iv) volume in volume,

and if the maximum percentage calculated in those ways differs, the higher or highest such percentage;

[<sup>F31</sup>"medical exposure" has the same meaning as in the Ionising Radiation (Medical Exposure) Regulations 2000;]

"medicinal product" includes any article or substance in respect of which section 58 of the Act has effect by virtue of an order made under section 104 of the Act<sup>F32</sup>...

"the Misuse of Drugs Regulations" means, in relation to England, Wales and Scotland, the Misuse of Drugs Regulations 1985(8) and in relation to Northern Ireland, the Misuse of Drugs (Northern Ireland) Regulations 1986(9);

[<sup>F5</sup>"NHS trust"—

- (a) in relation to England and Wales, has the same meaning as in the National Health Service and Community Care Act 1990;
- (b) in relation to Scotland, has the same meaning as in the National Health Service (Scotland) Act 1978; and
- (c) in relation to Northern Ireland, means a Health and Social Services trust established under article 10 of the Health and Social Services (Northern Ireland) Order 1991;]

[<sup>F33</sup>"nurse independent prescriber" means a person—

- (a) who is a registered nurse or a registered midwife, and
- (b) against whose name is recorded in the professional register an annotation signifying that he is qualified to order drugs, medicines and appliances as a nurse independent prescriber or a nurse independent/supplementary prescriber;]

[<sup>F34</sup>"NHS foundation trust" has the same meaning as in section 1(1) of the Health and Social Care (Community Health and Standards) Act 2003;]

[<sup>F35</sup>"nursing home" has the meaning given by article 16 [<sup>F36</sup>article 11 of the Health and Personal Social Services (Quality, Improvement and Regulation) (Northern Ireland) Order 2003];]

"occupational health scheme" means a scheme in which a person, in the course of a business carried on by him, provides facilities for his employees for the treatment or prevention of disease;

"offshore installation" means an offshore installation within the meaning of the Mineral Workings (Offshore Installations) Act 1971(10) which is within–

- (a) tidal waters and parts of the sea in or adjacent to the United Kingdom up to the seaward limits of territorial waters;
- (b) waters in any area designated under section 1(7) of the Continental Shelf Act 1964(11);

[<sup>F37</sup>"operator"—

- (a) in relation to an aircraft, means the person for the time being having management of the aircraft, and
- (b) for the purposes of article 7B, has the same meaning as in the Ionising Radiation (Medical Exposure) Regulations 2000;]

[<sup>F38</sup>"optometrist independent prescriber" means a person—

<sup>(</sup>**8**) S.I. 1985/2066.

<sup>(9)</sup> SR 1986 No. 52.

<sup>(10) 1971</sup> c. 61; section 1 was substituted by section 24 of the Oil and Gas (Enterprise) Act 1982 (c. 23).

<sup>(</sup>**11**) 1964 c. 29.

- (a) who is a registered optometrist, and
- (b) against whose name is recorded in the relevant register an annotation signifying that he is qualified to order drugs, medicines and appliances as an optometrist independent prescriber;]

"parenteral administration" means administration by breach of the skin or mucous membrane;

[<sup>F5</sup>"Patient Group Direction" means—

- (a) in connection with the [<sup>F39</sup>sale or] supply of a prescription only medicine as referred to in article 12A(2), [<sup>F40</sup>12B, 12C, 12D or 12E], a written direction relating to the [<sup>F41</sup>sale or supply, and administration,] of a description or class of prescription only medicine; or
- (b) in connection with the administration of a prescription only medicine as referred to in article 12A(2), 12B or 12C, a written direction relating to the administration of a description or class of prescription only medicine,

and which, in the case of either (a) or (b)-

- (i) is signed by a doctor or dentist, and by a pharmacist; and
- (ii) relates to [<sup>F42</sup>sale or] supply and administration, or to administration, to persons generally (subject to any exclusions which may be set out in the Direction);]

[<sup>F43</sup>"pharmacist independent prescriber" means a person—

- (a) who is a pharmacist, and
- (b) against whose name is recorded in the relevant register an annotation signifying that he is qualified to order drugs, medicines and appliances as a pharmacist independent prescriber;]

"prescription only medicine" means a medicinal product of a description or falling within a class specified in article 3 of this Order;

[<sup>F5</sup>"Primary Care Trust" has the same meaning as in the National Health Service Act 1977;]

[<sup>F44</sup>"prison service" means—

- (a) in relation to England and Wales, a Minister of the Crown exercising functions in relation to prisons (within the meaning of the Prison Act 1952),
- (b) in relation to Scotland, the Scottish Ministers exercising functions in relation to prisons (within the meaning of the Prisons (Scotland) Act 1989), and
- (c) in relation to Northern Ireland, the Northern Ireland Department exercising functions in relation to prisons (within the meaning of the Prison Act (Northern Ireland) 1953);]

[<sup>F45</sup>"professional register" means the register maintained by the Nursing and Midwifery Council [<sup>F46</sup>under article 5 of] the Nursing and Midwifery Order 2001;]

"prolonged release" in relation to a medicinal product means a formulation of that product which-

- (a) is used to reduce the rate at which the active ingredient in that product is released after administration, and
- (b) is sold or supplied as a prolonged, controlled or sustained release medicinal product;

[<sup>F47</sup>"radioactive medicinal product" means a medicinal product which is, which contains or which generates a radioactive substance and which is, contains or generates that substance in order, when administered, to utilize the radiation emitted therefrom;]

 $[^{F48}$  (registered chiropodist" means a person who is registered in Part 2 of the register maintained by the Health Professions Council under article 5 of the  $[^{F49}$ Health and Social Work Professions Order 2001];]

[<sup>F50</sup>"registered dietitian" means a person who is registered in Part 4 of the register maintained by the Health Professions Council under article 5 of the [<sup>F49</sup>Health and Social Work Professions Order 2001];]

[<sup>F51</sup>"registered midwife" means a person registered in the Midwives' Part of the professional register;]

[<sup>F52</sup>"registered nurse" means a person registered in the Nurses' Part [<sup>F53</sup> or Specialist Community Public Health Nurses' Part] of the professional register;]

[<sup>F50</sup>"registered occupational therapist" means a person who is registered in Part 6 of the register maintained by the Health Professions Council under article 5 of the [<sup>F49</sup>Health and Social Work Professions Order 2001];]

[ $^{F54}$  "registered optometrist" means a person whose name is registered in the register of optometrists maintained under section 7(a) of the Opticians Act 1989, or in the register of visiting optometrists from relevant European States maintained under section 8B(1)(a) of that Act;]

[<sup>F48</sup>"registered orthoptist" means a person who is registered in Part 7 of the register maintained by the Health Professions Council under article 5 of the [<sup>F49</sup>Health and Social Work Professions Order 2001];]

[<sup>F50</sup>"registered orthotist and prosthetist" means a person who is registered in Part 10 of the register maintained by the Health Professions Council under article 5 of the [<sup>F49</sup>Health and Social Work Professions Order 2001];]

[<sup>F48</sup>"registered paramedic" means a person who is registered in Part 8 of the register maintained by the Health Professions Council under article 5 of the [<sup>F49</sup>Health and Social Work Professions Order 2001];]

[<sup>F48</sup>"registered physiotherapist" means a person who is registered in Part 9 of the register maintained by the Health Professions Council under article 5 of the [<sup>F49</sup>Health and Social Work Professions Order 2001];]

[F55" registered provider" means-

- (a) in relation to an independent hospital, an independent clinic [<sup>F56</sup>in Wales, Scotland or Northern Ireland] or an independent medical agency—
  - (i) [<sup>F57</sup>in relation—
    - (aa) to England, the person who is registered under Chapter 2 of Part 1 of the Health and Social Care Act 2008 in respect of regulated activities (within the meaning of that Part) carried on in that hospital or agency, and
    - (bb) to Wales, the person who is registered under Part 2 of the Care Standards Act 2000 as the person carrying on the establishment or agency,]
  - (ii) in relation to Scotland, the person who is registered under [<sup>F58</sup>section 10P of the National Health Service (Scotland) Act 1978] as the person providing the establishment or agency, and
  - (iiii) [<sup>F59</sup>in relation to Northern Ireland, the person registered under Part III of the Health and Personal Social Services (Quality, Improvement and Regulation) (Northern Ireland) Order 2003 as the person carrying on the establishment or agency, and]

(b) [<sup>F60</sup>in relation to a nursing home, the person registered under Part III of the Health and Personal Social Services (Quality, Improvement and Regulation) (Northern Ireland) Order 2003 as the person carrying on the establishment;]]

[<sup>F48</sup>"registered radiographer" means a person who is registered in Part 11 of the register maintained by the Health Professions Council under article 5 of the [<sup>F49</sup>Health and Social Work Professions Order 2001];]

[<sup>F50</sup>"registered speech and language therapist" means a person who is registered in Part 12 of the register maintained by the Health Professions Council under article 5 of the [<sup>F49</sup>Health and Social Work Professions Order 2001];]

[<sup>F55</sup>"relevant manager" means—

- (a) in relation to an independent hospital, an independent clinic [<sup>F61</sup>in Wales, Scotland or Northern Ireland] or an independent medical agency—
  - (i) [<sup>F62</sup>in relation to England, a person who is registered under Chapter 2 of Part 1 of the Health and Social Care Act 2008 as a manager in respect of regulated activities (within the meaning of that Part) carried on in that hospital or agency, and
  - (ia) in relation to Wales—]
    - (aa) a person who is registered under Part II of the Care Standards Act 2000 as the manager of the establishment or agency, but who is not the registered provider for that establishment or agency, or
    - (bb) if there is no such person, but the registered provider has appointed a person to manage the establishment or agency, that appointed person,
  - (ii) in relation to Scotland, a person, other than the registered provider, who was identified as an individual who is to manage the establishment or agency on the application for registration of that establishment or agency under [<sup>F58</sup>section 10P of the National Health Service (Scotland) Act 1978], and
  - (iii) [<sup>F63</sup>in relation to Northern Ireland—
    - (aa) a person who is registered under Part III of the Health and Personal Social Services (Quality, Improvement and Regulation) (Northern Ireland) Order 2003 as the manager of the establishment or agency, but who is not the registered provider for that establishment or agency, or
    - (bb) if there is no such person, but the registered provider has appointed a person to manage the establishment or agency, that appointed person, and]
- (b) [<sup>F64</sup>in relation to a nursing home—
  - a person who is registered under Part III of the Health and Personal Social Services (Quality, Improvement and Regulation) (Northern Ireland) Order 2003 as the manager of the establishment or agency, but who is not the registered provider for that establishment or agency, or
  - (ii) if there is no such person, but the registered provider has appointed a person to manage the establishment or agency, that appointed person;]]

[<sup>F55</sup>"relevant register" means—

(a) in relation to a [<sup>F65</sup>registered] nurse [<sup>F66</sup>or registered midwife], the professional register, F67...

- (b) in relation to a pharmacist, [<sup>F68</sup>Part 1 of the register maintained under article 19 of the Pharmacy Order 2010] or the register maintained in pursuance of Articles 6 and 9 of the Pharmacy (Northern Ireland) Order 1976; <sup>F69</sup>...
- (c) [<sup>F70</sup>in relation to a person whose name is registered in the part of the register maintained by the Health Professions Council in pursuance of article 5 of the [<sup>F71</sup>Health and Social Work Professions Order 2001] relating to—
  - (i) chiropodists and podiatrists;
  - (ii) physiotherapists; or
  - (iii) radiographers diagnostic or therapeutic

that register [<sup>F72</sup>; and]

(d) [<sup>F73</sup>in relation to a registered optometrist, the register of optometrists maintained under section 7(a) of the Opticians Act 1989;]]

"repeatable prescription" means a prescription which contains a direction that it may be dispensed more than once;

"sell" means sell by retail as defined in section 131 of the Act and "sale" has a corresponding meaning;

"soap" means any compound of a fatty acid with an alkali or amine;

[<sup>F5</sup>"Special Health Authority"—

- (a) in relation to England and Wales, has the same meaning as in the National Health Service Act 1977;
- (b) in relation to Scotland, means a Special Health Board constituted under section 2 of the National Health Service (Scotland) Act 1978; and
- (c) in relation to Northern Ireland, means a Special Health and Social Services Agency established under the Health and Personal Social Services (Special Agencies) (Northern Ireland) Order 1990;]

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[<sup>F75</sup>"Strategic Health Authority" means a Strategic Health Authority established under section 8 of the National Health Service Act 1977;]

[<sup>F76</sup>"supplementary prescriber" means—

- (a) a [<sup>F77</sup>registered] nurse, <sup>F78</sup>...
- (b) a pharmacist, <sup>F79</sup>[<sup>F80</sup>...
- (c) a registered midwife,]<sup>F81</sup>...
- (d) [<sup>F82</sup>a person whose name is registered in the part of the register maintained by the Health Professions Council in pursuance of article 5 of the [<sup>F83</sup>Health and Social Work Professions Order 2001] relating to—
  - (i) chiropodists and podiatrists;
  - (ii) physiotherapists; or
  - (iii) radiographers diagnostic or therapeutic [<sup>F84</sup>or]]
- (e) [<sup>F85</sup>a registered optometrist, ]

against whose name is recorded in the relevant register an annotation [<sup>F86</sup>or entry] signifying that he is qualified to order drugs, medicines and appliances as a supplementary prescriber [<sup>F87</sup>or, in the case of a nurse or midwife, as a nurse independent/supplementary prescriber]; ]

"supply" means supply in circumstances corresponding to retail sale as defined in section 131 of the Act;

"unit preparation" means a preparation, including a mother tincture, prepared by a process of solution, extraction or trituration with a view to being diluted tenfold or one hundredfold, either once or repeatedly, in an inert diluent, and then used either in this diluted form or, where applicable, by impregnating tablets, granules, powders or other inert substances.

[<sup>F56</sup> United Kingdom marketing authorization" means a marketing authorization granted by the licensing authority under the Medicines For Human Use (Marketing Authorisations Etc.) Regulations 1994 (including a product licence having effect as such an authorization by virtue of Schedule 6 to those Regulations [<sup>F88</sup> or an authorization granted by the licensing authority in accordance with Article 126a of Directive 2001/83/EC]).]

[

<sup>F89</sup>(2A) In paragraph (2), for the purposes of the definition of "independent medical agency", "undertaking" includes any business or profession and—

- (a) in relation to a public or local authority, includes the exercise of any functions of that authority; and
- (b) in relation to any other body of persons, whether corporate or unincorporate, includes any of the activities of that body.]

(3) For the purposes of this Order, the equivalence of a substance to a reference material shall be determined by calculating the amount of that reference material which is contained in that substance either by weight or, where the amount of the reference material is specified in terms of international units of activity, those units.

(4) In this Order, unless the context otherwise requires, a reference-

- (a) to a numbered section is to the section of the Act which bears that number,
- (b) to a numbered article or Schedule is to the article of, or Schedule to, this Order which bears that number,
- (c) in an article or in a Part of a Schedule to a numbered paragraph is to the paragraph of that article or Part of that Schedule which bears that number, and
- (d) in a paragraph to a lettered sub-paragraph is to the sub-paragraph of that paragraph which bears that letter.
- (5) In [<sup>F90</sup>Schedules 1, 2, 3A and 5]-
  - (a) entries specified in columns 2 to 5 relate to the substances listed in column 1 against which they appear and where, in relation to a particular substance listed in column 1, an entry in columns 2 to 5 bears a number or letter it relates only to such entries in the other of those columns as bear the same number or letter;
  - (b) the following abbreviations are used:

"g" for gram,

"iu" for international unit of activity,

"mcg" for microgram,

"mg" for milligram,

"ml" for millilitre.

(6) In Schedule 3, the abbreviation "NPF" means the Nurse Prescribers' Formulary Appendix in the British National Formulary.

<sup>F91</sup>(7) In articles 12 to [ $^{F92}12E$ ], a reference to a prescription only medicine being sold or supplied for the purpose of being administered in accordance with the written directions of a doctor or dentist relating to a particular person, or in accordance with a Patient Group Direction, includes a reference to it being sold or supplied in accordance with such directions or such a Direction.

(8) In articles 12A and 12C, a reference to an arrangement for the [<sup>F93</sup>sale,] supply or administration of prescription only medicines includes a reference to an arrangement which covers such [<sup>F94</sup>sale,] supply or administration and other matters.

(9) In Schedule 7, Part I, a reference to treatment of a clinical situation includes a reference to any form of management of that situation.]]

#### **Textual Amendments**

- F1 Words in art. 1(2) inserted (30.6.2005) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), **3(a)**
- F2 Words in art. 1(2) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **2(2)(a)**
- F3 Words in art. 1(2) substituted (1.5.2004) by The Medicines for Human Use (Clinical Trials) Regulations 2004 (S.I. 2004/1031), reg. 1, Sch. 10 Pt. 2 para. 13(a)
- F4 Words in art. 1 omitted (1.4.2002) by virtue of The Prescription Only Medicines (Human Use) Amendment Order 2002 (S.I. 2002/549), arts. 1(1), **2(2)(a)**
- **F5** Words in art. 1(2) inserted (9.8.2000) by The Prescription Only Medicines (Human Use) Amendment Order 2000 (S.I. 2000/1917), arts. 1(1), **2(a)**
- **F6** Words in art. 1(2) inserted (20.11.2005) by The Medicines (Marketing Authorisations Etc.) Amendment Regulations 2005 (S.I. 2005/2759), reg. 1(b), **Sch. para. 5(a)**
- **F7** Words in art. 1(2) inserted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **2(a)**
- **F8** Words in art. 1(2) inserted (1.6.2010) by The Medicines for Human Use (Miscellaneous Amendments) Order 2010 (S.I. 2010/1136), arts. 1(1), **3(2**)
- **F9** Words in art. 1(2) omitted (1.5.2006) by virtue of The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **2(b)**
- **F10** Words in art. 1(2) omitted (1.5.2006) by virtue of The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **2(c)**
- F11 Words in art. 1(2) omitted (1.5.2006) by virtue of The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), 2(d)
- **F12** Words in art. 1(2) omitted (1.5.2006) by virtue of The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **2(e)**
- F13 Words in art. 1(2) repealed (8.5.2009) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2009 (S.I. 2009/1165), arts. 1, 2
- F14 Words in art. 1(2) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 2(2)(d)
- F15 Words in art. 1(2) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **2(2)(e)**
- **F16** Words in art. 1(2) substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **2(f)**
- F17 Words in art. 1(2) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **2(2)(f)**
- **F18** Words in art. 1(2) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **2(2)**(g)

- **F19** Words in art. 1(2) substituted (1.10.2010) by The Health and Social Care Act 2008 (Miscellaneous Consequential Amendments) Order 2010 (S.I. 2010/1881), arts. 1(1), **9(2)(a)**
- F20 Words in art. 1(2) 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. 24(a)
- F21 Words in art. 1(2) substituted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 2(2)(a)(i)
- F22 Words in art. 1(2) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 2(2)(a)(ii)
- F23 Words in art. 1(2) substituted (1.10.2010) by The Health and Social Care Act 2008 (Miscellaneous Consequential Amendments) Order 2010 (S.I. 2010/1881), arts. 1(1), 9(2)(b)
- F24 Words in art. 1(2) substituted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 2(2)(b)(i)
- F25 Words in art. 1(2) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 2(2)(b)(ii)
- **F26** Words in art. 1(2) substituted (1.10.2010) by The Health and Social Care Act 2008 (Miscellaneous Consequential Amendments) Order 2010 (S.I. 2010/1881), arts. 1(1), **9(2)(c)**
- **F27** Words in art. 1(2) substituted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **2(2)(c)(i)**
- **F28** Words in art. 1(2) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **2(2)(c)(ii)**
- **F29** Words in art. 1(2) inserted (17.11.2006) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2006 (S.I. 2006/2807), arts. 1(1), **2(a)**
- F30 Words in art. 1(2) inserted (19.11.2004) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 2004 (S.I. 2004/2693), arts. 1(1), 2
- **F31** Words in art. 1(2) inserted (17.11.2006) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2006 (S.I. 2006/2807), arts. 1(1), **2(b)**
- **F32** Words in art. 1(2) omitted (17.11.2006) by virtue of The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2006 (S.I. 2006/2807), arts. 1(1), **2(c)**
- **F33** Words in art. 1(2) inserted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **2(g)**
- F34 Words in art. 1(2) inserted (1.4.2004) by The Health and Social Care (Community Health and Standards) Act 2003 (Supplementary and Consequential Provision) (NHS Foundation Trusts) Order 2004 (S.I. 2004/696), arts. 1(1)(b), 3(8), Sch. 8
- **F35** Words in art. 1(2) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **2(2)(h)**
- **F36** Words in art. 1(2) substituted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **2(2)(d)**
- **F37** Words in art. 1(2) substituted (17.11.2006) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2006 (S.I. 2006/2807), arts. 1(1), **2(d)**
- **F38** Words in art. 1(2) inserted (4.6.2008) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2008 (S.I. 2008/1161), arts. 1(1), **2**
- F39 Words in art. 1(2) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 2(2)(e)(i)(aa)
- **F40** Words in art. 1 substituted (10.12.2003) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2003 (S.I. 2003/2915), arts. 1(1), **2**
- **F41** Words in art. 1(2) substituted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **2(2)(e)(i)(bb)**
- F42 Words in art. 1(2) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 2(2)(e)(ii)
- **F43** Words in art. 1(2) inserted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **2(h)**

- F44 Words in art. 1(2) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 2(2)(i)
- F45 Words in art. 1 inserted (1.4.2002) by The Prescription Only Medicines (Human Use) Amendment Order 2002 (S.I. 2002/549), arts. 1(1), **2(2)(e)**
- F46 Words in art. 1(2) substituted (1.8.2004) by The Health Act 1999 (Consequential Amendments) (Nursing and Midwifery) Order 2004 (S.I. 2004/1771), art. 1(1), Sch. para. 39(a)(iv)
- **F47** Words in art. 1(2) inserted (17.11.2006) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2006 (S.I. 2006/2807), arts. 1(1), **2(e)**
- F48 Words in art. 1 inserted (9.7.2003) by The Health Professions Order 2001 (Consequential Amendments) Order 2003 (S.I. 2003/1590), art. 1, Sch. para. 21(2)(a)
- **F49** Words in art. 1(2) substituted (1.8.2012) by The Health and Social Care Act 2012 (Consequential Provision—Social Workers) Order 2012 (S.I. 2012/1479), art. 1(2), Sch. para. 13(2)(a)
- **F50** Words in art. 1(2) inserted (18.5.2004) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2004 (S.I. 2004/1189), arts. 1(1), **2**
- F51 Words in art. 1(2) substituted (1.8.2004) by The Health Act 1999 (Consequential Amendments) (Nursing and Midwifery) Order 2004 (S.I. 2004/1771), art. 1(1), Sch. para. 39(a)(v)
- **F52** Words in art. 1(2) substituted (1.8.2004) by The Health Act 1999 (Consequential Amendments) (Nursing and Midwifery) Order 2004 (S.I. 2004/1771), art. 1(1), Sch. para. 39(a)(vi)
- **F53** Words in art. 1(2) inserted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **2(i)**
- F54 Words in art. 1(2) substituted (3.12.2007) by The European Qualifications (Health and Social Care Professions) Regulations 2007 (S.I. 2007/3101), regs. 1(2), 197
- **F55** Words in art. 1(2) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **2(2)(j)**
- F56 Words in art. 1(2) inserted (1.10.2010) by The Health and Social Care Act 2008 (Miscellaneous Consequential Amendments) Order 2010 (S.I. 2010/1881), arts. 1(1), 9(2)(d)(i)
- **F57** Words in art. 1(2) substituted (1.10.2010) by The Health and Social Care Act 2008 (Miscellaneous Consequential Amendments) Order 2010 (S.I. 2010/1881), arts. 1(1), **9(2)(d)(ii)**
- F58 Words in art. 1(2) 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. 24(b)
- **F59** Words in art. 1(2) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **2(2)(f)(i)**
- **F60** Words in art. 1(2) substituted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **2(2)(f)(ii)**
- **F61** Words in art. 1(2) inserted (1.10.2010) by The Health and Social Care Act 2008 (Miscellaneous Consequential Amendments) Order 2010 (S.I. 2010/1881), arts. 1(1), **9(2)(e)(i)**
- **F62** Words in art. 1(2) substituted (1.10.2010) by The Health and Social Care Act 2008 (Miscellaneous Consequential Amendments) Order 2010 (S.I. 2010/1881), arts. 1(1), **9(2)(e)(ii)**
- F63 Words in art. 1(2) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 2(2)(g)(i)
- **F64** Words in art. 1(2) substituted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **2(2)(g)(ii)**
- **F65** Word in art. 1(2) substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **2(j)**
- **F66** Words in art. 1(2) inserted (1.8.2004) by The Health Act 1999 (Consequential Amendments) (Nursing and Midwifery) Order 2004 (S.I. 2004/1771), art. 1(1), Sch. para. 39(a)(vii)
- **F67** Word in art. 1(2) omitted (7.4.2005) by virtue of The Medicines for Human Use (Prescribing) Order 2005 (S.I. 2005/765), arts. 1(1), **3(a)(i)**
- F68 Words in art. 1(2) substituted (27.9.2010) by The Pharmacy Order 2010 (S.I. 2010/231), art. 1(5), Sch. 4 para. 27 (with Sch. 5); S.I. 2010/1621, art. 2(1), Sch.
- F69 Word in art. 1(b) omitted (30.6.2005) by virtue of The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), 3(b)(i)

- **F70** Words in art. 1(2) inserted (7.4.2005) by The Medicines for Human Use (Prescribing) Order 2005 (S.I. 2005/765), arts. 1(1), **3(a)(iii)**
- **F71** Words in art. 1(2) substituted (1.8.2012) by The Health and Social Care Act 2012 (Consequential Provision—Social Workers) Order 2012 (S.I. 2012/1479), art. 1(2), **Sch. para. 13(2)(b)**
- **F72** Word in art. 1(c) inserted (30.6.2005) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), **3(b)(ii)**
- F73 Words inserted (30.6.2005) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), 3(b)(iii)
- F74 Words in art. 1 omitted (9.7.2003) by virtue of The Health Professions Order 2001 (Consequential Amendments) Order 2003 (S.I. 2003/1590), art. 1, Sch. para. 21(2)(b)
- F75 Words in art. 1(2) inserted (1.10.2002) by The National Health Service Reform and Health Care Professions Act 2002 (Supplementary, Consequential etc. Provisions) Regulations 2002 (S.I. 2002/2469), regs. 1, 7, Sch. 4
- **F76** Words in art. 1(2) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **2(2)(k)**
- **F77** Word in art. 1(2) substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **2(k)(i)**
- F78 Word in art. 1(2) 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. 39(a)(viii)
- F79 Word in art. 1(2) omitted (7.4.2005) by virtue of The Medicines for Human Use (Prescribing) Order 2005 (S.I. 2005/765), arts. 1(1), 3(b)(i)
- **F80** Words in art. 1(2) inserted (1.8.2004) by The Health Act 1999 (Consequential Amendments) (Nursing and Midwifery) Order 2004 (S.I. 2004/1771), art. 1(1), Sch. para. 39(a)(viii)
- F81 Word in art. 1(2)(c) omitted (30.6.2005) by virtue of The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), 3(c)(i)
- **F82** Words in art. 1(2) inserted (7.4.2005) by The Medicines for Human Use (Prescribing) Order 2005 (S.I. 2005/765), arts. 1(1), **3(b)(iii)**
- **F83** Words in art. 1(2) substituted (1.8.2012) by The Health and Social Care Act 2012 (Consequential Provision—Social Workers) Order 2012 (S.I. 2012/1479), art. 1(2), Sch. para. 13(2)(c)
- **F84** Word in art. 1(2)(d) inserted (30.6.2005) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), **3(c)(ii)**
- **F85** Art. 1(2)(e) inserted (30.6.2005) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), **3(c)(iii)**
- **F86** Words in art. 1(2) inserted (7.4.2005) by The Medicines for Human Use (Prescribing) Order 2005 (S.I. 2005/765), arts. 1(1), **3(c)**
- **F87** Words in art. 1(2) inserted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **2(k)(ii)**
- **F88** Words in art. 1(2) inserted (20.11.2005) by The Medicines (Marketing Authorisations Etc.) Amendment Regulations 2005 (S.I. 2005/2759), reg. 1(b), **Sch. para. 5(b)**
- **F89** Art. 1(2A) inserted (1.10.2010) by The Health and Social Care Act 2008 (Miscellaneous Consequential Amendments) Order 2010 (S.I. 2010/1881), arts. 1(1), **9(2)(f)**
- **F90** Words in art. 1(5) substituted (1.4.2002) by The Prescription Only Medicines (Human Use) Amendment Order 2002 (S.I. 2002/549), arts. 1(1), **2(3)**
- **F91** Words in art. 1(7)-(9) inserted (9.8.2000) by The Prescription Only Medicines (Human Use) Amendment Order 2000 (S.I. 2000/1917), arts. 1(1), **2(b)**
- **F92** Word in art. 1(7) substituted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **2(3)**
- **F93** Word in art. 1(8) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **2(3)(a)**
- **F94** Word in art. 1(8) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **2(3)(b)**

#### **Appropriate practitioners**

**2.** For the purposes of section 58 (medicinal products on prescription only), the following shall be appropriate practitioners–

- (a) in relation to the descriptions and classes of medicinal products specified in article
   3, doctors, dentists [<sup>F95</sup>, supplementary prescribers], [<sup>F96</sup>nurse independent prescribers, pharmacist independent prescribers,] veterinary surgeons and veterinary practitioners;
- [<sup>F97</sup>(b) in relation to the descriptions and classes of medicinal products specified in Schedule 3, [<sup>F98</sup>community practitioner nurse prescribers];
- <sup>F99</sup>(c) .....]
- [<sup>F100</sup>(c) in relation to the descriptions and classes of medicinal products specified in article 3, other than medicinal products that are controlled drugs or for parenteral administration or both, optometrist independent prescribers.]
- <sup>F99</sup>(d) .....

#### **Textual Amendments**

- **F95** Words in art. 2(a) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **3**
- **F96** Words in art. 2(a) added (1.4.2008) by The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), **2(a)**
- **F97** Art. 2(b)(c) substituted for art. 2(b) (1.4.2002) by The Prescription Only Medicines (Human Use) Amendment Order 2002 (S.I. 2002/549), arts. 1(1), **3**
- **F98** Words in art. 2(b) substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), **3(a)**
- **F99** Art. 2(c)(d) omitted (1.4.2008) by virtue of The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), **2(b)**
- F100 Art. 2(c) inserted (4.6.2008) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2008 (S.I. 2008/1161), arts. 1(1), **3**

### [<sup>F101</sup>Medicinal products on prescription only

**3.** The following descriptions and classes of medicinal products are specified for the purposes of section 58, namely—

- (a) medicinal products in respect of which a marketing authorization has been granted, which in the marketing authorization are classified as being prescription only medicines;
- (b) medicinal products in respect of which no marketing authorization has been granted consisting of or containing a substance listed in column 1 of Schedule 1;
- (c) medicinal products that are for parenteral administration;
- (d) medicinal products that are controlled drugs unless a marketing authorization has been granted in respect of that medicinal product in which the product is classified as being a pharmacy only or on general sale list medicine;
- (e) cyanogenetic substances, other than preparations for external use;
- (f) medicinal products that on administration emit radiation, or contain or generate any substance which emits radiation, in order that radiation may be used.
- [<sup>F102</sup>(g) medicinal products in respect of which a marketing authorization has been granted consisting of or containing aloxiprin, aspirin or paracetamol in the form of non-

effervescent tablets or capsules which in the marketing authorization are classified as being pharmacy only or general sale list medicines.]

[<sup>F103</sup>(h) medicinal products in respect of which a marketing authorization has been granted consisting of or containing pseudoephedrine salts or ephedrine base or salts in all pharmaceutical forms which in the marketing authorization are classified as being pharmacy only medicines.]]

#### **Textual Amendments**

- **F101** Art. 3 substituted (1.4.2002) by The Prescription Only Medicines (Human Use) Amendment Order 2002 (S.I. 2002/549), arts. 1(1), 4
- F102 Art. 3(g) added (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 4
- **F103** Art. 3(h) inserted (1.4.2008) by The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), **3**

#### Prescribing and administration by nurse independent prescribers

<sup>F104</sup>3A.

#### **Textual Amendments**

F104 Art. 3A omitted (1.4.2008) by virtue of The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), 4

#### [<sup>F105</sup>Prescribing and administration by supplementary prescribers

**3B.**—(1) Subject to paragraph (2), a supplementary prescriber may—

- (a) give a prescription for a medicinal product referred to in article 3; or
- (b) if that medicinal product is for parenteral administration—
  - (i) administer that medicinal product, or
  - (ii) give directions for the administration of that medicinal product,

only where he complies with the conditions as to the cases or circumstances in which he may do so specified in paragraph (3).

[<sup>F106</sup>(2) Paragraph (1) does not apply if the supplementary prescriber is a community practitioner nurse prescriber and the medicinal product prescribed or administered, or in respect of which he gives directions for administration, falls within a description or class of medicinal products specified in Schedule 3.]

(3) The conditions referred to in paragraph (1) are that—

- (a) the supplementary prescriber is acting in accordance with the terms of a clinical management plan which—
  - (i) relates to the patient for whom the product is prescribed or to whom it is, or is to be, administered,
  - (ii) is in effect at the time the prescription or direction is given or, as the case may be, the product is administered, and
  - (iii) includes the particulars specified in Schedule 3B;

 $F^{107}(b)$  ....

(c) the supplementary prescriber has access to the health records of the patient to whom the plan relates which are used by any doctor or dentist who is a party to the plan.

#### **Textual Amendments**

- F105 Arts. 3B, 3C inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 6
- F106 Art. 3B(2) substituted (1.4.2008) by The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), 5
- F107 Art. 3B(3)(b) omitted (7.4.2005) by virtue of The Medicines for Human Use (Prescribing) Order 2005 (S.I. 2005/765), arts. 1(1), 4

# Exemptions from conditions in respect of the cases or circumstances in which a <sup>F108</sup>... supplementary prescriber may administer a medicinal product

**3C.** The conditions specified by virtue of  $^{F109}$ ... article 3B(3) shall not apply in relation to the administration of a medicinal product by  $^{F110}$ ... a supplementary prescriber where—

- (a) that person is acting in accordance with the directions of another person who is an appropriate practitioner, or
- (b) that administration is exempt from the restriction in section 58(2)(b) (restriction on administration) by virtue of any of the subsequent provisions of this Order.]

#### **Textual Amendments**

- F105 Arts. 3B, 3C inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 6
- **F108** Words in art. 3C heading omitted (1.4.2008) by virtue of The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), **6(a)**
- **F109** Words in art. 3C omitted (1.4.2008) by virtue of The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), **6(b)(i)**
- **F110** Words in art. 3C omitted (1.4.2008) by virtue of The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), **6(b)(ii)**

#### Duration of special provisions in relation to new medicinal products

#### **Textual Amendments**

F111 Art. 4 revoked (1.4.2002) by The Prescription Only Medicines (Human Use) Amendment Order 2002 (S.I. 2002/549), arts. 1(1), 11

#### **Exempt medicinal products**

**5.**—(1) A medicinal product shall be exempt from the restrictions imposed by section 58(2)(a) (restrictions on sale or supply) if it, or a substance in it, is listed in column 1 of Schedule 1 and there-

(a) is an entry in column 2, 3, 4 or 5 of that Schedule which contains a condition and that condition is satisfied in accordance with the following provisions of this article; or

(b) there is more than one such condition which applies where that substance is used in that product and each of those conditions is so satisfied.

(2) Where a maximum strength is specified in column 2 in relation to a substance, a medicinal product which consists of or contains that substance satisfies the condition where the maximum strength of that substance in that medicinal product, or where specified in that column the maximum strength of a medicinal product which contains that substance, does not exceed that specified maximum or, where the medicinal product consists of more than one of the substances sodium fluoride, sodium monofluorophosphate or stannous fluoride combined in a dentifrice, where the maximum strength of that combination of substances in a product does not exceed the equivalent of 0.15 per cent of fluorine.

(3) Where a route of administration is specified in column 3 in relation to a substance, a medicinal product which consists of or contains that substance satisfies the condition where it is sold or supplied for administration only by that route.

(4) Where a use is specified in column 3 in relation to a substance, a medicinal product which consists of or contains that substance satisfies the condition in respect of use where it is sold or supplied only for use-

- (a) where a purpose for which it may be used is so specified, for that purpose;
- (b) where the class of persons in whom it may be used is so specified, in persons of that class  $I^{F112}$ ,

provided that, where the entry in column 3 contains a condition to the effect that the product is to be wholly or mainly for use in a specified class of persons, the product satisfies that condition where it is sold or supplied wholly or mainly for use in persons of that class].

(5) Where a pharmaceutical form is specified in column 3 in relation to a substance, a medicinal product which consists of or contains that substance satisfies the condition where it is sold or supplied in that pharmaceutical form.

(6) Where a maximum dose is specified in column 4 in relation to a substance, a medicinal product which consists of or contains that substance satisfies the condition where it is sold or supplied for use at a maximum dose which does not exceed that specified maximum dose.

(7) Subject to paragraph (8), where a maximum daily dose is specified in column 4 in relation to a substance, a medicinal product which consists of or contains that substance satisfies the condition where it is sold or supplied for use at a maximum daily dose which does not exceed that specified maximum daily dose.

(8) A medicinal product which contains more than one of the substances-

Atropine Atropine Methobromide Atropine Methonitrate Atropine Oxide Hydrochloride Atropine Sulphate Hyoscine Hyoscine Butylbromide Hyoscine Hydrobromide Hyoscine Methobromide Hyoscyamine Hyoscyamine Hyoscyamine Sulphate,

satisfies the condition only where it is sold or supplied for use at a maximum daily dose which does not exceed 1 milligram in total of the alkaloids derived from belladonna, hyoscyamus, stramonium or other solanaceous plant which are contained in that medicinal product.

(9) Where a maximum period of use or a maximum frequency of use is specified in column 4 in relation to a substance, a medicinal product which consists of or contains that substance satisfies the condition where it is sold or supplied for use for a maximum period or frequency, as the case may be, which does not exceed the maximum period of use or the maximum frequency of use which is so specified.

(10) Where a maximum quantity is specified in column 5 in relation to a substance, a medicinal product which consists of or contains that substance satisfies the condition where it, or where so specified in that column, the medicinal product which contains that substance is sold or supplied in a quantity which does not exceed that specified maximum quantity.

(11) In paragraphs (2) to (7) and (9) and (10) a reference to a numbered column is a reference to the column bearing that number in Schedule 1.

#### **Textual Amendments**

F112 Words in art. 5(4) added (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 2

## [<sup>F113</sup>Exemption for products consisting of or containing aloxiprin, aspirin or paracetamol

**5A.** A medicinal product falling within article 3(g) shall be exempt from the restrictions imposed by section 58(2) (restrictions on sale, supply and administration) if the quantity of that product sold or supplied to a person at any one time does not exceed 100 tablets or capsules.]

#### **Textual Amendments**

F113 Art. 5A inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 7

# [<sup>F114</sup>Exemption for products consisting of or containing pseudoephedrine salts or ephedrine base or salts

**5B.**—(1) A medicinal product falling within article 3(h) shall be exempt from the restrictions imposed by section 58(2) (restrictions on sale, supply and administration) if the conditions in paragraph (2) are satisfied—

(2) The conditions referred to in this paragraph are that—

- (a) the medicinal product sold or supplied to a person must not be sold or supplied at the same time as another medicinal product that consists of or contains—
  - (i) in the case of pseudoephedrine salts, ephedrine base or salts;
  - (ii) in the case of ephedrine base or salts, pseudoephedrine salts; and
- (b) the medicinal product or products sold or supplied to a person at any one time must not in total contain more than—
  - (i) in the case of pseudoephedrine salts, 720mg pseudoephedrine salts;
  - (ii) in the case of ephedrine base or salts, 180mg ephedrine base or salts. ]

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Textual Amendments

F114 Art. 5B inserted (1.4.2008) by The Prescription Only Medicines (Human Use) Amendment Order 2008

(S.I. 2008/464), arts. 1(1), 7
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# Circumstances in which controlled drugs and medicinal products authorized by the European Community are not prescription only medicines

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Textual Amendments
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F115 Art. 6 revoked (1.4.2002) by The Prescription Only Medicines (Human Use) Amendment Order 2002
(S.I. 2002/549), arts. 1(1), 11
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# Exemption for parenteral administration in an emergency to human beings of certain prescription only medicines

7. The restriction imposed by section 58(2)(b) (restriction on administration) shall not apply to the administration to human beings of any of the following medicinal products for parenteral administration–

Adrenaline Injection 1 in 1000 (1 mg in 1 ml)

Atropine Sulphate Injection

[<sup>F116</sup>Atropine sulphate and obidoxime chloride injection]

[<sup>F116</sup>Atropine sulphate and pralidoxime chloride injection]

[<sup>F116</sup>Atropine sulphate, pralidoxime mesilate and avizafone injection]

[<sup>F117</sup>Chlorphenamine Injection]

[<sup>F118</sup>Dicobalt Edetate Injection]

F119 ...

F119 ...

Glucagon Injection

[F120Glucose Injection]

Hydrocortisone Injection

[<sup>F121</sup>Naloxone Hydrochloride]

[<sup>F116</sup>Pralidoxime chloride injection]

[<sup>F116</sup>Pralidoxime mesilate injection]

Promethazine Hydrochloride Injection

Snake Venom Antiserum

Sodium Nitrite Injection

Sodium Thiosulphate Injection

Sterile Pralidoxime

where the administration is for the purpose of saving life in an emergency.

#### **Textual Amendments**

- F116 Words in art. 7 inserted (19.11.2004) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 2004 (S.I. 2004/2693), arts. 1(1), 3
- F117 Words in art. 7 substituted (30.6.2005) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), 4(a)
- F118 Words in art. 7 substituted (30.6.2005) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), 4(b)
- F119 Words in art. 7 omitted (30.6.2005) by virtue of The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), 4(c)
- **F120** Words in art. 7 substituted (17.1.2011) by The Prescription Only Medicines (Human Use) Amendment Order 2010 (S.I. 2010/2998), arts. 1(1), 2
- F121 Words in art. 7 substituted (30.6.2005) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), 4(e)

## [<sup>F122</sup>Exemptions for administration of smallpox vaccine

**7A.**—(1) The restriction imposed by section 58(2)(b) (restriction on administration) shall not apply to the administration to human beings of smallpox vaccine where the conditions specified in paragraph (2) or (3) are satisfied.

- (2) The conditions referred to in this paragraph are—
  - (a) the vaccine has been supplied by, or on behalf of, or under arrangements made by-
    - (i) the Secretary of State,
    - (ii) the Scottish Ministers,
    - (iii) the National Assembly for Wales,
    - (iv) the Department of Health, Social Services and Public Safety,
    - (v) an NHS body; and
  - (b) the vaccine is administered for the purpose of providing protection against smallpox virus in the event of a suspected or confirmed case of smallpox in the United Kingdom.
- (3) The conditions referred to in this paragraph are—
  - (a) the vaccine has been supplied by, or on behalf of, or under arrangements made by Her Majesty's Forces;
  - (b) the vaccine is administered for the purpose of providing protection against smallpox virus to—
    - (i) members of Her Majesty's Forces; or
    - (ii) other persons employed or engaged by those Forces.
- (4) For the purposes of this regulation, "NHS body" means-
  - (a) the Common Services Agency,
  - (b) a Strategic Health Authority, Health Authority or Special Health Authority,
  - (c) a Primary Care Trust,
  - (d) a Local Health Board, or
  - (e) an NHS trust or NHS foundation trust]

#### **Textual Amendments**

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F122 Art. 7A inserted (19.11.2004) by The Prescription Only Medicines (Human Use) Amendment (No. 3)
Order 2004 (S.I. 2004/2693), arts. 1(1), 4
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## [<sup>F123</sup>Exemption for administration by operators

**7B.**—(1) The restriction imposed by section 58(2)(b) (restriction on administration) shall not apply to—

- (a) a radioactive medicinal product, administration of which results in a medical exposure; or
- (b) any other prescription only medicine if it is being administered in connection with a medical exposure,

where the conditions specified in paragraph (2) are satisfied.

(2) The conditions referred to in paragraph (1) are that—

- (a) the radioactive medicinal product or other prescription only medicine is administered by an operator acting in accordance with the procedures and protocols referred to in regulation 4(1) and (2) of the Ionising Radiation (Medical Exposure) Regulations 2000 which apply to the exposure referred to in paragraph (1);
- (b) that medical exposure has been authorised by an IRME practitioner or, where it is not practicable for an IRME practitioner to authorise the exposure, by an operator acting in accordance with written guidelines issued by an IRME practitioner;
- (c) the IRME practitioner is the holder of a certificate granted pursuant to the Medicines (Administration of Radioactive Substances) Regulations 1978(5);
- (d) the radioactive medicinal product or other prescription only medicine is not a controlled drug; and
- (e) in the case of a prescription only medicine which is not a radioactive medicinal product, it is specified in the protocols referred to in sub-paragraph (a).]

#### **Textual Amendments**

F123 Art. 7B inserted (17.11.2006) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2006 (S.I. 2006/2807), arts. 1(1), 3

#### Exemptions for emergency sale or supply

**8.**—(1) The restrictions imposed by section 58(2)(a) (restriction on sale and supply) shall not apply to the sale or supply of a prescription only medicine by a person lawfully conducting a retail pharmacy business where the conditions specified in paragraph (2) are satisfied.

- (2) The conditions referred to in paragraph (1) are-
  - (a) that the pharmacist by or under whose supervision the prescription only medicine is to be sold or supplied is satisfied that the sale or supply has been requested by a doctor [<sup>F124</sup>, a supplementary prescriber][<sup>F125</sup>a community practitioner nurse prescriber, a nurse independent prescriber [<sup>F126</sup>, an optometrist independent prescriber][<sup>F127</sup>, dentist] or a pharmacist independent prescriber] who by reason of an emergency is unable to furnish a prescription immediately;

- (b) that the doctor [<sup>F128</sup>, supplementary prescriber], [<sup>F129</sup>community practitioner nurse prescriber, nurse independent prescriber [<sup>F130</sup>, optometrist independent prescriber][<sup>F127</sup>, dentist] or pharmacist independent prescriber] has undertaken to furnish the person lawfully conducting a retail pharmacy business with a prescription within 72 hours of the sale or supply;
- (c) that the prescription only medicine is sold or supplied in accordance with the directions of the doctor [<sup>F131</sup>, supplementary prescriber], [<sup>F132</sup>community practitioner nurse prescriber, nurse independent prescriber [<sup>F133</sup>, optometrist independent prescriber][<sup>F127</sup>, dentist] or pharmacist independent prescriber] requesting it;
- (d) subject to paragraph (5), that the prescription only medicine is not a controlled drug specified in Schedule 1, 2 or 3 to the Misuse of Drugs Regulations;
- (e) that an entry is made in the record kept under regulation 6 of the Medicines (Sale or Supply) (Miscellaneous Provisions) Regulations 1980(12) within the time specified in that regulation stating the particulars required under paragraph 1 of Schedule 2 to those Regulations.

(3) The restrictions imposed by section 58(2)(a) shall not apply to the sale or supply of a prescription only medicine by a person lawfully conducting a retail pharmacy business where the conditions specified in paragraph (4) are satisfied.

- (4) The conditions referred to in paragraph (3) are-
  - (a) [<sup>F134</sup>subject to paragraph (6)] that the pharmacist by or under whose supervision the prescription only medicine is to be sold or supplied has interviewed the person requesting a prescription only medicine and has satisfied himself-
    - (i) that there is an immediate need for the prescription only medicine requested to be sold or supplied and that it is impracticable in the circumstances to obtain a prescription without undue delay,
    - (ii) that treatment with the prescription only medicine requested has on a previous occasion been prescribed by a doctor [<sup>F135</sup>, supplementary prescriber], [<sup>F136</sup>community practitioner nurse prescriber, nurse independent prescriber [<sup>F137</sup>, optometrist independent prescriber][<sup>F138</sup>, dentist] or pharmacist independent prescriber] for the person requesting it, and
    - (iii) as to the dose which in the circumstances it would be appropriate for that person to take;
  - (b) that no greater quantity of the prescription only medicine than will provide 5 days' treatment [<sup>F139</sup>in the case of a controlled drug or 30 days in any other case] is sold or supplied except that where the prescription only medicine-
    - (i) is [<sup>F140</sup>a preparation of insulin,] an aerosol for the relief of asthma, an ointment or cream, and has been made up for sale in a container elsewhere than at the place of sale or supply, the smallest pack that the pharmacist has available for sale or supply may be sold or supplied,
    - (ii) is an oral contraceptive, a quantity sufficient for a full treatment cycle may be sold or supplied,
    - (iii) is an antibiotic for oral administration in liquid form, the smallest quantity that will provide a full course of treatment may be sold or supplied;

<sup>(12)</sup> S.I. 1980/1923, amended by S.I. 1997/1831.

- (c) subject to paragraph (5), that the prescription only medicine does not consist of or contain a substance specified in Schedule 4 to this Order and is not a controlled drug specified in Schedule 1, 2 or 3 to the Misuse of Drugs Regulations;
- (d) that an entry is made in the record kept under regulation 6 of the Medicines (Sale or Supply) (Miscellaneous Provisions) Regulations 1980 within the time specified in that regulation stating the particulars required under paragraph 3 of Schedule 2 to those Regulations;
- (e) that the container or package of the prescription only medicine is labelled so as to show-
  - (i) the date on which the prescription only medicine is sold or supplied,
  - (ii) the name, quantity and, except where it is apparent from the name, the pharmaceutical form and strength of the prescription only medicine,
  - (iii) the name of the person requesting the prescription only medicine,
  - (iv) the name and address of the registered pharmacy from which the prescription only medicine is sold or supplied, and
  - (v) the words "Emergency Supply".

(5) The conditions specified in paragraphs (2)(d) and (4)(c) shall not apply where the prescription only medicine consists of or contains phenobarbitone or phenobarbitone sodium (but no other substance specified in Schedule 4 to this Order or Schedule 1, 2 or 3 to the Misuse of Drugs Regulations) and is sold or supplied for use in the treatment of epilepsy.

 $[^{F141}(6)$  Paragraph (4)(a) does not apply in relation to the sale or supply of a prescription only medicine where—

(a) the sale or supply is made—

(i) whilst a disease is, or

(ii) in anticipation of a disease being imminently,

pandemic and a serious risk, or potentially a serious risk, to human health; and

- (b) the pharmacist by or under whose supervision the prescription only medicine is to be sold or supplied has satisfied himself—
  - (i) that treatment with the prescription only medicine requested has on a previous occasion been prescribed by a doctor, supplementary prescriber, community practitioner nurse prescriber, nurse independent prescriber, optometrist independent prescriber or pharmacist independent prescriber for the person to be treated with it; and
  - (ii) as to the dose which in the circumstances it would be appropriate for that person to take.]

#### **Textual Amendments**

- **F124** Words in art. 8(2)(a) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 8(a)(i)
- F125 Words in art. 8(2)(a) substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), 7(a)(i)
- F126 Words in art. 8(2)(a) inserted (4.6.2008) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2008 (S.I. 2008/1161), arts. 1(1), 4(a)(i)
- F127 Word in art. 8(2) inserted (8.5.2009) by The Medicines for Human Use (Prescribing)(Miscellaneous Amendments) Order 2009 (S.I. 2009/1165), arts. 1, 3(a)
- **F128** Words in art. 8(2)(b) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 8(a)(ii)

F129	Words in art. 8(2)(b) substituted (1.5.2006) by The Medicines for Human Use (Prescribing)	
	(Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), 7(a)(ii)	
F130	Words in art. 8(2)(b) inserted (4.6.2008) by The Medicines for Human Use (Prescribing)	
	(Miscellaneous Amendments) Order 2008 (S.I. 2008/1161), arts. 1(1), 4(a)(ii)	
F131	Words in art. 8(2)(c) inserted (4.4.2003) by The Prescription Only Medicines (Human Use)	
	Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 8(a)(iii)	
F132	2 Words in art. 8(2)(c) substituted (1.5.2006) by The Medicines for Human Use (Prescribing)	
	(Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), <b>7(a)(iii)</b>	
F133	Words in art. 8(2)(c) inserted (4.6.2008) by The Medicines for Human Use (Prescribing)	
	(Miscellaneous Amendments) Order 2008 (S.I. 2008/1161), arts. 1(1), 4(a)(iii)	
F134	Words in art. 8(4)(a) inserted (8.5.2009) by The Medicines for Human Use (Prescribing)	
-	(Miscellaneous Amendments) Order 2009 (S.I. 2009/1165), arts. 1, <b>3(b)(i)</b>	
F135	Words in art. 8(4)(a)(ii) inserted (4.4.2003) by The Prescription Only Medicines (Human Use)	
	Amendment Order 2003 (S.I. 2003/696), arts. 1(1), <b>8(b)</b>	
F136	Words in art. 8(4)(a)(ii) substituted (1.5.2006) by The Medicines for Human Use (Prescribing)	
	(Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), <b>7(b</b> )	
F137	Words in art. 8(4)(a)(ii) inserted (4.6.2008) by The Medicines for Human Use (Prescribing)	
-	(Miscellaneous Amendments) Order 2008 (S.I. 2008/1161), arts. 1(1), <b>4(b)</b>	
F138	Word in art. 8(4)(a)(ii) inserted (8.5.2009) by The Medicines for Human Use (Prescribing)	
	(Miscellaneous Amendments) Order 2009 (S.I. 2009/1165), arts. 1, <b>3(b)(ii)</b>	
F139	Words in art. 8(4)(b) inserted (8.5.2009) by The Medicines for Human Use (Prescribing)	
	(Miscellaneous Amendments) Order 2009 (S.I. 2009/1165), arts. 1, <b>3(c)</b>	
F140	Words in art. 8(4)(b)(i) inserted (16.9.1998) by The Prescription Only Medicines (Human Use)	
	Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), <b>2</b>	
F141	Art. 8(6) inserted (8.5.2009) by The Medicines for Human Use (Prescribing)(Miscellaneous	
	Amendments) Order 2009 (S.I. 2009/1165), arts. 1, <b>3(d)</b>	

#### Exemption for non-parenteral administration to human beings

**9.** The restriction imposed by section 58(2)(b) (restriction on administration) shall not apply to the administration to human beings of a prescription only medicine which is not for parenteral administration.

#### Exemption for medicinal products at high dilutions

10.— $[^{F142}(1)]$  The restrictions imposed by section 58(2) (restrictions on sale, supply and administration) shall not apply to the sale, supply or administration of a medicinal product which is not for parenteral administration and which consists of or contains, any of the substances listed in column 1 of Schedule 1 [<sup>F143</sup>or in Schedule 2], only one or more unit preparation of such substances, if–

- (a) each such unit preparation has been diluted to at least one part in a million (6x), and the person selling, supplying or administering the medicinal product has been requested by or on behalf of a particular person and in that person's presence to use his own judgment as to the treatment required; or
- (b) each such unit preparation has been diluted to at least one part in a million million (6c).

 $[^{F144}(2)$  The restrictions imposed by section 58(2) (restrictions on sale, supply and administration) shall not apply to the sale, supply or administration of a medicinal product which is not for parenteral administration and which consists of or contains solely one or more unit preparations of the following substances—

Aconite

Arsenic Trioxide

Belladonna Herb

Ignatia Bean

Nux Vomica Seed,

if each such unit preparation has been diluted to at least one part in a million (6x)]

#### **Textual Amendments**

- F142 Art. 10 renumbered as art. 10(1) (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 9
- F143 Words in art. 10(1) substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), 8
- F144 Art. 10(2) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 9

#### **Exemptions for certain persons**

**11.**—(1) The restrictions imposed by section 58(2)(a) (restrictions on sale and supply) shall not apply–

- (a) to the sale or supply by a person listed in column 1 of Part I of Schedule 5 of the prescription only medicines listed in relation to that person in column 2 of that Part where the conditions specified in the corresponding paragraphs in column 3 of that Part are satisfied;
- (b) to the supply by a person listed in column 1 of Part II of Schedule 5 of the prescription only medicines listed in relation to that person in column 2 of that Part where the conditions specified in the corresponding paragraphs in column 3 of that Part are satisfied.

(2) The restriction imposed by section 58(2)(b) (restriction on administration) shall not apply to the administration by a person listed in column 1 of Part III of Schedule 5 of the prescription only medicines for parenteral administration listed in relation to that person in column 2 of that Part where the conditions specified in the corresponding paragraphs in column 3 of that Part are satisfied.

## [<sup>F145</sup>Exemption for sale or supply in hospitals

**12.**—(1) Subject to paragraph (3), the restrictions imposed by section 58(2)(a) (restrictions on sale and supply) shall not apply to the sale or supply of a prescription only medicine in the course of the business of a hospital where the medicine is sold or supplied for the purpose of being administered (whether in the hospital or elsewhere) to a particular person in accordance with directions satisfying the conditions specified in paragraph (2).

(2) The conditions specified in paragraph (1) are that the directions—

- (a) are in writing;
- (b) relate to the particular person to whom the medicine is to be administered; and
- (c) are given by a person (other than a veterinary surgeon or veterinary practitioner) who is an appropriate practitioner in relation to that medicine.

(3) Such directions may be given by <sup>F146</sup>... a supplementary prescriber only where he complies with any condition as to the cases or circumstances in which he may give a prescription for that medicine specified by virtue of article <sup>F147</sup>... 3B, as if the directions are a prescription.

(4) The exemption in paragraph (1) applies notwithstanding that the directions do not satisfy the conditions specified in article 15(2).]

#### **Textual Amendments**

- F145 Art. 12 substituted (31.1.2004) by The Prescription Only Medicines (Human Use) Amendment Order 2004 (S.I. 2004/2), arts. 1(1), 2
- F146 Words in art. 12(3) omitted (1.4.2008) by virtue of The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), 8(a)
- F147 Words in art. 12(3) omitted (1.4.2008) by virtue of The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), **8(b)**

# [<sup>F148</sup>Exemptions for the supply and administration of prescription only medicines by national health service bodies

**12A.**—(1) The restrictions imposed by section 58(2)(a) (restrictions on sale and supply) shall not apply to the supply of a prescription only medicine by—

- (a) the Common Services Agency;
- (b) a [<sup>F149</sup>Strategic Health Authority,] Health Authority or Special Health Authority;
- (c) an NHS trust [<sup>F150</sup>or NHS foundation trust];
- (d) a Primary Care Trust; or
- (e) where sub-paragraphs (a) to (d) do not apply, a person other than an excepted person, pursuant to an arrangement made with one of the persons specified in those sub-paragraphs for the supply of prescription only medicines,

where the medicine is supplied for the purpose of being administered to a particular person in accordance with the written directions of a doctor or dentist relating to that person notwithstanding that those directions do not satisfy the conditions specified in article 15(2).

(2) The restrictions imposed by section 58(2) (restrictions on sale, supply and administration) shall not apply to the supply or, as the case may be, the administration of a prescription only medicine by—

- (a) the Common Services Agency;
- (b) a [<sup>F151</sup>Strategic Health Authority,] Health Authority or Special Health Authority;
- (c) an NHS trust [<sup>F152</sup> or NHS foundation trust];
- (d) a Primary Care Trust; or
- (e) where sub-paragraphs (a) to (d) do not apply, a person other than an excepted person, pursuant to an arrangement made with one of the persons specified in those sub-paragraphs for the supply or, as the case may be, the administration of prescription only medicines,

where the medicine is supplied for the purpose of being administered, or, as the case may be, is administered, to a particular person in accordance with a Patient Group Direction and where the conditions specified in paragraph (3) are satisfied.

- (3) The conditions referred to are that—
  - (a) the Patient Group Direction relates to the supply or, as the case may be, the administration, by the person who supplies or administers the medicine, of a description or class of prescription only medicine, and the Direction has effect at the time at which the medicine is supplied or, as the case may be, is administered;
  - (b) the Patient Group Direction contains the particulars specified in Part I of Schedule 7 to this Order (but with the omission of paragraph (c) of that Part in the case of a Direction which relates to administration only);

- (c) the Patient Group Direction is signed on behalf of the person specified in column 2 of the Table in Part II of Schedule 7 to this Order ("the authorising person") against the entry in column 1 of that Table for the class of person by whom the medicine is supplied or administered;
- (d) the individual who supplies or, as the case may be, administers the medicine belongs to one of the classes of individual specified in Part III of Schedule 7 to this Order, and is designated in writing, on behalf of the authorising person, for the purpose of the supply or, as the case may be, the administration of prescription only medicines under the Patient Group Direction; and
- (e) at the time at which the medicine is supplied or, as the case may be, is administered, a product licence, a marketing authorization or a homoeopathic certificate of registration has effect in respect of it.
- (4) In this article, "excepted person" means-
  - (a) a doctor or dentist; or
  - (b) a person lawfully conducting a retail pharmacy business within the meaning of section 69.

#### **Textual Amendments**

- F148 Arts. 12A-12C inserted (9.8.2000) by The Prescription Only Medicines (Human Use) Amendment Order 2000 (S.I. 2000/1917), arts. 1(1), 2(d)
- **F149** Words in art. 12A(1)(b) inserted (1.10.2002) by The National Health Service Reform and Health Care Professions Act 2002 (Supplementary, Consequential etc. Provisions) Regulations 2002 (S.I. 2002/2469), reg. 1, Sch. 1 para. 73(2)(a)
- F150 Words in art. 12A(1)(c) inserted (1.4.2004) by The Health and Social Care (Community Health and Standards) Act 2003 (Supplementary and Consequential Provision) (NHS Foundation Trusts) Order 2004 (S.I. 2004/696), arts. 1(1)(b), 3(2), Sch. 2
- F151 Words in art. 12A(2)(b) inserted (1.10.2002) by The National Health Service Reform and Health Care Professions Act 2002 (Supplementary, Consequential etc. Provisions) Regulations 2002 (S.I. 2002/2469), reg. 1, Sch. 1 para. 73(2)(b)
- F152 Words in art. 12A(2)(c) inserted (1.4.2004) by The Health and Social Care (Community Health and Standards) Act 2003 (Supplementary and Consequential Provision) (NHS Foundation Trusts) Order 2004 (S.I. 2004/696), arts. 1(1)(b), 3(2), Sch. 2

# Exemption for health professionals who supply or administer prescription only medicines under a Patient Group Direction in order to assist doctors or dentists in providing national health services

**12B.**—(1) The restrictions imposed by section 58(2) (sale, supply and administration) shall not apply to the supply or, as the case may be, the administration of a prescription only medicine by an individual belonging to one of the classes specified in Part III of Schedule 7 to this Order where—

- (a) the individual supplies or, as the case may be, administers the medicine in order to assist a doctor or dentist in the provision of, respectively, NHS primary medical services or NHS primary dental services;
- (b) the medicine is supplied for the purpose of being administered, or, as the case may be, is administered, in accordance with a Patient Group Direction; and
- (c) the conditions specified in paragraph (2) are satisfied.

(2) The conditions referred to are that—

(a) the Patient Group Direction relates to the supply or, as the case may be, the administration of a description or class of prescription only medicine in order to assist the doctor or dentist

in question in providing the services referred to in paragraph (1)(a) (whether or not it also relates to such supply or administration in order to assist any other doctor or dentist);

- (b) the Patient Group Direction has effect at the time at which the medicine is supplied or, as the case may be, is administered;
- (c) the Patient Group Direction contains the particulars specified in Part I of Schedule 7 to this Order (but with the omission of paragraph (c) of that Part in the case of a Direction which relates to administration only);
- (d) the Patient Group Direction is signed—
  - (i) by the doctor or dentist in question or, alternatively, where the Direction also relates to supply or administration in order to assist one or more other doctors or dentists, as referred to in sub-paragraph (a), by one of those other doctors or dentists; and
  - [<sup>F153</sup>(ii) on behalf of the Primary Care Trust, Local Health Board, Health Board or Health and Social Services Board that is responsible for the arrangements under which the services are provided]
- (e) the individual referred to in paragraph (1) is designated in writing for the purpose of the supply or, as the case may be, the administration of prescription only medicines under the Patient Group Direction, by the doctor or dentist in question, or, alternatively, where the Direction also relates to supply or administration in order to assist one or more other doctors or dentists, as referred to in sub-paragraph (a), by one of those other doctors or dentists; and
- (f) at the time at which the medicine is supplied or, as the case may be, is administered, a product licence, a marketing authorization or a homoeopathic certificate of registration has effect in respect of it.
- (3) In this article—
  - (a) a reference to the provision of NHS primary dental services shall be construed as a reference to—
    - [<sup>F154</sup>(i) in relation to England and Wales, the provision of primary dental services under Part 1 of the National Health Service Act 1977;]
      - (ii) in relation to Scotland, the provision of general dental services under Part II of the National Health Service (Scotland) Act 1978, or the performance of personal dental services in connection with a pilot scheme under the National Health Service (Primary Care) Act 1997; and
      - (iii) in relation to Northern Ireland, the provision of general dental services under the Health and Personal Social Services (Northern Ireland) Order 1972, or the performance of personal dental services in connection with a pilot scheme under the Health Services (Primary Care) (Northern Ireland) Order 1997;
  - (b) a reference to the provision of NHS primary medical services shall be construed as a reference to—
    - [<sup>F155</sup>(i) in relation to England and Wales, the provision of primary medical services under Part I of the National Health Service Act 1977;]
    - [<sup>F156</sup>(ii) in relation to Scotland, the provision of primary medical services under Part I of the National Health Service (Scotland) Act 1978; and]
    - [<sup>F157</sup>(iii) in relation to Northern Ireland, the provision of primary medical services under Article 15B or Part VI of the Health and Personal Social Services (Northern Ireland) Order 1972.]

#### Textual Amendments

- F148 Arts. 12A-12C inserted (9.8.2000) by The Prescription Only Medicines (Human Use) Amendment Order 2000 (S.I. 2000/1917), arts. 1(1), 2(d)
- **F153** Art. 12B(2)(d)(ii) substituted (1.4.2006) by The General Dental Services, Personal Dental Services and Abolition of the Dental Practice Board Transitional and Consequential Provisions Order 2006 (S.I. 2006/562), art. 1(1), Sch. 1 para. 6(2)(a)
- F154 Art. 12B(3)(a)(i) substituted (1.4.2006) by The General Dental Services, Personal Dental Services and Abolition of the Dental Practice Board Transitional and Consequential Provisions Order 2006 (S.I. 2006/562), art. 1(1), Sch. 1 para. 6(2)(b)
- F155 Art. 12B(3)(b)(i) substituted (E.) (1.4.2004) by The General Medical Services and Personal Medical Services Transitional and Consequential Provisions Order 2004 (S.I. 2004/865), art. 1(1), Sch. 1 para. 18(2); and substituted (W.) by The General Medical Services Transitional and Consequential Provisions (Wales) (No. 2) Order 2004 (S.I. 2004/1016), art. 1(1), Sch. 1 para. 18(2)
- F156 Art. 12B(3)(b)(ii) substituted (1.10.2004) by The Primary Medical Services (Scotland) Act 2004 (Consequential Amendments) Order 2004 (S.I. 2004/2261), art. 1(1), Sch. para. 2
- F157 Art. 12B(3)(b)(iii) substituted (18.11.2004) by The Primary Medical Services (Northern Ireland) Order 2004 (Consequential Amendments) Order 2004 (S.I. 2004/3038), art. 1(1), Sch. para. 3

# Exemption for persons conducting a retail pharmacy business who supply or administer prescription only medicines under a Patient Group Direction

12C.—(1) The restrictions imposed by section 58(2) (restrictions on sale, supply and administration) shall not apply to the [<sup>F158</sup>sale or] supply or, as the case may be, the administration of a prescription only medicine by a person lawfully conducting a retail pharmacy business within the meaning of section 69 where—

[<sup>F159</sup>(a) the medicine—

- (i) is supplied, or as the case may be, is administered by such a person pursuant to an arrangement made with—
  - (aa) a body referred to in article 12A(a) to (d),
  - (bb) a force or service referred to in article 12E(1)(a)(i) to (iii), or
  - (cc) Her Majesty's Forces,

for the supply or, as the case may be, the administration of prescription only medicines; or

- (ii) is sold or supplied or, as the case may be, is administered by such a person pursuant to an arrangement made with an authority or person carrying on the business of an establishment or agency referred to in article 12D(1) for the sale or supply or, as the case may be, the administration of prescription only medicines;]
- (b) the medicine is [<sup>F160</sup>sold or] supplied for the purpose of being administered, or, as the case may be, is administered, to a particular person in accordance with a Patient Group Direction; and
- (c) the conditions specified in paragraph (2) are satisfied.
- (2) The conditions referred to are that—
  - (a) the Patient Group Direction relates to the [<sup>F161</sup>sale or] supply or, as the case may be, the administration of a description or class of prescription only medicine by the person lawfully conducting a retail pharmacy business who [<sup>F162</sup>sells or] supplies or, as the case

may be, administers the prescription only medicine, and the Direction has effect at the time at which the medicine is [<sup>F163</sup>sold or] supplied or, as the case may be, is administered;

- (b) the Patient Group Direction contains the particulars specified in Part I of Schedule 7 to this Order (but with the omission of paragraph (c) of that Part in the case of a Direction which relates to administration only);
- [<sup>F164</sup>(c) the Patient Group Direction is signed—
  - (i) in the case of an arrangement with a body referred to in article 12A(a) to (d), on behalf of that body,
  - (ii) in the case of an arrangement with an authority or person carrying on the business of an establishment or agency referred to in article 12D(1), by or on behalf of the relevant provider and, if there is a relevant manager for the establishment or agency, that manager,
  - (iii) in the case of an arrangement with a prison service, by or on behalf of the person specified in column 2 of the Table in Part IIA of Schedule 7 to this Order against the entry in column 1 of that Table for that service,
  - (iv) in the case of an arrangement with a police force or the Police Service of Northern Ireland—
    - (a) by or on behalf of the person specified in column 2 of the Table in Part IIA of Schedule 7 to this Order against the entry in column 1 of that Table for that force or service, and
    - (b) a doctor who is not employed or engaged by, and who does not provide services under arrangements made with, any police force or the Police Service of Northern Ireland, and
  - (v) in the case of an arrangement with Her Majesty's Forces, by or on behalf of a person specified in column 2 of the Table in Part IIA of Schedule 7 to this Order against the entry in column 1 of that Table for Her Majesty's Forces;]
- [ where the prescription only medicine is administered by the person lawfully conducting a <sup>F165</sup>(cc) retail pharmacy business within the meaning of section 69, the individual who administers the medicine belongs to one of the classes of individual specified in Part III of Schedule 7 to this Order, and is [<sup>F166</sup>designated in writing—
  - (i) in the case of an arrangement with a body referred to in article 12A(a) to (d), on behalf of that body,
  - (ii) in the case of an arrangement with an authority or person carrying on the business of an establishment or agency referred to in article 12D(1), by or on behalf of the relevant provider or, if there is a relevant manager for the establishment or agency, that relevant manager,
  - (iii) in the case of an arrangement with a force or service referred to in article 12E(1)(a)
    (i) to (iii), by or on behalf of the person specified in column 2 of the Table in Part IIA of Schedule 7 to this Order against the entry in column 1 of that Table for that force or service, or
  - (iv) in the case of an arrangement with Her Majesty's Forces, by or on behalf of a person specified in column 2 of the Table in Part IIA of Schedule 7 to this Order against the entry in column 1 of that Table for Her Majesty's Forces,

for the purpose of the administration of prescription only medicines under the Patient Group Direction; and]]

(d) at the time at which the medicine is [<sup>F167</sup>sold or] supplied or, as the case may be, is administered, a product licence, a marketing authorization or a homoeopathic certificate of registration has effect in respect of it.]

#### **Textual Amendments**

- F148 Arts. 12A-12C inserted (9.8.2000) by The Prescription Only Medicines (Human Use) Amendment Order 2000 (S.I. 2000/1917), arts. 1(1), 2(d)
- **F158** Words in art. 12C(1) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **3(a)(i)**
- F159 Art. 12C(1)(a) substituted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 3(a)(ii)
- **F160** Words in art. 12C(1)(b) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **3(a)(iii)**
- F161 Words in art. 12C(2)(a) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 3(b)(i)(aa)
- F162 Words in art. 12C(2)(a) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 3(b)(i)(bb)
- **F163** Words in art. 12C(2)(a) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **3(b)(i)(cc)**
- F164 Art. 12C(2)(c) substituted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 10(b)(i)
- **F165** Art. 12C(2)(cc) inserted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), **3(1)**
- **F166** Words in art. 12C(2)(cc) substituted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **10(b)(ii)**
- **F167** Words in art. 12C(2)(d) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **3(b)(ii)**

# [<sup>F168</sup>Exemption for the supply and administration of prescription only medicines by independent hospitals, clinics and agencies

**12D.**—(1) The restrictions imposed by section 58(2) (restrictions on sale, supply and administration) shall not apply to the [<sup>F169</sup>sale or] supply or, as the case may be, the administration of a prescription only medicine in the course of the business of—

- [<sup>F170</sup>(a) an independent hospital,
  - (b) [<sup>F171</sup>in Wales, Scotland or Northern Ireland,] an independent clinic,
  - (c) an independent medical agency, or
  - (d) in Northern Ireland, a nursing home,]

where the medicine is  $[^{F172}$ sold or] supplied for the purpose of being administered, or, as the case may be, is administered, to a particular person in accordance with a Patient Group Direction and where the conditions specified in paragraph (2) are satisfied.

- (2) The conditions referred to are that—
  - (a) the Patient Group Direction relates to the [<sup>F173</sup>sale or] supply or, as the case may be, the administration, by the person who [<sup>F174</sup>sells or] supplies or administers the medicine, of a description or class of prescription only medicine, and the Direction has effect at the time at which the medicine is [<sup>F175</sup>sold or] supplied or, as the case may be, is administered;

- (b) the Patient Group Direction contains the particulars specified in Part I of Schedule 7 to this Order (but with the omission of paragraph (c) of that Part in the case of a Direction which relates to administration only);
- (c) the Patient Group Direction is signed—
  - (i) by or on behalf of the registered provider, and
  - (ii) if there is a relevant manager for the independent hospital, clinic or agency, or nursing home, by that manager;
- (d) the individual who [<sup>F176</sup>sells or] supplies or, as the case may be, administers the medicine belongs to one of the classes of individual specified in Part III of Schedule 7 to this Order, and is designated in writing—
  - (i) by or on behalf of the registered provider, or
  - (ii) if there is a relevant manager for the independent hospital, clinic or agency, or nursing home, by that manager,

for the purpose of the [<sup>F177</sup>sale or] supply or, as the case may be, the administration of prescription only medicines under the Patient Group Direction; and

(e) at the time at which the medicine is [<sup>F178</sup>sold or] supplied or, as the case may be, is administered, a product licence, a marketing authorization or a homoeopathic certificate of registration has effect in respect of it.

#### **Textual Amendments**

- F168 Arts. 12D, 12E inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 11
- F169 Words in art. 12D(1) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 4(a)(i)
- F170 Art. 12D(1)(a)-(d) substituted for art. 12D(1)(a)(b) (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 4(a)(ii)
- F171 Words in art. 12D(1)(b) inserted (1.10.2010) by The Health and Social Care Act 2008 (Miscellaneous Consequential Amendments) Order 2010 (S.I. 2010/1881), arts. 1(1), 9(3)
- F172 Words in art. 12D(1) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 4(a)(iii)
- F173 Words in art. 12D(2)(a) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 4(b)(i)(aa)
- F174 Words in art. 12D(2)(a) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 4(b)(i)(bb)
- F175 Words in art. 12D(2)(a) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 4(b)(i)(cc)
- F176 Words in art. 12D(2)(d) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 4(b)(ii)(aa)
- F177 Words in art. 12D(2)(d) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 4(b)(ii)(bb)
- **F178** Words in art. 12D(2)(e) inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), **4(b)(iii)**

#### Exemption for health professionals who supply or administer prescription only medicines under a Patient Group Direction in order to assist the provision of health care by or on behalf of the police, the prison services or the armed forces

**12E.**—(1) The restrictions imposed by section 58(2) (sale, supply and administration) shall not apply to the supply or, as the case may be, the administration of a prescription only medicine by an individual belonging to one of the classes specified in Part III of Schedule 7 to this Order where—

- (a) the individual supplies or, as the case may be, administers the medicine in order to assist the provision of health care by, on behalf of, or under arrangements made by—
  - (i) a police force in England, Wales or Scotland,
  - (ii) the Police Service of Northern Ireland,
  - (iii) a prison service, or
  - (iv) Her Majesty's Forces;
- (b) the medicine is supplied for the purpose of being administered, or, as the case may be, is administered, in accordance with a Patient Group Direction; and
- (c) the conditions specified in paragraph (2) are satisfied.
- (2) The conditions referred to are that—
  - (a) the Patient Group Direction relates to the supply or, as the case may be, the administration of a description or class of prescription only medicine in order to assist the provision of health care by, or on behalf of, or under arrangements made by the police force or service, the prison service or, as the case may be, Her Majesty's Forces;
  - (b) the Patient Group Direction has effect at the time at which the medicine is supplied or, as the case may be, is administered;
  - (c) the Patient Group Direction contains the particulars specified in Part I of Schedule 7 to this Order (but with the omission of paragraph (c) of that Part in the case of a Direction which relates to administration only);
  - (d) the Patient Group Direction is signed-
    - (i) by or on behalf of a person specified in column 2 of the Table in Part IIA of Schedule 7 to this Order ("the authorising person") against the entry in column 1 of that Table for the police force or service, the prison service or Her Majesty's Forces by whom, or on whose behalf, the health care is provided, or with whom arrangements are made for the provision of such care; and
    - (ii) in the case of a police force or the Police Service of Northern Ireland, by a doctor who is not employed or engaged by, and who does not provide services under arrangements made with, any police force or the Police Service of Northern Ireland;
  - (e) the individual referred to in paragraph (1) is designated in writing, by or on behalf of the authorising person, for the purpose of the supply or, as the case may be, the administration of prescription only medicines under the Patient Group Direction; and
  - (f) at the time at which the medicine is supplied or, as the case may be, is administered, a product licence, a marketing authorization or a homoeopathic certificate of registration has effect in respect of it.]

#### **Textual Amendments**

F168 Arts. 12D, 12E inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 11

# [<sup>F179</sup>Exemption for the supply of prescription only medicines in the event or anticipation of pandemic disease

**12F.** The restrictions imposed by section 58(2)(a) (restrictions on sale and supply) shall not apply to the supply of a prescription only medicine by an individual where that supply is—

- (a) made—
  - (i) whilst a disease is, or
  - (ii) in anticipation of a disease being imminently,
- pandemic and a serious risk, or potentially a serious risk, to human health; and
- (b) in accordance with a protocol which-
  - (i) is approved by-
    - (aa) the Ministers;
    - (bb) an NHS body (within the meaning of article 7A(4)); or
    - (cc) the Health Protection Agency established under section 1 of the Health Protection Agency Act 2004; and
  - (ii) contains criteria as to-
    - (aa) symptoms of, and treatment for, that disease;
    - (bb) the recording of the name of the person who supplies the prescription only medicine to the person to be treated (or to a person acting on that person's behalf) and of the evidence that the medicine was supplied to the person to be treated (or to a person acting on that person's behalf).]

#### **Textual Amendments**

**F179** Art. 12F inserted (8.5.2009) by The Medicines for Human Use (Prescribing)(Miscellaneous Amendments) Order 2009 (S.I. 2009/1165), arts. 1, 4

#### Exemption in cases involving another's default

**13.** The restrictions imposed by section 58(2)(a) (restrictions on sale and supply) shall not apply to the sale or supply of a prescription only medicine by a person who, having exercised all due diligence, believes on reasonable grounds that the product sold or supplied is not a prescription only medicine.

## [<sup>F180</sup>Exemptions relating to prescriptions given by [<sup>F181</sup>certain health professionals]

**13A.**— $[^{F182}(1)$  The restrictions imposed by section 58(2)(a) (restrictions in sale and supply) shall not apply to the sale or supply of a prescription only medicine by a pharmacist in accordance with a prescription given by—

- (a) another pharmacist,
- (b) a registered nurse, <sup>F183</sup>...
- (c) a registered midwife, <sup>F184</sup>...
- [<sup>F185</sup>(d) a person whose name is registered in the part of the register maintained by the Health Professions Council in pursuance of article 5 of the [<sup>F186</sup>Health and Social Work Professions Order 2001] relating to—

(i) chiropodists and podiatrists;

(ii) physiotherapists; or

(iii) radiographers: diagnostic or therapeutic, [<sup>F187</sup>or]]

[<sup>F188</sup>(e) a registered optometrist,]

who is not an appropriate practitioner in relation to that medicine where the pharmacist selling or supplying the medicine, having exercised all due diligence, believes on reasonable grounds that the person is such a practitioner.]

(2) The restrictions imposed by section 58(2)(a) (restrictions on sale and supply) shall not apply to the sale or supply of a prescription only medicine by a pharmacist in accordance with a prescription given by a <sup>F189</sup>... [<sup>F190</sup> supplementary prescriber] where the pharmacist, having exercised all due diligence, believes on reasonable grounds that [<sup>F191</sup>the <sup>F189</sup>...][<sup>F190</sup> supplementary prescriber] has complied with any condition with which he is required to comply by virtue of [<sup>F192</sup>[<sup>F193</sup>article] 3B].]

#### **Textual Amendments**

- **F180** Art. 13A inserted (1.4.2002) by The Prescription Only Medicines (Human Use) Amendment Order 2002 (S.I. 2002/549), arts. 1(1), 7
- F181 Words in art. 13A heading substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), 10(a)
- **F182** Art. 13A(1) substituted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **12(a)**
- **F183** Word in art. 13A(1)(b) omitted (7.4.2005) by virtue of The Medicines for Human Use (Prescribing) Order 2005 (S.I. 2005/765), arts. 1(1), **5(a)**
- F184 Word in art. 13A(1)(c) omitted (30.6.2005) by virtue of The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), 5(a)
- **F185** Art. 13A(1)(d) inserted (7.4.2005) by The Medicines for Human Use (Prescribing) Order 2005 (S.I. 2005/765), arts. 1(1), **5(c)**
- **F186** Words in art. 13A(1)(d) substituted (1.8.2012) by The Health and Social Care Act 2012 (Consequential Provision—Social Workers) Order 2012 (S.I. 2012/1479), art. 1(2), Sch. para. 13(3)
- F187 Word in art. 13A(d) inserted (30.6.2005) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), 5(b)
- **F188** Art. 13A(e) inserted (30.6.2005) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), **5(c)**
- **F189** Words in art. 13A(2) omitted (1.4.2008) by virtue of The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), 9(a)
- **F190** Words in art. 13A(2) inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **12(b)(i)**
- F191 Words in art. 13A(2) substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), 10(b)(ii)
- **F192** Words in art. 13A(2) substituted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), **12(b)(ii)**
- **F193** Word in art. 13A(2) substituted (1.4.2008) by The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), **9(b)**

#### Exemption in the case of a forged prescription

14. The restrictions imposed by section 58(2)(a) (restrictions on sale and supply) shall not apply to the sale or supply of a prescription only medicine by a pharmacist in accordance with a forged prescription where the pharmacist, having exercised all due diligence, believes on reasonable grounds that the prescription is genuine.

### Prescriptions

 $[^{F194}15.-(1)$  For the purposes of section 58(2)(a), and subject to paragraph (3), a prescription only medicine shall not be taken to be sold or supplied in accordance with a prescription given by an appropriate practitioner unless the conditions specified in paragraph (2) are fulfilled.

(2) The conditions referred to in paragraph (1) are that the prescription—

- (a) shall be signed in ink with his own name by the appropriate practitioner giving it;
- (b) shall, without prejudice to sub-paragraph (a), be written in ink or otherwise so as to be indelible, unless it is a health prescription which is not for a controlled drug specified in Schedule 1, 2 or 3 to the Misuse of Drugs Regulations, in which case it may be written by means of carbon paper or similar material;
- (c) shall contain the following particulars—
  - (i) the address of the appropriate practitioner giving it,
  - (ii) the appropriate date,
  - (iii) such particulars as indicate whether the appropriate practitioner giving it is a doctor, a dentist, a supplementary prescriber, [<sup>F195</sup>a community practitioner nurse prescriber, a nurse independent prescriber[<sup>F196</sup>, an optometrist independent prescriber], a pharmacist independent prescriber], a veterinary surgeon or a veterinary practitioner,
  - (iv) where the appropriate practitioner giving it is a doctor, dentist, a supplementary prescriber, [<sup>F197</sup>a community practitioner nurse prescriber, a nurse independent prescriber]<sup>F198</sup>, an optometrist independent prescriber] or a pharmacist independent prescriber] the name, address and the age, if under 12, of the person for whose treatment it is given, and
  - (v) where the appropriate practitioner giving it is a veterinary surgeon or a veterinary practitioner, the name and the address of the person to whom the prescription only medicine is to be delivered and a declaration by the veterinary surgeon or veterinary practitioner giving it that the prescription only medicine is prescribed for an animal or herd under his care;
- (d) shall not be dispensed after the end of the period of 6 months from the appropriate date, unless it is a repeatable prescription in which case it shall not be dispensed for the first time after the end of that period nor otherwise than in accordance with the directions contained in the repeatable prescription;
- (e) in the case of a repeatable prescription which does not specify the number of times it may be dispensed, shall not be dispensed on more than two occasions unless it is a prescription for an oral contraceptive in which case it may be dispensed 6 times before the end of the period of 6 months from the appropriate date.

(3) For the purposes of paragraph (1) the prescription may, as an alternative to fulfilling the conditions specified in paragraph (2)(a) and (b), fulfil instead the conditions specified in paragraph (4), unless the prescription is a health prescription for a controlled drug specified in Schedule 1, 2 or 3 to the Misuse of Drugs Regulations or is given by a veterinary surgeon or a veterinary practitioner.

(4) The conditions referred to in paragraph (3) are that the prescription shall be created in electronic form and signed with an advanced electronic signature and transferred to the person by whom it is dispensed as an electronic communication (including where it is so transferred through one or more intermediaries).

(5) The prohibition on sale or supply imposed by section 58(2)(a) shall not apply where a prescription only medicine is sold or supplied other than in accordance with a prescription given by an appropriate practitioner and –

- (a) the reason the sale or supply is not in accordance with such a prescription is that a condition specified in paragraph (2) or (4) is not fulfilled; and
- (b) the person selling or supplying the prescription only medicine has exercised all due diligence and believes on reasonable grounds that the condition is fulfilled.
- (6) In paragraph (2) "appropriate date" means-
  - (a) in the case of a health prescription, the date on which it was signed by the appropriate practitioner giving it or a date indicated by him as being the date before which it shall not be dispensed; and
  - (b) in every other case, the date on which the prescription was signed by the appropriate practitioner giving it,

and, for the purposes of sub-paragraphs (d) and (e) of that paragraph, where the health prescription bears both the date on which it was signed and a date indicated as being that before which it shall not be dispensed, the appropriate date is the later of those dates.

(7) In this Article—

"advanced electronic signature" means an electronic signature which is-

- (a) uniquely linked to the signatory,
- (b) capable of identifying the signatory,
- (c) created using means that the signatory can maintain under his sole control, and
- (d) which is linked to the data to which it relates in such a manner that any subsequent change of data is detectable;

"electronic communication" means a communication transmitted (whether from one person to another, from one device to another or from a person to a device or vice versa)—

- (a) by means of a telecommunication system (within the meaning of the Telecommunications Act 1984, or
- (b) by other means but while in an electronic form; and

"signatory" means the appropriate practitioner giving the prescription.]

#### **Textual Amendments**

- **F194** Art. 15 substituted (7.4.2005) by The Medicines for Human Use (Prescribing) Order 2005 (S.I. 2005/765), arts. 1(1), **6**
- F195 Words in art. 15(2)(c)(iii) substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), 11(a)
- **F196** Words in art. 15(2)(c)(iii) inserted (4.6.2008) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2008 (S.I. 2008/1161), arts. 1(1), **5(a)**
- F197 Words in art. 15(2)(c)(iv) substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), 11(b)
- F198 Words in art. 15(2)(c)(iv) inserted (4.6.2008) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2008 (S.I. 2008/1161), arts. 1(1), 5(b)

### Revocations

**16.**—(1) The Orders specified in Schedule 6 are revoked.

(2) In the Medicines (Prescription Only, Pharmacy and General Sale) Amendment Order 1989(13) articles 2 to 6 and Schedules 1 and 2 are revoked.

<sup>(</sup>**13**) S.I. 1989/1852.

Signed by authority of the Secretary of State for Health

Baroness Jay Minister of State, Department of Health

Win Griffiths Parliamentary Under Secretary of State, Welsh Office

Sam Galbraith Parliamentary Under Secretary of State, The Scottish Office

*Jeff Rooker* Minister of State, Ministry of Agriculture, Fisheries and Food

Sealed with the Official Seal of the Department of Health and Social Services for Northern Ireland on 22nd July 1997.



D. C. Gowdy Permanent Secretary Sealed with the Official Seal of the Department of Agriculture for Northern Ireland on 18th July 1997.



*P. Small* Permanent Secretary

### SCHEDULE 1

Articles 3(a), 5(1) and 10

# SUBSTANCES WHICH IF INCLUDED IN MEDICINAL PRODUCTS MAKE THOSE PRODUCTS PRESCRIPTION ONLY MEDICINES AND EXEMPTIONS FROM RESTRICTIONS ON THE SALE AND SUPPLY OF PRESCRIPTION ONLY MEDICINES

		from the restric only medicine	ctions on the sale and sup	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
[ <sup>F199</sup> Acampro	osate]			
Acarbose				
Acebutolol Hydrochlorid	de			
[ <sup>F199</sup> Aceclofe	enac			
Acemetacin				
Acetarsol				
Acetazolami	de			
Acetazolami Sodium	de			
Acetohexam	ide			
Acetylcholin Chloride	e0.2 per cent	External		
Acetylcystei	ne			
Acipimox				
Aciclovir	5.0 per cent	External For treatment of herpes simplex virus infections of the lips and face (Herpes labialis)		Container or package containing not more than 2g of medicinal product
Acitretin				
Aclarubicin Hydrochlorid	de			
Aconite	1.3 per cent	External		

		from the restr	ictions on the sale and sup	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratiuse or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity
Acrivastine			24 mg (MDD)	Container or package containing not more than 240mg of Acrivastine
Acrosoxacin				
Actinomycin C				
Actinomycin D				
[ <sup>F200</sup> Adapalen	e]			
Adenosine				
Adrenaline		(1) By inhaler		
		(2) External [ <sup>F201</sup> (except ophthalmic)]	l	
Adrenaline Acid		(1) By inhaler		
Tartrate		(2) External		
Adrenaline Hydrochlorid	e	(1) By inhaler		
		(2) External		
Adrenocortica Extract	al			
Albendazole				
Alclofenac				
Alclometason Dipropionate	le			
Alcuronium Chloride				
Aldesleukin				
Aldosterone				

		from the restrictions on the sale and supply of only medicines			
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceution form	Column 4 Treatment limitations on,	Column 5 Maximum quantity	
[ <sup>F199</sup> Alendron Sodium]	nate				
Alfacalcidol					
Alfuzosin Hydrochlorid	le				
Allergen Extracts					
Allopurinol					
Allyloestrend	ol				
[ <sup>F202</sup> Aloxiprin	1(1) 620 mg	(1) Non- effervescent tablets and capsules		<ul> <li>(1) The quantity sold or supplied in one container or package shall not exceed 32</li> <li>The quantity</li> </ul>	
				of non- effervescent tablets, capsules or a combination of both sold or supplied to a person at any one time shall not exceed 100	
		(2) All preparations other than non- effervescent tablets or capsules]			

	<i>Exemptions from the restrictions on the sale and supply of prescription only medicines</i>				
Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity	
Alphadolone Acetate					
Alphaxalone					
Alprenolol					
Alprenolol Hydrochloride					
Alprostadil					
Alseroxylon					
[ <sup>F200</sup> Altretamin	e]				
Amantadine Hydrochloride					
Ambenonium Chloride					
Ambutonium Bromide					
Amcinonide					
Ametazole Hydrochloride					
Amethocaine		Non- ophthalmic use			
Amethocaine Gentisate		Non- ophthalmic use			
Amethocaine Hydrochloride		Non- ophthalmic use			
Amikacin Sulphate					
Amiloride Hydrochloride					
Aminocaproic Acid					
Aminoglutethin	mide				

		from the restri	ictions on the sale and suppose	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceution form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
Aminopterin Sodium				
Amiodarone Hydrochlorid	le			
Amiphenazo Hydrochloric				
[ <sup>F203</sup> Amisulp	ride]			
Amitriptyline	e			
Amitriptyline Embonate	2			
Amitriptyline Hydrochloric				
Amlodipine Besylate				
Ammonium Bromide				
Amodiaquine Hydrochloric				
Amorolfine Hydrochloric	le			
Amoxapine				
Amoxycillin				
Amoxycillin Sodium				
Amoxycillin Trihydrate				
Amphomycin Calcium	1			
Amphoterici	n			
Ampicillin				
Ampicillin Sodium				
Ampicillin Trihydrate				

	Exemptions from the restrictions on the sale and supply of prescription only medicines				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity	
Amsacrine					
Amygdalin					
Amyl Nitrite					
Amylocaine Hydrochloride	2	Non- ophthalmic use			
[ <sup>F199</sup> Anastrozo	ole]				
Ancrod					
Androsterone					
Angiotensin Amide					
Anistreplase					
Anterior Pituitary Extract					
Antimony Barium Tartrate					
Antimony Dimercaptosu	ccinate				
Antimony Lithium Thiomalate					
Antimony Pentasulphide					
Antimony Potassium Tartrate					
Antimony Sodium Tartrate					
Antimony Sodium Thioglycollate	2				

	Exemptions from the restrictions on the sale and supply of prescription only medicines				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceuti form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity	
Antimony Sulphate					
Antimony Trichloride					
Antimony Trioxide					
Antimony Trisulphide					
Apiol					
Apomorphine	e				
Apomorphine Hydrochlorid					
[ <sup>F200</sup> Apraclon Hydrochlorid					
Aprotinin					
Arecoline Hydrobromic	le				
Argipressin					
Aristolochia					
Aristolochia Clematitis					
Aristolochia Contorta					
Aristolochia Debelis					
Aristolochia Fang-chi					
Aristolochia Manshuriens	is				
Aristolochia Serpentaria					
Arsenic					
Arsenic Triiodide					

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity		
Arsenic Trioxide						
Arsphenami	ne					
[ <sup>F204</sup> Aspirin	[	[ s] <sup>F205</sup> (1) Non- effervescent tablets and capsules]		[ F205(1) The quantity sold or supplied in one container or package shall not exceed 100 The quantity of non- effervescent tablets, capsules or a combination of both sold or supplied to a person at any one time shall not exceed 100		
	[ <sup>F206</sup> [ <sup>F2</sup> 57(0 mg]	()] [ <sup>F207</sup> ( <b>B</b> ))]n- effervescent tablets and capsules		[ <sup>F207</sup> (2]]he quantity sold or supplied in one container or package		

	Exemptions from the restrictions on the sale and supply of prescription only medicines				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity	
				shall not exceed 32	
		[ <sup>F207</sup> (3)All preparations other than non- effervescent tablets or capsules		The quantity of non- effervescent tablets, capsules or a combination of both sold or supplied to a person at any one time shall not exceed 100]	
Astemizole		F208	F208	F208	
		F208  F208			
Atenolol		• • •			
Atracurium Besylate					
Atropine		<ul><li>(1) Internal</li><li>(a) by</li><li>inhaler</li></ul>			
		(b) otherwise	(b) 300mcg (MD)		

			ictions on the sale and supp	ply of	
Column 1 Substance	prescription Column 2 Maximum strength	n only medicine Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity	
		than by inhaler			
		million	1mg (MDD)		
		(2) External (except ophthalmic)			
Atropine	1	(1) Internal			
Methobromic	le	(a) by inhaler			
		(b) otherwise than by inhaler	(b) 400mcg (MD)		
			1.3mg (MDD)		
		(2) External (except ophthalmic)			
Atropine		Internal			
Methonitrate		(a) by inhaler			
		(b) otherwise than by inhaler	(b) 400mcg (MD)		
			1.3mg (MDD)		
Atropine Oxide		(1) Internal			
Hydrochloric	le	(a) by inhaler			
		(b) otherwise than by inhaler	(b) 360mcg (MD)		
			1.2mg (MDD) 3		
		(2) External (except ophthalmic)			

	prescription	n only medicine	ictions on the sale and supplies	
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity
Atropine Sulphate		(1) Internal		
Sulphate		(a) by inhaler		
		(b) otherwise than by inhaler	(b) 360mcg (MD)	
			1.2mg (MDD)	
		(2) External (except ophthalmic)		
Auranofin				
Azapropazon	e			
Azathioprine				
Azathioprine Sodium				
Azelaic Acid				
Azelastine Hydrochlorid	e	For nasal administratio	140mcg per nostril (MD)	Container or package
nyarochionae		For the treatment of seasonal allergic rhinitis [ <sup>F209</sup> or perennial allergic rhinitis] For use in	280mcg per nostril (MDD)	containing not more than 5,040mcg of Azelastine Hydrochloride
		adults and children not less than [ <sup>F210</sup> 5 years]		
		As a non- aerosol,		

		from the restring only medicine	ictions on the sale and supp	ly of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceuti form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity
		aqueous form		
Azidocillin Potassium				
Azithromycir	1			
Azlocillin Sodium				
Aztreonam				
Bacampicillin Hydrochlorid				
Bacitracin				
Bacitracin Methylene Disalicylate				
Bacitracin Zinc				
Baclofen				
[ <sup>F203</sup> Balsalazi Sodium]	de			
Bambuterol Hydrochlorid	e			
Barium Carbonate				
Barium Chloride				
Barium Sulphide				
Beclamide				
Beclomethas	one			
Beclomethase Dipropionate	one	For nasal administratio (non- aerosol)	100mcg per nostril (MD)	Container or package containing not more than [ <sup>F211</sup> 20,000 mcg] of

	-	from the restruction from the restruction from the from the second secon	ictions on the sale and supp es	ly of
Column 1	Column 2	Column 3	Column 4	Column 5
Substance	Maximum strength	Route of administrati use or pharmaceut		Maximum quantity
		form	200	Beclomethasone
		For the prevention	200 mcg per nostril (MDD)	Dipropionate
		and treatment of allergic rhinitis	[ <sup>F212</sup> For a maximum period of 3 months]	
		[ <sup>F213</sup> For use in persons aged 18 years and over]		
Belladonna Herb		(1) Internal	(1) 1mg of the alkaloids (MDD)	
		(2) External		
Belladonna Root		(1) Internal	(1) 1mg of the alkaloids (MDD)	
		(2) External		
Bemegride				
Bemegride Sodium				
Benapryzine Tydrochloric	le			
Bendrofluazi	de			
Benethamine Penicillin				
Benoxaprofe	n			
Benperidol				
<sup>F203</sup> Benseraz	ide]			
Benserazide Tydrochlorid	le			
Bentiromide				
Benzathine Penicillin				
Benzbromarc	one			

	Exemptions from the restrictions on the sale and supply of prescription only medicines				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity	
Benzhexol Hydrochlorid	le				
Benzilonium Bromide					
Benzocaine		Any use except ophthalmic use			
Benzoctamin Hydrochlorid					
Benzoyl Peroxide	10.0 per cent	External			
N-Benzoyl Sulphanilami	de				
Benzquinami	de				
Benzquinami Hydrochlorid					
Benzthiazide					
Benztropine Mesylate					
Benzylpenici Calcium	llin				
Benzylpenici Potassium	llin				
Benzylpenici Sodium	llin				
Beractant					
Betahistine Hydrochlorid	le				
Betamethason	ne				
Betamethason Adamantoate					
Betamethason Benzoate	ne				

		from the restric only medicines	ctions on the sale and sup	ply of	
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratic use or pharmaceutic form	Column 4 Treatment limitations on,	Column 5 Maximum quantity	
Betamethaso Dipropionate					
Betamethaso Sodium Phosphate	ne				
Betamethaso Valerate	ne				
Betaxolol Hydrochloric	le				
Bethanechol Chloride					
Bethanidine Sulphate					
Bezafibrate					
[ <sup>F200</sup> Bicalutar	nide]				
Biperiden Hydrochlorid	le				
Biperiden Lactate					
Bismuth Glycollylarsa	nilate				
Bisoprolol Fumarate					
Bleomycin					
Bleomycin Sulphate					
Bretylium Tosylate					
[ <sup>F203</sup> Brimonic Tartrate]	line				
Bromhexine Hydrochloric	le				
Bromocriptir Mesylate	ne				
Bromperidol					

		from the restruction from the restruction from the from the second secon	ictions on the sale and suppl	ly of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity
Bromvaleton	e			
Brotizolam				
Budesonide		For nasal administratio	200mcg per nostril (MD)	Container or package
		For the prevention or treatment of seasonal allergic rhinitis	[ <sup>F212</sup> For a maximum period of 3 months]	containing not more than 10mg of Budesonide
		200 mcg per nostril (MDD)		
		[ <sup>F213</sup> For use in persons aged 18 years and over]		
		As a non- aerosol, aqueous form		
Bufexamac				
Bumetanide				
Buphenine			6mg (MD)	
Hydrochlorid	le		18mg (MDD)	
Bupivacaine		Any use except ophthalmic use		
Bupivacaine Hydrochlorid	e	Any use except ophthalmic use		
Buserelin Acetate				

		from the restrict only medicine	ctions on the sale and sup s	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
Buspirone Hydrochloric	le			
Busulphan				
Butacaine Sulphate		Any use except ophthalmic use		
Butorphanol Tartrate				
Butriptyline Hydrochloric	le			
[ <sup>F214</sup> Cabergol	ine]			
Calcipotriol				
[ <sup>F200</sup> Calcipotr Hydrate]	riol			
Calcitonin				
Calcitriol				
Calcium Amphomycin	n			
Calcium Benzamidosa	alicylate			
Calcium Bromide				
Calcium Bromidolacte	obionate			
Calcium Carbimide				
Calcium Folinate				
Calcium Metrizoate				
Calcium Sulphaloxate				
[ <sup>F215</sup> Candesar Cilexetil]	rtan			

	prescription	from the restri only medicine		ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceuti form		Column 5 Maximum quantity
Candicidin				
Canrenoic Acid				
Cantharidin	0.01 per cent	External		
Capreomycir Sulphate	1			
Captopril				
Carbachol				
Carbamazepi	ine			
Carbaryl				
[ <sup>F203</sup> Carbasal Calcium]	ate			
Carbenicillin Sodium	L			
Carbenoxolo Sodium	ne	(1) Pellet	(1) 5mg (MD) 25mg (MDD)	
	(2) 2.0 per cent	(2) Gel		
	(3) 1.0 per cent	(3) Granules for mouthwash in adults and children not less than 12 years	(3) 20mg (MD) 80mg (MDD)	(3) Container or package containing not more than [ <sup>F216</sup> 560mg] of Carbenoxolone Sodium
Carbidopa				
Carbimazole				
Carbocistein	e			
Carbon Tetrachloride	e			
Carboplatin				

58

<i>Exemptions from the restrictions on the sale and supply of prescription only medicines</i>			pply of	
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
Carboprost Trometamol				
Carbuterol Hydrochloride	5			
Carfecillin Sodium				
Carindacillin Sodium				
Carisoprodol				
Carmustine				
Carperidine				
Carteolol Hydrochloride	5			
Cefaclor				
Cefadroxil				
Cefazedone Sodium				
[ <sup>F203</sup> Cefdinir]				
Cefixime				
Cefodizime Sodium				
Cefotaxime Sodium				
Cefoxitin Sodium				
Cefpodoxime Proxetil				
[ <sup>F214</sup> Cefprozil]	l			
Cefsulodin Sodium				
Ceftazidime				
Ceftizoxime Sodium				

		from the restruction only medicine	ictions on the sale and sup	ply of	
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity	
Ceftriaxone Sodium					
Cefuroxime Axetil					
Cefuroxime Sodium					
Celiprolol Hydrochlorid	e				
Cephalexin					
Cephalexin Sodium					
Cephaloridine	2				
Cephalothin Sodium					
Cephamandol Nafate	e				
Cephazolin Sodium					
Cephradine					
Cerium Oxalate					
Cerivastatin					
[ <sup>F203</sup> Cerivasta Sodium]	tin				
Ceruletide Diethylamine					
Cetirizine Hydrochlorid	e		10mg (MDD)	F217	
Chenodeoxyc Acid	holic				
Chloral Hydrate		External			
Chlorambucil					
Chloramphen	icol				

		from the restruction only medicine	ictions on the sale and sup	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity
Chloramphen Cinnamate	icol			
Chloramphen Palmitate	icol			
Chloramphen Sodium Succinate	icol			
Chlorhexadol				
Chlormadinoi Acetate	ne			
Chlormerodri	n			
Chlormethiaz	ole			
Chlormethiaz Edisylate	ole			
Chlormezano	ne			
Chloroform(1	(4)) 5.0 per cent	(1) Internal		
		(2) External		
Chloroquine Phosphate		Prophylaxis of malaria		
Chloroquine Sulphate		Prophylaxis of malaria		
Chlorothiazid	le			
Chlorotrianise	ene			
Chlorphenoxa Hydrochlorid				
Chlorpromazi	ine			
Chlorpromazi Embonate	ine			
Chlorpromazi Hydrochlorid				
Chlorpropam	ide			

<sup>(14)</sup> SeeS.I. 1979/382 amended by S.I. 1980/263 and S.I. 1989/1124.

		from the restring from the restrict from the res	ctions on the sale and suppl	y of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceute form		Column 5 Maximum quantity
Chlorprothix	ene			
Chlorprothix Hydrochlorid				
Chlortetracy	cline			
Chlortetracy Calcium	cline			
Chlortetracy Hydrochlorid				
Chlorthalido	ne			
Chlorzoxazo	ne			
Cholestyram	ine			
Ciclacillin				
Ciclobendaz	ole			
Cilastatin Sodium				
Cilazapril				
Cimetidine		(a) For the	(a) 200mg (MD)	
		short-term symptomatic	800mg (MDD)	
		relief of heartburn, dyspepsia, indigestion, acid indigestion and hyperacidity and for the prophylaxis of meal- induced	For a maximum period of 14 days	
		heartburn	$(1) 100 \dots (100) + 1$	
		management	(b) 100mg (MD) to be taken as a single dose at night	
		of nocturnal heartburn by a single	For a maximum period of 14 days	
			62	

		from the restri only medicine	ctions on the sale and sup	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceution form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
		dose taken at night		
Cimetidine Hydrochlorid	le			
Cinchocaine	3.0 per cent	Non- ophthalmic use		
Cinchocaine Hydrochlorid		Non- ophthalmic use		
Cinchophen				
Cinoxacin				
Ciprofibrate				
Ciprofloxaci	n			
Ciprofloxaci Hydrochlorid				
Cisapride				
Cisplatin				
[ <sup>F200</sup> Citalopra Hydrobromio				
Clarithromy	cin			
Clavulanic Acid				
Clidinium Bromide				
Clindamycin				
Clindamycin Hydrochlorid				
Clindamycin Palmitate Hydrochlorid				
Clindamycin Phosphate				

		from the restri only medicine	ctions on the sale and sup	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceution form		Column 5 Maximum quantity
Clioquinol		(1) External (other than treatment of mouth ulcers)		
	(2) 35mg	(2) Treatment of mouth ulcers	(2) 350mg (MDD)	
Clobetasol Propionate				
Clobetasone Butyrate	[ <sup>F218</sup> 0.05 per cent]	[ <sup>F218</sup> Cream for use in adults and in children aged 12 years and over, for external use for the short term symptomatic treatment and control of patches of eczema and dermatitis (excluding seborrhoeic dermatitis)]		[ <sup>F218</sup> Container or package containing not more than 15g of medicinal product]
Clofazimine				
Clofibrate				
Clomiphene Citrate				
Clomipramin	e			
Clomipramin Hydrochlorid				
Clomocycline	e			

	Exemptions from the restrictions on the sale and supply of prescription only medicines						
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceution form	Column 4 Treatment limitations on,	Column 5 Maximum quantity			
Clomocycline Sodium	2						
Clonidine							
Clonidine Hydrochlorid	e						
Clopamide							
Clopenthixol Decanoate							
Clopenthixol Hydrochlorid	e						
Clorexolone							
Clotrimazole		External but in the case of vaginal use only external use for the treatment of vaginal candidiasis					
Cloxacillin Benzathine							
Cloxacillin Sodium							
Clozapine							
Cocculus Indicus							
Co- dergocrine Mesylate							
Colaspase							
Colchicine							
Colestipol Hydrochlorid	e						
Colfosceril Palmitate							

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceution form	Column 4 Treatment limitations on,	Column 5 Maximum quantity		
Colistin Sulphate						
Colistin Sulphometha	ite					
Colistin Sulphometha Sodium	ıte					
Coniine						
Conium Leaf	7.0 per cent	External				
Corticotroph	in					
Cortisone						
Cortisone Acetate						
Co- tetroxazine						
Co- trimoxazole						
Cropropamic	le					
Crotethamide	e					
Croton Oil						
Croton Seed						
Curare						
Cyclofenil						
Cyclopenthia	zide					
Cyclopentola Hydrochloric						
Cyclophosph	amide					
Cycloserine						
Cyclosporin						
Cyclothiazid	e					
Cyproterone Acetate						

		Exemptions from the restrictions on the sale and supply of prescription only medicines			
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceutio form	Column 4 Treatment limitations on,	Column 5 Maximum quantity	
Cytarabine					
Cytarabine Hydrochlorid	e				
Dacarbazine					
Dalteparin Sodium					
Danazol					
Danthron					
Dantrolene Sodium					
Dapsone					
Dapsone Ethane Ortho Sulphonate					
Daunorubicir Hydrochlorid					
Deanol Bitartrate			26mg (MDD)		
Debrisoquine Sulphate					
Demecarium Bromide					
Demeclocycl	ine				
Demeclocycl Calcium	ine				
Demeclocycl Hydrochlorid					
Deoxycortone Acetate	e				
Deoxycorton Pivalate	e				
Deptropine Citrate					

		from the restrict only medicines	ions on the sale and sup	pply of
Column 1 Substance	Column 2 Maximum strength	Column 3		Column 5 Maximum quantity
Dequalinium Chloride	n (1) 0.25mg	(1) Internal: throat lozenges or throat pastilles		
	(2) 1.0 per cent	(2) External: paint		
Deserpidine				
Desferrioxan Mesylate	nine			
Desflurane				
Desipramine Hydrochlorid				
Deslanoside				
Desmopressi	n			
Desmopressi Acetate				
Desogestrel				
Desonide				
Desoxymeth	asone			
Dexamethas				
Dexamethase Acetate	one			
Dexamethase Isonicotinate				
Dexamethase Phenylpropie				
Dexamethase Pivalate	one			
Dexamethase Sodium Metasulphob				
Dexamethase Sodium Phosphate				

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form		Column 5 Maximum quantity		
Dexamethase Troxundate	one					
Dexfenflurar Hydrochloric						
Dextromethorphan Hydrobromide		Internal	<ul><li>(a) In the case</li><li>of a prolonged</li><li>release preparation:</li><li>equivalent of 30mg of</li><li>Dextromethorphan (MD)</li></ul>			
			equivalent of 75mg of Dextromethorphan (MDD)			
			(b) in any other case: equivalent of 15mg of Dextromethorphan (MD)			
			equivalent of 75mg of Dextromethorphan (MDD)			
Dextrothyrox Sodium	kine					
Diazoxide						
Dibenzepin Hydrochloric	le					
Dichloralphe	nazone					
Dichlorphen	amide					
Diclofenac	1.16 per	External	For maximum period of 7	Container		
Diethylamm	Di <b>urant</b> i	For local symptomatic relief of pain and inflammation in trauma of the tendons, ligaments, muscles and joints and in localised	days	or package containing not more than 30g of medicinal product		

69

		from the restriction only medicine	ctions on the sale and sup	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
		soft tissue rheumatism		
		For use in adults and children not less than 12 years		
Diclofenac Potassium				
Diclofenac Sodium				
Dicyclomine			10mg (MD)	
Hydrochloride			60mg (MDD)	
[ <sup>F199</sup> Didanosin	e]			
Dienoestrol				
Diethanolamin Fusidate	ne			
Diflucortolone Valerate	e			
Diflunisal				
Digitalin				
Digitalis Leaf				
Digitalis Prepared				
Digitoxin				
Digoxin				
Dihydralazine Sulphate				
Dihydroergota Mesylate	mine			
Dihydrostrept	omycin			
Dihydrostrept Sulphate	omycin			

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceutit form	,	Column 5 Maximum quantity		
Diloxanide Furoate						
Diltiazem Hydrochloric	le					
Dimercaprol						
Dimethisoqu Hydrochloric		Non- ophthalmic use				
Dimethistero	ne					
Dimethothiaz Mesylate	zine					
Dimethyl Sulphoxide						
Dimethyltub Bromide	ocurarine					
Dimethyltub Chloride	ocurarine					
Dimethyltub Iodide	ocurarine					
Dinoprost						
Dinoprost Trometamol						
Dinoprostone	e					
[ <sup>F202</sup> Diphenh <u>y</u> Hydrochlorid	ydialmine lepreparations except liquid-filled capsules]					
[ <sup>F219</sup> Dipheno: Hydrochlorid	x∳fål2.5 mg] le]	[ <sup>F219</sup> In combination with Atropine Sulphate for short term use as an adjunctive therapy to	[ <sup>F219</sup> 25 mg (MDD)]	[ <sup>F219</sup> Container or package containing not more than 20 tablets]		

		xemptions from the restrictions on the sale and supply of rescription only medicines				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity		
		appropriate rehydration in acute diarrhoea		_		
		For use in persons aged 16 years and over				
		Tablets]		_		
Dipivefrin Hydrochlorid	e					
Dipyridamole	e					
Disodium Etidronate						
Disodium Pamidronate						
Disopyramid	e					
Disopyramide Phosphate	5					
Distigmine Bromide						
Disulfiram						
Dithranol	1.0 per cent					
Dobutamine Hydrochlorid	e					
[ <sup>F219</sup> Dolasetro Mesilate]	n					
Domperidone	;	[ <sup>F220</sup> For the relief of post- prandial symptoms of excessive fullness, nausea, epigastric	[ <sup>F220</sup> 10mg of Domperidone (MD)] [ <sup>F220</sup> 40mg of Domperidone (MDD)]	[ <sup>F220</sup> Container or package containing not more than 200mg of Domperidone]		

		from the restri only medicine	ctions on the sale and sup	ply of
Column 1 Substance	prescription Column 2 Maximum strength	Column 3 Route of administration	Column 4 Treatment limitations	Column 5 Maximum quantity
		use or pharmaceuti	ical	
		form	cui	
		bloating and belching, occasionally accompanied by epigastric discomfort and heartburn]		
Domperidone Maleate		[ <sup>F221</sup> For the relief of post- prandial symptoms of excessive fullness, nausea, epigastric bloating and belching, occasionally accompanied by epigastric discomfort and heartburn,]	[ <sup>F222</sup> 10 mg of Domperidone as Domperidone Maleate (MD)] [ <sup>F222</sup> 40 mg of Domperidone as Domperidone Maleate (MDD)]	[ <sup>F221</sup> Container or package containing not more than [ <sup>F223</sup> 200mg] of Domperidone as Domperidone Maleate;]
[ <sup>F203</sup> Donepezil Hydrochloride	]			
Dopamine Hydrochloride				
Dopexamine Hydrochloride				
[ <sup>F200</sup> Dorzolami Hydrochloride				
Dothiepin				
Dothiepin Hydrochloride				
Doxapram Hydrochloride				
			73	

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity		
Doxazosin Mesylate						
Doxepin Hydrochlorid	e					
Doxorubicin						
Doxorubicin Hydrochlorid	e					
Doxycycline						
Doxycycline Calcium Chelate						
Doxycycline Hydrochlorid	e					
Droperidol						
Dydrogestero	ne					
Dyflos						
Econazole		External but in the case of vaginal use only external use for the treatment of vaginal candidiasis				
Econazole Nitrate		External but in the case of vaginal use only external use for the treatment of vaginal candidiasis				
Ecothiopate Iodide						
Edrophonium Chloride						

		from the restruction only medicine	ictions on the sale and supp	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity
Eflornithine Hydrochlorid	le			
[ <sup>F199</sup> Eformote Fumarate]	erol			
Embutramide	e			
Emepronium Bromide				
Emetine	1.0 per cent			
Emetine Bismuth Iodide				
Emetine Hydrochloric	Equivalent leof 1.0 per cent of Emetine			
Enalapril Maleate				
Encephalitis Virus, Tick- borne, Cent Eur				
Enoxacin				
Enoxaparin Sodium				
Enoximone				
Ephedrine		(1) Internal (other than nasal sprays or nasal drops)	(1) 30mg (MD)	
			60mg (MDD)	
	(2) 2.0 per cent	<ul><li>(2) Nasal sprays or nasal drops</li><li>(3) External</li></ul>		

		from the restri only medicine	ictions on the sale and supply	v of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity
Ephedrine Hydrochlori	de	(1) Internal (other than	(1) Equivalent of 30mg of Ephedrine (MD)	
		nasal sprays or nasal drops)	Equivalent of 60mg of Ephedrine (MDD)	
	(2) Equivalent of 2.0 per cent of Ephedrine	(2) Nasal sprays or nasal drops		
		(3) External		
Ephedrine Sulphate		(1) Internal (other than nasal sprays or nasal drops)	(1) Equivalent of 30mg of Ephedrine (MD)	
			Equivalent of 60mg of Ephedrine (MDD)	
	(2) Equivalent of 2.0 per cent of Ephedrine	(2) Nasal sprays or nasal drops		
		(3) External		
Epicillin				
Epirubicin				
Epirubicin Hydrochlori	de			
Epithiazide				
Epoetin Alfa				
Epoetin Beta				
Epoprosteno Sodium	1			
Ergometrine Maleate				

		from the restriction only medicine		ons on the sale and supply of		
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity		
Ergometrine Tartrate						
Ergot, Prepared						
Ergotamine Tartrate						
Erythromycir	1					
Erythromycir Estolate	1					
Erythromycir Ethylcarbona						
Erythromycir Ethyl Succinate	1					
Erythromycir Lactobionate	1					
Erythromycir Phosphate	1					
Erythromycir Stearate	1					
Erythromycir Thiocyanate	1					
Esmolol Hydrochlorid	e					
Estramustine Phosphate						
[ <sup>F224</sup> Estramus Sodium Phosphate]	tine					
Etafedrine Hydrochlorid	e					
Ethacrynic Acid						
Ethambutol Hydrochlorid	e					

Column 1 Substance		from the restri only medicine Column 3 Route of administratio use or pharmaceuti	Column 4 Treatment limitations on,	oly of Column 5 Maximum quantity
Ethamivan		form		
Ethamsylate Ethiazide				
Ethinyl Androstened	iol			
Ethinyloestra				
Ethionamide				
Ethisterone				
Ethoglucid				
Ethoheptazin Citrate	ie			
Ethopropazir Hydrochloric				
Ethosuximid	e			
Ethotoin				
Ethyl Biscoumacet	ate			
Ethynodiol Diacetate				
Etodolac				
Etomidate				
Etomidate Hydrochloric	le			
Etoposide				
Etretinate				
[ <sup>F200</sup> Exemesta	ane]			
Famciclovir				
Famotidine		For the	10mg (MD)	
		short-term symptomatic	20mg (MDD)	
		relief of heartburn, dyspepsia, indigestion,	For maximum period of 14 days	
		margestion,	78	

		from the restring only medicine	ctions on the sale and suppl	ly of
Column 1	Column 2	Column 3	Column 4	Column 5
Substance	Maximum	Route of	Treatment limitations	Maximum
	strength	administrati	on,	quantity
	U	use or		1 2
		pharmaceut	ical	
		form		
		acid		
		indigestion		
		and		
		hyperacidity,		
		and		
		prevention		
		of these		
		symptoms		
		when associated		
		with food		
		and drink,		
		including		
		nocturnal		
		symptoms		
т I''		~)p		
Fazadinium Bromide				
Felbinac	3.17 per	External	For maximum period of 7	Container
	cent	[ <sup>F226</sup> For the	days	or package
		relief of		containing
		rheumatic		not more than
		pain, pain of		
		non-serious		[ <sup>F225</sup> 50g] of medicinal
		arthritic		product
		conditions		product
		and soft		
		tissue		
		injuries such		
		as sprains,		
		strains and		
		contusions]		
		For use in		
		adults and		
		children not		
		less than 12		
		years		
Felodipine				
Felypressin				
Fenbufen				
Fenclofenac				
Fenclofenac				

		from the restr	ictions on the sale and suppers	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity
Fenfluramine Hydrochlorie				
Fenofibrate				
Fenoprofen				
Fenoprofen Calcium				
Fenoterol Hydrobromi	de			
Fenticonazol Nitrate	le	[ <sup>F218</sup> External use (but in the case of vaginal use, only for the treatment of vaginal candidiasis)]		
Feprazone				
Ferrous Arsenate				
[ <sup>F200</sup> Ferumox	sil]			
[ <sup>F203</sup> Fexofena Hydrochlorid				
Filgrastim				
Finasteride				
Flavoxate Hydrochlorid	de			
Flecainide Acetate				
Flosequinan				
Fluanisone				
Flubendazol	e			
Fluclorolone Acetonide	;			
Flucloxacilli Magnesium	n			

	prescription	from the restr n only medicin	ictions on the sale and supper	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrat use or pharmaceut form		Column 5 Maximum quantity
Flucloxacilli Sodium	n			
Fluconazole		For oral administration for the treatment of vaginal candidiasis [ <sup>F227</sup> or associated candidal balanitis] in persons aged not less than 16 but less than 60 years		Container or package containing not more than 150mg of Fluconazole
Flucytosine				
Fludrocortise Acetate	one			
Flufenamic Acid				
Flumazenil				
Flumethasor	ie			
Flumethasor Pivalate	ie			
Flunisolide	0.025 per cent	(a) For the prevention and treatment of seasonal allergic rhinitis, including hay fever	<ul><li>(a) 50mcg per nostril</li><li>(MD)</li><li>100mcg per nostril</li><li>(MDD)</li></ul>	(a) Container or package containing not more than 6,000mcg of Flunisolide
		[ <sup>F228</sup> For use in persons aged 18	[ <sup>F229</sup> For a maximum period of 3 months]	

		from the restr only medicin	rictions on the sale and sup es	ply of	
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrat use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity	
		years and over]			
		In the form of a non- pressurised nasal spray			
		F230	F230	F230	
			F230		
		F230			
		F230			
Fluocinolone Acetonide					
Iuocinonide					
Fluocortin Butyl					
Fluocortolone	e				
Fluocortolone Hexanoate	2				
Fluocortolone Pivalate	2				
Fluorescein Dilaurate					
Fluoromethol	one				
Fluorouracil					
Fluorouracil Frometamol					
Fluoxetine Hydrochlorid	e				
Flupenthixol Decanoate					

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity		
Flupenthixol Hydrochloride	e					
Fluperolone Acetate						
Fluphenazine Decanoate						
Fluphenazine Enanthate						
Fluphenazine Hydrochloride	e					
Fluprednidene Acetate	2					
Fluprednisolo	ne					
Fluprostenol Sodium						
Flurandrenolo	one					
Flurbiprofen		[ <sup>F232</sup> Throat lozenges]	[ <sup>F233</sup> 43.75 mg (MDD)]	[ <sup>F234</sup> Container or package containing not more than 140 mg of Flurbiprofen]		
Flurbiprofen Sodium						
Fluspirilene						
Flutamide						
Fluticasone Propionate						
[ <sup>F203</sup> Flutrimaz	ole]					
Fluvastatin Sodium						
Fluvoxamine Maleate						

	<i>Exemptions from the restrictions on the sale and supply of prescription only medicines</i>				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form		Column 5 Maximum quantity	
Formestane					
Formocortal					
Foscarnet Sodium					
Fosfestrol Sodium					
Fosfomycin Trometamol					
Fosinopril Sodium					
Framycetin Sulphate					
Frusemide					
Furazolidone					
Fusafungine					
Fusidic Acid					
Gabapentin					
Gadoteridol					
Gallamine Triethiodide					
Ganciclovir					
Ganciclovir Sodium					
Gelsemine	0.1 per cent				
Gelsemium			25mg (MD)		
			75mg (MDD)		
Gemeprost					
Gemfibrozil					
Gentamicin					
Gentamicin Sulphate					
Gestodene					

		from the restriction only medicine.	ctions on the sale and sup	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
Gestrinone				
Gestronol				
Gestronol Hexanoate				
Glibenclamic	le			
Glibornuride				
Gliclazide				
Glipizide				
Gliquidone				
Glisoxepide				
Glucagon				
Glycopyrron Bromide	ium		1mg (MD) 2mg (MDD)	
Glymidine			2g (10.00)	
Gonadorelin				
Goserelin Acetate				
Gramicidin	0.2 per cent	External		
Granisetron Hydrochlorid	le			
Griseofulvin				
Growth Hormone				
Guanethidine Monosulphat				
Guanfacine Hydrochlorid	le			
Guanoclor Sulphate				
Guanoxan Sulphate				
Halcinonide				

		from the restruction only medicine	ictions on the sale and supp	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity
Halofantrine Hydrochlorid				
Haloperidol				
Haloperidol Decanoate				
Heparin		External		
Heparin Calcium		External		
Heparin Sodium				
Hexachlorop	hane	External		
	(a) 2.0 per cent	(a) Soaps		
	(b) 0.1 per cent	(b) Aerosols		
	(c) 0.75 per cent	(c) preparations other than soaps and aerosols		
Hexamine Phenylcinch	oninate			
Hexobarbito	ne			
Hexobarbito Sodium	ne			
Hexoestrol				
Hexoestrol Dipropionate	e			
L-Histidine Hydrochlorid	de	Dietary supplementa	tion	
Homatropine	e	(1) Internal	(1) 0.15mg (MD)	
			0.45mg (MDD)	
		(2) External (except ophthalmic)		

		from the restri only medicine	ctions on the sale and sup s	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceution form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
Homatropine		0	0.2mg (MD)	
Hydrobromide			0.6mg (MDD)	
Homatropine Methylbromid	e		2mg (MD) 6mg (MDD)	
Hydralazine Hydrochloride				
Hydrargaphen		Local application to skin		
Hydrobromic Acid				
Hydrochloroth	iazide			
Hydrocortison	per cent]	<ul> <li>[<sup>F235</sup>(1) Ex</li> <li>(a) For use in combin with Nystatin of maximus strength 3.0 per cent for intertrig</li> <li>(b) For use in adults and children not less than 10 years]</li> </ul>	ation n um n	(Containing or package containing not more than 15g of medicinal product]
]	[ <sup>F236</sup> (2)].0 per cent	[ <sup>F236</sup> (2)] E (a) For use either alone	xternal 87	(Con(a)))er or package containing not more than 15g

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Column 4 Route of Treatment lim administration, use or	Column 5 itations Maximum quantity			
		pharmaceutical				
		form				
		or in	of medicinal			
		conjunction with	product			
		Crotamiton	(cream or ointment) or			
		in	30ml (spray)			
		irritant				
		dermatitis,				
		contact				
		allergic				
		dermatitis,				
		insect bite				
		reactions,				
		mild				
		to				
		moderate				
		eczema,				
		and				
		either				
		in combination				
		with				
		Clotrimazole				
		[ <sup>F237</sup> or				
		Miconazole				
		Nitrate]				
		for				
		athlete's				
		foot				
		and candidal				
		intertrigo				
		or in				
		combination				
		with				
		lignocaine				
		for				
		anal				
		and				
		perianal itch				
		associated				
		with				
		haemorrhoids				

			ctions on the sale and sup	pply of
Column		only medicines		Column 5
Column 1	Column 2	Column 3	Column 4	Column 5
Substance	Maximum	Route of	Treatment limitations	Maximum
	strength	administratio	on,	quantity
		use or		
		pharmaceuti	cal	
		form		
		(b) For		
		use in		
		adults		
		and		
		children	1	
		not		
		less		
		than		
		10		
		years		
		(c) Cream		
		ointmen	nt	
		or		
		spray		
Hydrocortis	onEquivalent	External		
Acetate	to 1.0			
	per cent	For use		Container
	Hydrocortiso	in irritant		or package
	)	dermatitis,		containing
		contact		not more
		allergic		than 15g of
		dermatitis,		medicinal
		insect bite		product
		reactions,		In the
		mild to		case of
		moderate		suppositories,
		eczema,		container
		and in		or package
		combination		containing
		with one or		no more
		more of the		than 12
		following:		
		Benzyl		
		Benzoate,		
		Bismuth		
		Oxide,		
		Bismuth		
		Subgallate,		
		Peru		
		Balsam,		
		Pramoxine		
		Hydrochlorid	e,	
		Zinc		
		Oxide, for		
		haemorrhoids	5	
			89	

		from the restric only medicines	ctions on the sale and s	supply of	
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceutio form		Column 5 Maximum quantity	
		[ <sup>F238</sup> or in combination with Miconazole Nitrate, for athlete's foot and candidal intertrigo]			
		For use in adults and children not less than 10 years			
Hydrocortiso	ne	Cream, ointment or suppositories			
Butyrate Hydrocortisor Caprylate					
Hydrocortison Hydrogen Succinate	ne				
Hydrocortiso Sodium Phosphate	ne				
Hydrocortisor Sodium Succinate	nEquivalent to 2.5mg Hydrocortison	External For aphthous ulceration of the mouth for adults and children not less than 12 years		Container or package containing not more than equivalent to 50mg of Hydrocortisone	
		In the form of pellets			

	prescription	n only medicine.	ctions on the sale and sup s	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form		Column 5 Maximum quantity
[ <sup>F202</sup> Hydrocy Acid]	anic			
Hydroflumet	hiazide			
Hydroxychlo Sulphate	oroquine	Prophylaxis of malaria		
Hydroxyprog	gesterone			
Hydroxyprog Enanthate	gesterone			
Hydroxyprog Hexanoate	gesterone			
Hydroxyurea	ı			
Hydroxyzine Embonate				
Hydroxyzine Hydrochlorid		(a) For the management of pruritis associated with acute or chronic urticaria or atopic dermatitis or contact dermatitis, in adults and in children not less than 12 years	(a) 25mg (MD) 75mg (MDD)	(a) Container or package containing not more than 750mg of Hydroxyzine Hydrochloride
		(b) For the management of pruritis associated with acute or chronic urticaria or atopic dermatitis or contact dermatitis,	(b) 25 mg (MD) 50mg (MDD)	(b) Container or package containing not more than 750mg of Hydroxyzine Hydrochloride

		from the restri only medicine	ctions on the sale and support	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceuth form in children not less than 6 years but less than 12		Column 5 Maximum quantity
Hyoscine	(1) 0.15 per	years (1) Internal		
	cent	(2) External (except ophthalmic)		
Hyoscine		(1) Internal		
Butylbromid				
		(b) otherwise than by inhaler	(b) 20mg (MD) 80mg (MDD)	(b) Container or package containing not more than 240mg of Hyoscine Butylbromide
		(2) External		
Hyoscine		(1) Internal		
Hydrobromic	de	(a) by inhaler		
		(b) otherwise than by inhaler	(b) 300mcg (MD) 900mcg (MDD)	
		(2) External (except ophthalmic)		
Hyoscine		(1) Internal		
Methobromie	de	(a) by inhaler		
		(b) otherwise	(b) 2.5mg (MD) 7.5mg (MDD)	

		from the restruction only medicine	ictions on the sale and supples	y of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form than by	,	Column 5 Maximum quantity
		inhaler		
Hyoscine		<ul><li>(2) External</li><li>(1) Internal</li></ul>		
Methonitrate		(a) by inhaler		
		(b) otherwise than by inhaler	(b) 2.5mg (MD) 7.5mg (MDD)	
		(2) External		
Hyoscyamine	;	(1) Internal		
		(a) by inhaler		
		(b) otherwise than by inhaler	(b) 300mcg (MD) 1mg (MDD)	
		(2) External		
Hyoscyamine		(1) Internal		
Hydrobromid	e	(a) by inhaler		
		(b) otherwise than by inhaler	<ul><li>(b) Equivalent of 300mcg of Hyoscyamine (MD)</li><li>Equivalent of 1mg of Hyoscyamine (MDD)</li></ul>	
		(2) External		
Hyoscyamine Sulphate	;	(1) Internal		
Sulphate		(a) by inhaler		
		(b) otherwise than by inhaler	<ul><li>(b) Equivalent of 300mcg of Hyoscyamine (MD)</li><li>Equivalent of 1mg of Hyoscyamine (MDD)</li></ul>	
		(2) External		

		from the restri only medicine	ctions on the sale and supply	y of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceution form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
Ibuprofen		Rheumatic and muscular pain, pain of non-serious arthritic conditions, backache, neuralgia, migraine, headache, dental pain, dysmenorrho feverishness, symptoms of colds and influenza	ea,	
		(1) Internal	(1)(a) In the case of a prolonged release preparation 600mg (MD)	
			1,200mg (MDD)	
			(b) in any other case 400mg (MD)	
			1,200mg (MDD)	
	(2) 5.0 per cent	(2) External		
	[ <sup>F239</sup> (3) 10.0 per cent]	[ <sup>F239</sup> (3) External]	[ <sup>F239</sup> (3) 125 mg (MD) 500 mg (MDD)]	[ <sup>F239</sup> (3) Container or package containing not more than [ <sup>F240</sup> 50g] of medicinal product]
[ <sup>F202</sup> Ibuprofer Lysine	1	Rheumatic and muscular pain, pain of non-serious arthritic	(a) in the case of a prolonged release preparation 600 mg (MD) 1,200 mg (MDD)	

	prescription	from the restr only medicine	ictions on the sale and supp es	ly of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati	Column 4 Treatment limitations ion.	Column 5 Maximum quantity
	strength	use or		quantity
		pharmaceut	ical	
		<i>form</i> conditions,		
		backache,		
		neuralgia, migraine,		
		headache,		
		dental pain,		
		dysmenorrho feverishness		
		symptoms	,	
		of colds and influenza		
		Internal	(b) in any other case 400 mg (MD) 1,200 mg (MDD)]	
Idarubicin Hydrochloride	5			
Idoxuridine				
Ifosfamide				
Ignatius Bean				
[ <sup>F199</sup> Imidapril Hydrochloride	e]			
Imipenem Hydrochloride	2			
Imipramine				
Imipramine Hydrochloride	e			
Imipramine Ion Exchange				
Resin Bound Salt or Complex				
[ <sup>F214</sup> Indapamic	de]			
Indapamide Hemihydrate				
Indomethacin				

	prescription	from the restrict only medicine	ictions on the sale and suppers	ply of	
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity	
Indomethacir Sodium	1				
Indoprofen					
Indoramin Hydrochlorid	e				
Inosine Pranobex					
[ <sup>F241</sup> Insulin]					
Iodamide					
Iodamide Meglumine					
Iodamide Sodium					
Iohexol					
Iomeprol					
Iopamidol					
Iopentol					
Iothalamic Acid					
Ioversol					
Ioxaglic Acid					
Ipratropium Bromide					
Iprindole Hydrochlorid	e				
Iproniazid Phosphate					
[ <sup>F203</sup> Irbesartar	1]				
Isoaminile					
Isoaminile Citrate					
Isocarboxazio	1				

		from the restring from the restrict from the res	ictions on the sale and suppl	y of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity
Isoconazole Nitrate		External but in the case of vaginal use only external use for the treatment of vaginal candidiasis		
Isoetharine				
Isoetharine Hydrochlorid	e			
Isoetharine Mesylate				
Isoniazid				
Isoprenaline Hydrochlorid	е			
Isoprenaline Sulphate				
Isopropamide Iodide			Equivalent of 2.5mg of Isopropamide ion (MD)	
			Equivalent of 5.0mg of Isopropamide ion (MDD)	
Isotretinoin				
Isradipine				
Itraconazole				
Jaborandi		External		
Kanamycin Acid Sulphate				
Kanamycin Sulphate				
Ketamine Hydrochlorid	e			

		from the restri only medicine	ctions on the sale and suppl	y of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati- use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
Ketoconazol	e 2.0 per cent	[ <sup>F242</sup> (a)][ <sup>F243</sup> E For the prevention and treatment of dandruff and seborrhoeic dermatitis of the scalp In the form of a shampoo	xtEth(a)] Maximum frequency of application of once every 3 days	[ <sup>F242</sup> (a)] Container or package containing not more than 120ml of medicinal product and containing not more than 2,400mg of Ketoconazole
		[ <sup>F244</sup> (b) For the treatment of the following mycotic infections of the skin: tinea pedis, tinea cruris and candidal intertrigo]		
Ketoprofen	2.5 per cent	External For rheumatic and muscular pain in adults and children not less than 12	For maximum period of 7 days	Container or package containing not more than 30g of medicinal product
Ketorolac Trometamol Ketotifen Fumarate				
Labetalol Hydrocholor	ide			

	<i>Exemptions from the restrictions on the sale and supply of prescription only medicines</i>					
Column 1 Column 1 Column 1	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceutic form	Column 4 Treatment limitations n,	Column 5 Maximum quantity		
Lachesine Chloride						
Lacidipine						
Lamotrigine						
Lanatoside C						
Lanatoside Complex A, B and C						
[ <sup>F214</sup> Lansoprazo	le]					
Latamoxef Disodium						
[ <sup>F214</sup> Lercanidipi Hydrochloride]						
Levallorphan Tartrate						
Levobunolol Hydrochloride						
-		(1) Nasal sprays For the symptomatic treatment of seasonal allergic rhinitis		(1) Container or package containing not more than 10 ml of medicinal product		
		(2) Aqueous eye drops		(2) Container		
		For the symptomatic treatment of seasonal allergic conjunctivitis		or package containing not more than 4 ml of medicinal product]		
[ <sup>F245</sup> Levocarniti	ne]	[ <sup>F245</sup> For dietary supplementati	on]			

		from the restriction only medicine	ictions on the sale and supp es	bly of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity
Levodopa				
[ <sup>F203</sup> Levoflox	acin]			
Levonorgest	re <mark>[<sup>F246</sup>0.75mg]</mark>	[ <sup>F246</sup> for use as an emergency contraceptive in women aged 16 years and over]	2	
Lidoflazine				
Lignocaine		Non- ophthalmic use		
Lignocaine Hydrochlorid	le	Non- ophthalmic use		
Lincomycin				
Lincomycin Hydrochlorid	le			
Liothyronine Sodium	;			
Lisinopril				
Lithium Carbonate			Equivalent of 5mg of Lithium (MD)	
			Equivalent of 15mg of Lithium (MDD)	
Lithium Citrate				
Lithium Succinate				
Lithium Sulphate			Equivalent of 5mg of Lithium (MD)	
			Equivalent of 15mg of Lithium (MDD)	
Lobeline		(1) Internal	(1) 3mg (MD)	
			100	

			ctions on the sale and suppl	y of
Column 1 Substance	Column 2 Maximum	only medicine Column 3 Route of	Column 4 Treatment limitations	Column 5 Maximum
	strength	administrati use or pharmaceuti		quantity
		form	Oma (MDD)	
		(2) External	9mg (MDD)	
Lobeline Hydrochlorid	le	(1) Internal	(1) Equivalent of 3mg of Lobeline (MD)	
Try aroonion			Equivalent of 9mg of Lobeline (MDD)	
		(2) External		
Lobeline Sulphate		(1) Internal	(1) Equivalent of 3mg of Lobeline (MD)	
			Equivalent of 9mg of Lobeline (MDD)	
		(2) External		
Lodoxamide Trometamol	[ <sup>F247</sup> equivaler of 0.1 per cent Lodoxamide	treatment of ocular	S,	
Lofepramine				
Lofepramine Hydrochlorid				
Lofexidine Hydrochlorid	le			
Lomefloxaci Hydrochlorid				
Lomustine				
Loperamide Hydrochlorid	le	Treatment of acute diarrhoea		
Loratidine			10mg (MDD)	F249

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#### Status: Point in time view as at 01/08/2012. Changes to legislation: There are currently no known outstanding effects for the The Prescription Only Medicines (Human Use) Order 1997. (See end of Document for details)

	1	from the restr	ictions on the sale and supp es	ply of
Column 1	Column 2	Column 3	Column 4	Column 5
Substance	Maximum strength	Route of administrat use or pharmaceut form		Maximum quantity

[<sup>F215</sup>Lornoxicam]

[ <sup>F215</sup> Losartan Potassium]
Loxapine Succinate
Lung Surfactant Porcine
Luteinising Hormone
Lymecycline
Lynoestrenol
Lypressin
Lysuride Maleate
Mafenide
Mafenide Acetate
Mafenide Hydrochloride
Mafenide 5.0 per cent Eye drops Propionate
Magnesium Fluoride
Magnesium Metrizoate
Mandragora Autumnalis
Mannomustine Hydrochloride
Maprotiline Hydrochloride
Mebanazine

Substance       Maximum       Route of       Treatment limitations       Maximum         strength       administration,       administration,       quantity         use or       pharmaceutical       form       quantity         Webendazole       For oral       100mg (MD)       Container         use in the       treatment of       containing       containing         in adults       not more       in       not more         in adults       not more       than       and in         and in       s00mg of       children not       Mebendazole         less than 2       years       years       Mebendazole         Webeverine       [* <sup>250</sup> (a) For       [* <sup>250</sup> (b) 100 mg (MD)]       symptomatic         relief       of       irritable       bowel       symptomatic         symptomatic       relief       of       irritable       bowel         syndrome       (b)       For       [* <sup>250</sup> (b) 100 mg (MDD)]       than         Webeverine       amoate       weinstatic       irritable       bowel         syndrome       weinstatic       irritable       irritable       irritable         bowel       syndrome       irritable       irritable			from the restri only medicine	ctions on the sale and supp s	ly of
use in the       or package         reatment of       containing         enterobiasis       not more         in adults       than         and in       800mg of         children not       less than 2         years       years         Mebeverine       [*250(a) For       [*250(a) 135 mg (MD)         Hydrochloride       the       405 mg (MDD)]         symptomatic       relief of       of         irritable       bowel       synptomatic         vess other 300 mg (MDD)]       than       than         uses other 300 mg (MDD)]       than       than         yamoate       of       irritable       bowel         synptomatic       relief of       of       irritable         bowel       synptomatic       relief of       irritable         bowel       syndrome]       well       syndrome]         Mebeverine       amoate       irritable       bowel         'amoate       well       syndrome]       irritable         Mebhydrolin       Meconylamine       irritable       irritable         'tydrochloride       irritable       irritable       irritable         Meclofenoxate       irri	Substance	Maximum	Route of administratio use or pharmaceuti	Treatment limitations on,	Maximum
Hydrochloride       The (a) for ing (MDD)]         symptomatic       relief of         irritable       bowel         syndrome       (b) For [ <sup>F250</sup> (b) 100 mg (MD)]         uses other 300 mg (MDD)]       than the         symptomatic       relief of         irritable       bowel         syndrome       (b) For [ <sup>F250</sup> (b) 100 mg (MD)]         uses other 300 mg (MDD)]       than the         symptomatic       relief of         irritable       bowel         syndrome]       Webeverine         Pamoate       Webhydrolin         Mebhydrolin       Mecamylamine         Hydrochloride       Mecillinam         Meclofenoxate       Hydrochloride         Medigoxin       Medrogestone         Medroxyprogesterone       Sume Sume Sume Sume Sume Sume Sume Sume	Mebendazole		use in the treatment of enterobiasis in adults and in children not less than 2	100mg (MD)	or package containing not more than 800mg of
uses other 300 mg (MDD)] than the symptomatic relief of irritable bowel syndrome] Mebeverine Pamoate Mebhydrolin Mebhydrolin Mebhydrolin Napadisylate Mecamylamine Hydrochloride Mecillinam Meclofenoxate Hydrochloride Medigoxin Medrogestone Medroxyprogesterone	Mebeverine Hydrochloride		the symptomatic relief of irritable bowel		
Pamoate Mebhydrolin Mebhydrolin Napadisylate Mecamylamine Hydrochloride Mecillinam Meclofenoxate Hydrochloride Medigoxin Medigoxin Medrogestone			uses other than the symptomatic relief of irritable bowel		
Mebhydrolin Napadisylate Mecamylamine Hydrochloride Mecillinam Meclofenoxate Hydrochloride Medigoxin Medrogestone Medrogesterone	Mebeverine Pamoate				
Napadisylate Mecamylamine Hydrochloride Mecillinam Meclofenoxate Hydrochloride Medigoxin Medrogestone Medroxyprogesterone	Mebhydrolin				
Hydrochloride Mecillinam Meclofenoxate Hydrochloride Medigoxin Medrogestone Medroxyprogesterone	Mebhydrolin Napadisylate				
Meclofenoxate Hydrochloride Medigoxin Medrogestone Medroxyprogesterone	Mecamylamine Hydrochloride	2			
Hydrochloride Medigoxin Medrogestone Medroxyprogesterone	Mecillinam				
Medrogestone Medroxyprogesterone	Meclofenoxate Hydrochloride				
Medroxyprogesterone	Medigoxin				
	Medrogestone				
Acciaic	Medroxyproge Acetate	sterone			

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity		
Mefenamic Acid						
Mefloquine Hydrochlorid	e					
Mefruside						
Megestrol						
Megestrol Acetate						
Meglumine Gadopentetat	e					
Meglumine Iodoxamate						
Meglumine Ioglycamate						
Meglumine Iothalamate						
Meglumine Iotroxate						
Meglumine Ioxaglate						
[ <sup>F214</sup> Meloxica	m]					
Melphalan						
Melphalan Hydrochlorid	e					
Menotrophin						
Mepenzolate			25mg (MD)			
Bromide			75mg (MDD)			
Mephenesin						
Mephenesin Carbamate						
Mepivacaine Hydrochlorid	e	Any use except ophthalmic use				

		Exemptions from the restrictions on the sale and supply of prescription only medicines				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratic use or pharmaceutic form		Column 5 Maximum quantity		
Meptazinol Hydrochloric	le					
Mequitazine						
[ <sup>F203</sup> Mercapta Bitartrate]	amine					
Mercaptopur	ine					
Mersalyl						
Mersalyl Acid						
Mesalazine						
Mesna						
Mestranol						
Metaraminol Tartrate						
Metergoline						
Metformin Hydrochloric	le					
Methacycline	e					
Methacycline Calcium	9					
Methacycline Hydrochloric						
Methallenoes	stril					
Methicillin Sodium						
Methixene						
Methixene Hydrochloric	le					
Methocarban	nol					
Methocidin		Throat lozenges and throat pastilles				

	prescription	n only medicine	ctions on the sale and support	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceutin form		Column 5 Maximum quantity
Methohexito Sodium	ne			
Methoin				
Methoserpid	ine			
Methotrexate	9			
Methotrexate Sodium	5			
Methotrimer	orazine			
Methotrimer Hydrochlorid				
Methotrimep Maleate	orazine			
Methoxamin Hydrochlorid		Nasal sprays or nasal drops not containing liquid paraffin as a vehicle		
Methsuximi	le			
Methyclothia	azide			
Methyldopa				
Methyldopat Hydrochlorid				
Methylepheo			30mg (MD)	
Hydrochlorid	de		60mg (MDD)	
Methylpredn	isolone			
Methylpredn Acetate	isolone			
Methylpredn Sodium Succinate	isolone			
Methylthiou	racil			
Methysergid Maleate	e			
			101	

		tions from the restrictions on the sale and supply of iption only medicines			
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity	
Metipranolol					
Metirosine					
Metoclopran Hydrochloric					
Metolazone					
Metoprolol Fumarate					
Metoprolol Succinate					
Metoprolol Tartrate					
Metronidazo	le				
Metronidazo Benzoate	le				
Metyrapone					
Mexiletine Hydrochlorid	le				
Mezlocillin Sodium					
Mianserin Hydrochlorid	le				
Miconazole		External but in the case of vaginal use only external use for the treatment of vaginal candidiasis			
Miconazole Nitrate		External but in the case of vaginal use only external use for the treatment			

	Exemptions from the restrictions on the sale and supply of prescription only medicines			
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity
		of vaginal candidiasis		
Mifepristone	;			
Miglitol				
Milrinone				
Milrinone Lactate				
Minocycline				
Minocycline Hydrochlorid				
Minoxidil	[ <sup>F251</sup> (1) 2.0 per cent]	[ <sup>F251</sup> (1) External		
	[ <sup>F251</sup> (2) 5.0 per cent	(2) External use for the treatment of alopecia androgenetic in men aged 18 to 65 (but not in women);]	a,	
[ <sup>F199</sup> Mirtazap	oine]			
Misoprostol				
Mitobronitol				
Mitomycin				
Mitozantrone Hydrochlorie				
Mivacurium Chloride				
[ <sup>F224</sup> Mizolast	ine]			
Moclobemid	e			
[ <sup>F203</sup> Modafin	il]			
[ <sup>F200</sup> Moexipr Hydrochloric	il de]			

		from the restrictions on the only medicines	he sale and supp	ly of
Column 1 Substance	Column 2 Maximum strength	Column 3 Column	4 tt limitations	Column 5 Maximum quantity
Molgramosti	m			
Molindone Hydrochlorid	e			
Mometasone Furoate				
Moracizine Hydrochlorid	e			
Morazone Hydrochlorid	e			
[ <sup>F199</sup> Moxonid	ine]			
Mupirocin				
Mupirocin Calcium				
Mustine Hydrochlorid	e			
Nabilone				
Nabumetone				
Nadolol				
Nafarelin Acetate				
Naftidrofuryl Oxalate				
Naftifine Hydrochlorid	e			
Nalbuphine Hydrochlorid	e			
Nalidixic Acid				
Nalorphine Hydrobromid	le			
Naloxone Hydrochlorid	e			
Naltrexone Hydrochlorid	e			

			ctions on the sale and sup	ply of
Column 1 Substance	prescription Column 2 Maximum strength	only medicine Column 3 Route of administratio use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
Naphazoline Hydrochlorid		(1) Nasal sprays or nasal drops not containing liquid paraffin as a vehicle		
	(2) 0.015 per cent	(2) Eye drops		
Naphazoline Nitrate	0.05 per cent	Nasal sprays or nasal drops not containing liquid paraffin as a vehicle		
Naproxen				
Naproxen Sodium				
[ <sup>F203</sup> Naratript Hydrochlorid				
Natamycin				
[ <sup>F215</sup> Nebivolo Hydrochlorid				
Nedocromil Sodium	[ <sup>F252</sup> 2.0 per cent]	[ <sup>F252</sup> For the prevention, relief and treatment of seasonal and perennial allergic conjunctivitis	5]	[ <sup>F252</sup> Container or package containing not more than 3 ml of medicinal product]
Nefazodone Hydrochlorid	e			
Nefopam Hydrochlorid	e			
Neomycin				

110

Oleate Neomycin Palmitate Neomycin Sulphate Neomycin Undecanoate Neostigmine Bromide Smonide Neostigmine Methylsulphate Neostigmine Methylsulphate Netilmicin Sulphate Nicardipine Hydrochloride Nicardipine Hydrochloride Nicergoline I <sup>*24</sup> Niceritrol] Nicotinic Acid except for the treatment of hyperlipidaemia Nicoumalone Nifedipine Nifed			from the restri only medicine	ictions on the sale and suppers	ply of
Oleate Neomycin Palmitate Neomycin Sulphate Neomycin Undecanoate Neostigmine Bromide Smonide Neostigmine Methylsulphate Neostigmine Methylsulphate Netilmicin Sulphate Nicardipine Hydrochloride Nicardipine Hydrochloride Nicergoline I <sup>*24</sup> Niceritrol] Nicotinic Acid except for the treatment of hyperlipidaemia Nicoumalone Nifedipine Nifed		Maximum	Route of administrati use or pharmaceut	Treatment limitations ion,	Maximum
Palmitate Neomycin Sulphate Neomycin Undecanoate Neostigmine Bromide Neostigmine Methylsulphate Netilmicin Sulphate Nicardipine Hydrochloride Nicergoline (* <sup>224</sup> Niceritrol] Nicotinic Any use, 600mg (MDD) Acid except for the treatment of hyperlipidaemia Nicoumalone Nifedipine Nifenazone Nikethamide (* <sup>228</sup> Nilutamide] Nimodipine Niridazole (* <sup>228</sup> Nisoldipine] Nitrendipine Nitendipine	Neomycin Oleate				
Sulphate         Neomycin         Undecanoate         Neostigmine         Bromide         Neostigmine         Methylsulphate         Netilmicin         Sulphate         Nicardipine         Hydrochloride         Nicergoline         I*2*Niceritrol]         Nicotinic         Any use,       600mg (MDD)         Acid       except for the treatment of hyperlipidaemia         Nicoumalone         Nifedipine         Nimodipine         Niridazole         I*2**Nisoldipine]         Nirtendipine	Neomycin Palmitate				
Undecanoate Neostigmine Bromide Neostigmine Methylsulphate Netilmicin Sulphate Nicardipine Hydrochloride Nicergoline [ <sup>F224</sup> Niceritrol] Nicotinic Any use, 600mg (MDD) Acid except for the treatment of hyperlipidaemia Nicoumalone Nifedipine Nifenazone Nikethamide [ <sup>F202</sup> Nilutamide] Nimodipine Niridazole [ <sup>F205</sup> Nisoldipine] Nitrendipine	Neomycin Sulphate				
Bromide Bromide Neostigmine Methylsulphate Netilmicin Sulphate Nicardipine Hydrochloride Nicergoline I <sup>*224</sup> Niceritrol] Nicotinic Acid Any use, 600mg (MDD) Acid except for the treatment of hyperlipidaemia Nicoumalone Nifedipine Nifenazone Nikethamide I <sup>*202</sup> Nilutamide] Nimodipine Niridazole I <sup>*215</sup> Nisoldipine] Nitrendipine	Neomycin Undecanoate				
MethyIsulphate Netilmicin Sulphate Nicardipine Hydrochloride Nicergoline [ <sup>F224</sup> Niceritrol] Nicotinic Any use, 600mg (MDD) Acid except for the treatment of hyperlipidaemia Nicoumalone Nifedipine Nifedipine [ <sup>F202</sup> Nilutamide] Nimodipine Niridazole [ <sup>F215</sup> Nisoldipine] Nitrendipine	Neostigmine Bromide				
Sulphate         Nicardipine         Hydrochloride         Nicergoline         I*224         I*224         Niceritrol]         Nicotinic       Any use, 600mg (MDD)         Acid       except for the treatment of hyperlipidaemia         Nicoumalone         Nifedipine         Nifenazone         Nikethamide         I*202         I*204         Nimodipine         Niridazole         I*215         Nisoldipine]         Niterndipine	Neostigmine Methylsulpha	te			
Hydrochloride         Nicergoline         I <sup>*224</sup> Niceritrol]         Nicotinic       Any use, 600mg (MDD)         Acid       except         for the         treatment of         hyperlipidaemia         Nicoumalone         Nifedipine         Nifedipine         Nikethamide         I <sup>*202</sup> Nilutamide]         Nimodipine         Niridazole         I <sup>*215</sup> Nisoldipine]         Nitrendipine	Netilmicin Sulphate				
I*224 Niceritrol]         Nicotinic       Any use, 600mg (MDD)         Acid       except         for the       treatment of         hyperlipidaemia       Nicoumalone         Nifedipine       Nifedipine         Nikethamide       I*202 Nilutamide]         Nimodipine       I*202 Nilutamide]         Niridazole       I*215 Nisoldipine]         Nitrendipine       Nitrendipine	Nicardipine Hydrochloride	e			
Nicotinic       Any use, 600mg (MDD)         Acid       except         for the       treatment of         hyperlipidaemia       Nicoumalone         Nifedipine       Vifedipine         Nikethamide       Image: State St	Nicergoline				
Acid     except       for the     treatment of       hyperlipidaemia       Nicoumalone       Nifedipine       Nifenazone       Nikethamide       [ <sup>F202</sup> Nilutamide]       Nimodipine       Niridazole       [ <sup>F215</sup> Nisoldipine]       Nitrendipine	[F224Niceritrol]				
Nifedipine Nifenazone Nikethamide [ <sup>F202</sup> Nilutamide] Nimodipine Niridazole [ <sup>F215</sup> Nisoldipine] Nitrendipine	Nicotinic Acid		except for the treatment of		
Nifenazone Nikethamide [ <sup>F202</sup> Nilutamide] Nimodipine Niridazole [ <sup>F215</sup> Nisoldipine] Nitrendipine	Nicoumalone				
Nikethamide [ <sup>F202</sup> Nilutamide] Nimodipine Niridazole [ <sup>F215</sup> Nisoldipine] Nitrendipine	Nifedipine				
[ <sup>F202</sup> Nilutamide] Nimodipine Niridazole [ <sup>F215</sup> Nisoldipine] Nitrendipine	Nifenazone				
Nimodipine Niridazole [ <sup>F215</sup> Nisoldipine] Nitrendipine	Nikethamide				
Niridazole [ <sup>F215</sup> Nisoldipine] Nitrendipine	[ <sup>F202</sup> Nilutamid	e]			
[ <sup>F215</sup> Nisoldipine] Nitrendipine	Nimodipine				
Nitrendipine	Niridazole				
-	[ <sup>F215</sup> Nisoldipir	ne]			
	Nitrendipine				
Nitrofurantoin	Nitrofurantoir	1			

	prescription	from the restr n only medicine	ictions on the sale and supper	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity
Nitrofurazon	e			
Nizatidine		For the prevention [ <sup>F253</sup> and treatment] of the symptoms of food- related heartburn [ <sup>F253</sup> and meal- induced indigestion] For use in adults and children not less than 16	75mg (MD) [ <sup>F254</sup> 150mg (MDD)] [ <sup>F255</sup> For a maximum period of 14 days]	
Nomifensine Maleate		years		
Noradrenalin Noradrenalin Acid Tartrate				
Norethistero	ne			
Norethisteron Acetate	ie			
Norethisteron Enanthate	ne			
Norethynodr	el			
Norfloxacin				
Norgestimate				
Norgestrel				
Nortriptyline Hydrochlorid				
Noscapine				

		from the restrictions on the sale and sup only medicines	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Column 4 Route of Treatment limitations administration, use or pharmaceutical form	Column 5 Maximum quantity
Noscapine Hydrochlorid	le		
Novobiocin Calcium			
Novobiocin Sodium			
Nux Vomica Seed			
Nystatin	[ <sup>F256</sup> 3.0 per cent]	[ <sup>F256</sup> External For use in combination with Hydrocortisone of maximum strength 0.5 per cent for intertrigo For use in adults and children not less than 10 years]	[ <sup>F256</sup> Container or package containing not more than 15g of medicinal product]
Octacosactrir	1		
Octreotide			
Oestradiol Oestradiol Benzoate			
Oestradiol Cypionate			
Oestradiol Dipropionate			
Oestradiol Diundecanoa	te		
Oestradiol Enanthate			

	prescription	n only medicine		
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 5 Maximum quantity	
Oestradiol Phenylpropie	onate			
Oestradiol Undecanoate	;			
Oestradiol Valerate				
Oestriol				
Oestriol Succinate				
Oestrogenic Substances Conjugated				
Oestrone				
Ofloxacin				
Olsalazine Sodium				
Omeprazole				
[ <sup>F199</sup> Omepraz Magnesium]	zole			
Ondansetron Hydrochlorid				
Orciprenalin Sulphate	e			
Orphenadrin Citrate	e			
Orphenadrin Hydrochlorid				
Ouabain				
Ovarian Gland Dried				
Oxamniquin	e			
Oxantel Embonate				
Oxaprozin				

		from the restruction only medicine	ictions on the sale and sup	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity
Oxatomide				
Oxedrine Tartrate				
Oxethazaine			10mg (MD)	Container
			30mg (MDD)	or package containing not more than 400mg of Oxethazaine
Oxitropium Bromide				
Oxolinic Acid				
Oxpentifyllin	ne			
Oxprenolol Hydrochloric	le			
Oxybuprocai Hydrochloric		Non- ophthalmic use		
Oxybutynin Hydrochloric	le			
Oxypertine				
Oxypertine Hydrochloric	le			
Oxyphenbuta	azone			
Oxyphencyc Hydrochloric				
Oxyphenoniu Bromide	ım		5mg (MD) 15mg (MDD)	
Oxytetracycl	ine		/	
Oxytetracycl Calcium				
Oxytetracycl Dihydrate	ine			

		from the restri	ctions on the sale and suppl	y of	
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity	
Oxytetracycl Hydrochlorid					
Oxytocin, natural					
Oxytocin, synthetic					
Pancreatin	(1) 21,000 European Pharmacopo units of lipase per capsule	(1) capsules eia			
	(2) 25,000 European Pharmacopo units of lipase per gram	(2) powder eia			
Pancuroniun Bromide	-				
[ <sup>F214</sup> Pantopra Sodium]	zole				
Papaverine		(1) By inhaler			
		(2)	(2) 50mg (MD)		
		Otherwise than by inhaler	150mg (MDD)		
Papaverine Hydrochlorid	le	(1) By inhaler			
		(2) Otherwise	(2) Equivalent of 50mg of Papaverine (MD)		
		than by inhaler	Equivalent of 150mg of Papaverine (MDD)		
[ <sup>F204</sup> Paraceta	mol (1) [ <sup>F257</sup> 2	50m <b>g</b> ]) Non- effervescent tablets and capsules		(1) The quantity sold or supplied in	
			117		

	<i>Exemptions from the restrictions on the sale and supply of prescription only medicines</i>					
Column 1 Substance	Column 2 Maximum strength	Column 3 Column 4 Route of Treatment limitations administration, use or pharmaceutical form	Column 5 Maximum quantity			
	(2) 500 mg	[ <sup>F258</sup> wholly or mainly] for use in children aged less than 12 years	one container or package shall not exceed 32 The quantity of			
		(3) All preparations other than non- effervescent tablets and capsules	(2) The quantity sold or supplied in one container or package shall not exceed 32 The quantity of non- effervescent tablets, capsules or a			

	<i>Exemptions from the restrictions on the sale and supply of prescription only medicines</i>					
Column I Column 2 Substance Maximum strength	Column 3 Column 4 Route of Treatment limitations administration, use or pharmaceutical form	Column 5 Maximum quantity				
		combination of both sold or supplied to a person at any one time shall not exceed 100]				
araldehyde		-				
aramethadione						
aramethasone						
arathyroid Hand						
argyline Iydrochloride						
aroxetine Iydrochloride						
ecilocin						
enamecillin						
enbutolol ulphate						
<sup>7214</sup> Penciclovir]						
enicillamine						
enicillamine Iydrochloride						
entamidine sethionate						
enthienate Bromide	5mg (MD) 15mg (MDD)					

		from the restri only medicine	ictions on the sale and sup	oply of	
Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceuti form		Column 5 Maximum quantity	
Pentolinium Tartrate					
Perfluamine					
Pergolide Mesylate					
Perhexiline Maleate					
Pericyazine					
Perindopril					
Perindopril Erbumine					
Perphenazine					
Phenacetin (	0.1 per cent				
Phenazone		External			
Phenazone Salicylate					
Phenbutrazate Hydrochloride					
Phenelzine Sulphate					
Phenethicillin Potassium					
Phenformin Hydrochloride					
Phenglutarimic Hydrochloride	le				
Phenindione					
[ <sup>F260</sup> Phenolphth	nalein.]				
Phenoxybenza Hydrochloride					
Phenoxymethy	lpenicillin				
Phenoxymethy Calcium	lpenicillin				

		from the restring from the restrict from the res	ictions on the sale and supp	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
Phenoxymeth Potassium	ylpenicillin			
Phenprocoum	on			
Phensuximide	;			
Phentolamine Hydrochloride				
Phentolamine Mesylate				
Phenylbutazor	ne			
Phenylbutazor Sodium	ne			
Phenylpropan		Internal		
Hydrochloride	-	(1) all preparations except prolonged release capsules, nasal sprays and nasal drops	(1) 25mg (MD) 100mg (MDD)	
		(2) prolonged	(2) 50mg (MD)	
		release capsules	100mg (MDD)	
	(3) 2.0 per cent	(3) nasal sprays and nasal drops		
Phenytoin				
Phenytoin Sodium				
Phthalylsulph	athiazole			
Physostigmine	e			
Physostigmine Aminoxide Salicylate	e			

Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceution form	Column 4 Treatment limitations on,	Column 5 Maximum quantity	
Physostigmir Salicylate	ne				
Physostigmir Sulphate	ie				
[ <sup>F202</sup> Phytome	nadione	Any use except the prevention or treatment of haemorrhagic disorders]	2		
Picrotoxin					
Pilocarpine					
Pilocarpine Hydrochlorid	le				
Pilocarpine Nitrate					
Pimozide					
Pindolol					
Pipenzolate			5mg (MD)		
Bromide			15mg (MDD)		
Piperacillin Sodium					
Piperazine Oestrone Sulphate					
Piperidolate			50mg (MD)		
Hydrochloric	le		150mg (MDD)		
Pipothiazine Palmitate					
Piracetam					
Pirbuterol Acetate					
Pirbuterol Hydrochloric	le				

121

	· ·	from the restri only medicine	ctions on the sale and supples	y of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceution form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
[ <sup>F261</sup> Pirenzep: Dihydrochlor Monohydrate	ride			
Pirenzepine Hydrochlorid	le			
Piretanide				
Piroxicam	0.5 per cent	External For the relief of rheumatic pain, pain of non-serious arthritic conditions and muscular aches, pains and swellings such as strains, sprains and sports injuries For use in adults and children not less than 12 years	For maximum period of 7 days	Container or package containing not more than 30g of medicinal product
[ <sup>F224</sup> Piroxicar Beta- cyclodextrin]				
Pituitary Gland (Whole Dried)		By inhaler		
Pituitary Powdered (Posterior		By inhaler		

Column 1       Column 2       Column 3       Column 4       Column 5         Substance       Maximum       Route of       Treatment limitations       Maximum         guantity       administration,       use or       pharmaceutical       Maximum         Pivampicillin       pivampicillin       pivampicillin       pivampicillin         Pivampicillin       pivampicillin       pivampicillin         Pivampicillin       pivampicillin       pivampicillin         Pivampicillin       pivampicillin       pivampicillin         Pivampicillin       pivampicillin       pivampicillin         Pivampicillin       pivampicillin       pivampicillin         Pivampicillinam       pivampicillin       pivampicillin         Pivampicillinam       pivampicillin       pivampicillin         Pivampicillinam       pivation       picamica         Pizotifen       picamica       picamica         Picamycin       podophyllum       podophyllum       podophyllum         Podophyllum       20.0 per       External       Ointment or         mipregnated       plaster       6mg (MDD)       fomg (MDD)         Polidexide       Polymyxin       B       Sulphate         Polythiazide			from the restring only medicine	ctions on the sale and suppos	ply of	
Pivampicillin HydrochlorideEvenPivmecillinam		Column 2 Maximum	Column 3 Route of administrati use or pharmaceuti	Column 4 Treatment limitations on,	Maximum	
Hydrochloride         Pivmecillinam         Pivmecillinam         Hydrochloride         Pizotifen         Hydrochloride         Pizotifen         Malate         Polophyllotoxin         Podophyllotoxin         Podophyllum         Indian         Resin       Cent         Ointment or impregnated plaster         Poldine       Cent         Methylsulphate       2mg (MD) 6mg (MDD)         Polidexide       Sulphate         Polystradiol       Sulphate         Polytinizide       Sulphate         Poppy Capsule       Sulphate         Potassium       0.0127 per cent         Potassium       0.0127 per	Pivampicillin					
Pivmecillinami Hydrochloride   Pizotifen Pizotifen   Pizotifen Huter   Pizotifen Huter   Pizotifen Huter   Malate Huter   Picamycin Huter   Podophyllumindian External   Podophyllumindian Ointment or impregnated plaster   Podophyllumice External   Odiniment or impregnated plaster Ointment or impregnated plaster   Poldine Methylsulphate External Ointment or impregnated plaster   Polidexide External Ointment or impregnated plaster   Polidexide Samp (MD) Gmg (MDD)   Polidexide Huter   Polystradiol Huter   Polytinazide Huter   Polytinazide Huter   Poppy Capsule Huter   Potassium 0.0127 per Arsenite   Potassium 0.0127 per Arsenite						
Hydrochloride Pizotifen Malate Pizotifen Malate Pizotifen Malate Pizotifen Malate Pizotifen Malate Pizotifen Malate Pizotifen Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Podophyllum Polidexide Polyestradiol Polyestrad	Pivmecillinar	n				
Pizotifen Malate       Picamycin         Plicamycin       Podophyllotoxin         Podophyllotoxin       Podophyllom         Podophyllum Indian       External         Podophyllum Indian       20.0 per Content or impregnated plaster       External         Podophyllum Methylsulphate       External         Poldine Methylsulphate       2mg (MD)         Polidexide       3mg (MDD)         Polidexide       Sulphate         Polytinazide       Sulphate         Polythiazide       Sulphate         Poppy Capsule       0.0127 per Arsenite         Potassium       0.0127 per Cent						
MalatePlicamycinPodophyllotxirPodophyllumPodophyllumIndianPodophyllumPodophyllumContrent or ingregnated plasterPoldine MethylsulpharePolidexidePolidexidePolystradiol PhosphatePolytinazidePolytinazidePolytinazidePolytinazidePoppy CapsuleNontrent or indicePotassium0.0127 per ArsenitePotassium0.0127 per CapsulePotassium0.0127 per CapsulePotassium0.0127 per CapsulePotassium0.0127 per CapsulePotassium0.0127 per CapsulePotassium0.0127 per CapsulePotassium <t< td=""><td>Pizotifen</td><td></td><td></td><td></td><td></td><td></td></t<>	Pizotifen					
Podophyllum Podophyllum Indian Podophyllum 20.0 per Resin cent Biscenal Ointment or impregnated plaster Poldine Methylsulphate 200 per Polidexide Polidexide Polyestradiol Polidexide Polyestradiol Po						
Podophyllum Podophyllum Indian Podophyllum 20.0 per Resin cent Cintment or impregnated plaster Poldine Methylsulphate Polidexide Polyestradiol Phosphate Polyestradiol Phosphate Polythiazide Polythiazide Poppy Capsule O.0127 per Arsenite 0.0127 per cent Cintment or Potassium Cintment or Cintment	Plicamycin					
Podophyllum Indian Podophyllum 20.0 per Resin cent External Ointment or impregnated plaster Poldine Methylsulphate Polidexide Polyestradiol Phosphate Polymyxin B Sulphate Polythiazide Poppy Capsule Potassium 0.0127 per Arsenite 0.0127 per cent V	Podophylloto	xin				
Indian Podophyllum 20.0 per cent External Ointment or impregnated plaster Poldine Methylsulphate Polidexide Polyestradiol Phosphate Polyestradiol Phosphate Polythiazide Polythiazide Polythiazide Poppy Capsule Potassium 0.0127 per Arsenite 0.0127 per Arsenite Votassium Votas Votassium Votasu Votassium Votassium Votassium Vota	Podophyllum	L				
ResincentOintment or impregnated plasterPoldine2mg (MD)Methylsulphate6mg (MDD)Polidexide90Polyestradiol Phosphate90Polymyxin B Sulphate90Polythiazide90Polythiazide90Potassium0.0127 per centPotassium0.0127 per centPotassium0.0127 per cent		L				
Ontment or impregnated plaster2mg (MD)Poldine Methylsulphate2mg (MD)Polidexide6mg (MDD)Polyestradiol Phosphate		20.0 per	External			
Methylsulphate 6mg (MDD) Polidexide Polyestradiol Phosphate Polymyxin B Sulphate Polythiazide Poppy Capsule Potassium 0.0127 per Arsenite cent	Resin	cent	impregnated			
Polidexide Polyestradiol Phosphate Polymyxin B Sulphate Polythiazide Poppy Capsule Potassium 0.0127 per Arsenite cent Potassium	Poldine			2mg (MD)		
Polyestradiol PhosphatePolymyxin B SulphatePolythiazidePoppy CapsulePotassium Arsenite0.0127 per centPotassiumPotassiumSulphate	Methylsulpha	ate		6mg (MDD)		
PhosphatePolymyxin B SulphatePolythiazidePoppy CapsulePotassium Arsenite0.0127 per centPotassium b SulphatePotassium Cent	Polidexide					
B Sulphate Polythiazide Poppy Capsule Potassium 0.0127 per Arsenite cent Potassium						
Poppy Capsule Potassium 0.0127 per Arsenite cent Potassium						
Capsule Potassium 0.0127 per Arsenite cent Potassium	Polythiazide					
Arsenite cent Potassium						
		-				
Bromide	Potassium Bromide					
Potassium Canrenoate						

		from the restruction only medicine	ictions on the sale and suppers	ply of	
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity	
Potassium Clavulanate					
Potassium Perchlorate					
Practolol					
Pralidoxime Chloride					
Pralidoxime Iodide					
Pralidoxime Mesylate					
[ <sup>F203</sup> Pramipex Hydrochlorid					
Pravastatin Sodium					
Prazosin Hydrochlorid	e				
Prednisolone					
Prednisolone Acetate					
Prednisolone Butylacetate					
Prednisolone Hexanoate					
Prednisolone Metasulphob	enzoate				
Prednisolone Metasulphob Sodium	enzoate				
Prednisolone Pivalate					
Prednisolone Sodium Phosphate					

		from the restri only medicine	ctions on the sale and sup	ply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
Prednisolone Steaglate				
Prednisone				
Prednisone Acetate				
Prenalterol Hydrochloric	le			
Prenylamine Lactate				
Prilocaine Hydrochloric	le	Non- ophthalmic use		
Primidone				
Probenecid				
Probucol				
Procainamide Hydrochloric				
Procaine Hydrochloric	le	Non- ophthalmic use		
Procaine Penicillin				
Procarbazine Hydrochloric				
Prochlorpera	zine			
Prochlorpera Edisylate	zine			
Prochlorpera Maleate	z <b>[f<sup>f</sup>&amp;</b> 3mg]	[ <sup>F218</sup> Buccal tablets for the treatment of nausea and vomiting in cases of previously diagnosed migraine	[ <sup>F218</sup> 12mg (MDD)]	[ <sup>F218</sup> Container or package containing not more than 8 tablets]

prescription only medicines         Column 1       Column 2       Column 3       Column 4       Column 5         Substance       Maximum strength       Route of administration, use or pharmaceutical form       Maximum quantity         only. For use in persons aged 18 years and over.]       only. For use in persons       For use in persons         Prochlorperazine       Vears and over.]       Vears and over.]       Vears and over.]         Progesterone       Vears and over.]       Vears Vears         Proligestone       Vears       Vears         Proligestone       Vears       Vears         Promazine       Vears       Vears	
only. For use in persons aged 18 years and over.] Prochlorperazine Mesylate Procyclidine Hydrochloride Progesterone Prolactin Proligestone Proligestone Proligestone Prolintane Hydrochloride Promazine Embonate	
Procyclidine Hydrochloride Progesterone Prolactin Proligestone Prolintane Hydrochloride Promazine Embonate	
Prolactin Proligestone Prolintane Hydrochloride Promazine Embonate Promazine	
Proligestone Prolintane Hydrochloride Promazine Embonate Promazine	
Prolintane Hydrochloride Promazine Embonate Promazine	
Hydrochloride Promazine Embonate Promazine	
Embonate Promazine	
Propafenone	
Propafenone Hydrochloride	
Propanidid	
Propantheline 15mg (MD) Bromide 45mg (MDD)	
[ <sup>F215</sup> Propiverine Hydrochloride]	
Propofol	
Propranolol Hydrochloride	
Propylthiouracil	
Proquazone	
Protamine Sulphate	

		from the restri	ctions on the sale and supply	v of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceuti form		Column 5 Maximum quantity
Prothionamic	le			
Protirelin				
Protriptyline Hydrochlorid	e			
Proxymetaca Hydrochlorid		Non- ophthalmic use		
Pseudoephed Hydrochlorid		Internal	(a) In the case of a prolonged release preparation	
			120mg (MD)	
			240mg (MDD)	
			(b) in any other case 60mg (MD)	
			240mg (MDD)	
Pseudoephed	rine		60mg (MD)	
Sulphate			180mg (MDD)	
Pyrantel Embonate		(a) For the treatment of enterobiosis, in adults and children not less than 12 years	(a) 750mg MDD (as a single dose)	(a) Container or package containing not more than 750mg of Pyrantel Embonate
		(b) For the treatment of enterobiosis, in children less than 12 years but not less than 6 years	(b) 500mg MDD (as a single dose)	(b) Container or package containing not more than 750mg of Pyrantel Embonate
		(c) For the treatment of enterobiosis in children less than 6	(c) 250mg MDD (as a single dose)	(c) Container or package containing not more

		from the restrict only medicine.	ctions on the sale and supp	ply of
Column 1	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
		years but not less than 2 years		than 750mg of Pyrantel Embonate
Pyrantel Tartrate				
Pyrazinamide				
Pyridostigmine Bromide	e			
Pyrimethamine	e			
[ <sup>F215</sup> Quetiapine Fumarate]	2			
[ <sup>F200</sup> Quinagolic Hydrochloride				
Quinapril				
[ <sup>F261</sup> Quinapril Hydrochloride	1			
Quinestradol	-			
Quinestrol				
Quinethazone				
Quinidine				
Quinidine Bisulphate				
Quinidine Polygalacturor	nate			
Quinidine Sulphate				
Quinine			100mg (MD)	
			300mg (MDD)	
Quinine Bisulphate			Equivalent of 100mg of Quinine (MD)	
			Equivalent of 300mg of Quinine (MDD)	
Quinine Cinchophen			Equivalent of 100mg of Quinine (MD)	
			128	

	1	from the restrictions on the sale and sup only medicines	oply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Column 4 Route of Treatment limitations administration, use or pharmaceutical form	Column 5 Maximum quantity
		Equivalent of 300mg of Quinine (MDD)	
Quinine Dihydrochlo	ride	Equivalent of 100mg of Quinine (MD)	
		Equivalent of 300mg of Quinine (MDD)	
Quinine Ethyl		Equivalent of 100mg of Quinine (MD)	
Carbonate		Equivalent of 300mg of Quinine (MDD)	
Quinine Glycerophos	phate	Equivalent of 100mg of Quinine (MD)	
		Equivalent of 300mg of Quinine (MDD)	
Quinine Hydrobromio	de	Equivalent of 100mg of Quinine (MD)	
		Equivalent of 300mg of Quinine (MDD)	
Quinine Hydrochlorid	le	Equivalent of 100mg of Quinine (MD)	
		Equivalent of 300mg of Quinine (MDD)	
Quinine Iodobismuth	ate	Equivalent of 100mg of Quinine (MD)	
		Equivalent of 300mg of Quinine (MDD)	
Quinine Phosphate		Equivalent of 100mg of Quinine (MD)	
-		Equivalent of 300mg of Quinine (MDD)	
Quinine Salicylate		Equivalent of 100mg of Quinine (MD)	
-		Equivalent of 300mg of Quinine (MDD)	
Quinine Sulphate		Equivalent of 100mg of Quinine (MD) 129	

		from the restrict only medicine	ctions on the sale and supply	y of
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity
			Equivalent of 300mg of Quinine (MDD)	
Quinine Tannate			Equivalent of 100mg of Quinine (MD)	
			Equivalent of 300mg of Quinine (MDD)	
Quinine in combination with Urea Hydrochloride	;			
Ramipril				
[ <sup>F199</sup> Ranitidine Bismuth Citrate]				
Ranitidine Hydrochloride		For the short term	Equivalent to 75mg of Ranitidine (MD)	
		symptomatic relief of heartburn,	Equivalent to 300mg of Ranitidine (MDD)	
		dyspepsia, indigestion, acid indigestion and hyperacidity [ <sup>F262</sup> or the prevention of these symptoms when associated	For a maximum period of 14 days	
		with consuming food and drink]		
Rauwolfia Serpentina				
Rauwolfia Vomitoria				

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratuuse or pharmaceut form		Column 5 Maximum quantity		
Razoxane						
[ <sup>F203</sup> Reboxetin Mesilate]	ne					
Remoxipride Hydrochlorid	e					
Reproterol Hydrochlorid	e					
Rescinnamine	e					
Reserpine						
Rifabutin						
Rifampicin						
Rifampicin Sodium						
Rifamycin						
F <sup>199</sup> Rimexolo	one]					
Rimiterol Hydrobromid	e					
Risperidone						
Ritodrine Hydrochlorid	e					
Rolitetracycli Nitrate	ne					
( <sup>F219</sup> Ropinirol Hydrochlorid						
Sabadilla						
Salbutamol						
Salbutamol Sulphate						
Salcatonin						
Salcatonin Acetate						
Salmefamol						

	<i>Exemptions from the restrictions on the sale and supply of prescription only medicines</i>					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity		
Salmeterol Xinafoate						
Salsalate						
Saralasin Acetate						
Selegiline Hydrochlorid	e					
Semisodium Valproate						
[ <sup>F203</sup> Sertindol	e]					
[ <sup>F199</sup> Sertraline Hydrochlorid						
Serum Gonadotroph						
[ <sup>F199</sup> Sevoflura	ine]					
Silver Sulphadiazin						
Simvastatin						
Sissomicin						
Sissomicin Sulphate						
Snake Venoms						
Sodium Acetrizoate						
Sodium Aminosalicyl	ate					
Sodium Antimonylglu	uconate					
Sodium Arsanilate						
Sodium Arsenate						

		emptions from the restrictions on the sale and supply of scription only medicines		
Column 1 Substance	Column 2 Maximum strength	Column 3 Column 4 Route of Treatment limitations administration, use or pharmaceutical form	Column 5 Maximum quantity	
Sodium Arsenite	0.013 per cent			
Sodium Bromide				
Sodium Clodronate				
Sodium Cromoglyca	te	(a) For nasal admistration		
	(b) 2.0 per cent	(b) For the treatment of acute seasonal allergic conjunctivitis [ <sup>F263</sup> or perennial allergic conjunctivitis]	(b) Container or package containing not more than 10ml of medicinal product	
		In the form of aqueous eye drops		
	(c) 4.0 per cent	(c) For the treatment of acute seasonal allergic conjunctivitis	(c) Container or package containing not more than 5g of medicinal	
		In the form of an eye ointment	product	
Sodium Ethacrynate				
Sodium Fluoride	(1) 0.33 per cent	(1) Dentifrices		
		(2) Other preparations for use in the prevention		
		133		

		Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceuti form	Column 4 Treatment limitations on,	Column 5 Maximum quantity			
		of dental caries					
		In the form of					
		(a) tablets or drops	(a) 2.2 mg (MDD)				
	(b) 0.2 per cent	(b) mouth rinses other than those for daily use					
	(c) 0.05 per cent	(c) mouth rinses for daily use					
Sodium Fusidate							
Sodium Metrizoate							
Sodium Monofluoro	1.14 per phæspthate	Dentrifrice					
Sodium Oxidronate							
Sodium Stiboglucon	ate						
Sodium Valproate							
Somatorelin Acetate							
Sotalol Hydrochlori	de						
[ <sup>F200</sup> Sparflox	acin]						
Spectinomy	cin						
Spectinomy Hydrochlori							
Spiramycin							
Spiramycin Adipate							
			124				

		from the restrictions on the sale and s only medicines	supply of
Column 1 Substance	Column 2 Maximum strength	Column 3 Column 4 Route of Treatment limitations administration, use or pharmaceutical form	Column 5 Maximum quantity
Spironolacto	one		
Stannous Fluoride	([ <sup>F264</sup> 1]) 0.62 per cent	([ <sup>F264</sup> 1]) Dentifrice	
	[ <sup>F264</sup> (2) 0.4 per]	[ <sup>F264</sup> (2) Dental gels for use in the prevention and treatment of dental caries and decalcification of the teeth]	
Stilboestrol			
Stilboestrol Dipropionate	5		
Streptodorna	ise	External	
Streptokinas	e	External	
Streptomyci	n		
Streptomyci Sulphate	n		
Strychnine			
Strychnine Arsenate			
Strychnine Hydrochlori	de		
[ <sup>F202</sup> Strychni Nitrate]	ne		
Styramate			
Succinylsulp	ohathiazole		
Sucralfate			
Sulbactam Sodium			

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity		
Sulbenicillin						
Sulbenicillin Sodium						
Sulconazole Nitrate		External (except vaginal)				
[ <sup>F202</sup> Sulfaben	zamide]					
Sulfacytine						
Sulfadoxine						
Sulfamerazin	e					
Sulfamerazin Sodium	e					
Sulfametopy	razine					
Sulfamonom	ethoxine					
Sulindac						
Sulphacetam	ide					
Sulphacetam Sodium	ide					
Sulphadiazin	e					
Sulphadiazin Sodium	e					
Sulphadimet	noxine					
Sulphadimid	ine					
Sulphadimid Sodium	ine					
Sulphafurazo	le					
Sulphafurazo Diethanolam						
Sulphaguanic	line					
Sulphaloxic Acid						
Sulphamethiz	zole					

	<i>Exemptions from the restrictions on the sale and supply of prescription only medicines</i>					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administration use or pharmaceution form	Column 4 Treatment limitations on,	Column 5 Maximum quantity		
Sulphametho	oxazole					
Sulphametho	oxydiazine					
Sulphametho	oxypyridazine					
Sulphametho Sodium	oxypyridazine					
Sulphamoxo	le					
Sulphanilam	ide					
Sulphaphena	zole					
Sulphapyridi	ne					
Sulphapyridi Sodium	ne					
Sulphasalazi	ne					
Sulphathiazo	le					
Sulphathiazo Sodium	le					
Sulphaurea						
Sulphinpyraz	zone					
Sulpiride						
Sultamicillin						
Sultamicillin Tosylate						
Sulthiame						
Sumatriptan Succinate						
Suprofen						
Suxamethon Bromide	ium					
Suxamethon Chloride	ium					
Suxethonium Bromide	1					

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceuth form		Column 5 Maximum quantity		
[ <sup>F215</sup> Tacalcito Monohydrate						
Tacrine Hydrochloric	le					
Talampicillir	1					
Talampicillir Hydrochloric						
Talampicillin Napsylate	1					
Tamoxifen						
Tamoxifen Citrate						
[ <sup>F214</sup> Tamsulos Hydrochloric						
[ <sup>F199</sup> Tazarote	ne]					
Tazobactam Sodium						
Teicoplanin						
[ <sup>F203</sup> Temocap Hydrochloric						
Temocillin Sodium						
Tenoxicam						
Terazosin Hydrochloric	le					
Terbinafine	[ <sup>F265</sup> 1.0 per cent]	[ <sup>F266</sup> External use for the treatment of tinea pedis, tinea cruris and tinea corporis. In the form of a gel]		[ <sup>F267</sup> Container or package containing not more than 30 grams of medicinal product]		

		s from the restrictions on the sale and supply of			
Column 1 Substance	prescription Column 2 Maximum strength	only medicine Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity	
[ <sup>F268</sup> Terbinafin] <sup>F268</sup> 1.0 per Hydrochloride]ent]		([ <sup>F269</sup> 1]) [ <sup>F270</sup> Preparation other than spray solutions, for][ <sup>F268</sup> exter use for the treatment of tinea pedis and tinea cruris]		([ <sup>F269</sup> 1]) [ <sup>F268</sup> Container or package containing not more than 15 g of medicinal product.]	
		[ <sup>F271</sup> (2) Spray solutions for external use for the treatment of tinea corporis, tinea cruris and tinea pedis]		[ <sup>F271</sup> (2) Container containing not more than 30ml of medicinal product]	
Terbutaline					
Terbutaline Sulphate					
Terfenadine			F272	F272	
Terlipressin					
Terodiline Hydrochlorid	e				
[ <sup>F203</sup> Testoster	one]				
Tetrabenazine	e				
Tetracosactrin	1				
Tetracosactrin Acetate	1				
Tetracycline					

	<i>Exemptions from the restrictions on the sale and supply of prescription only medicines</i>				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrat use or pharmaceut form		Column 5 Maximum quantity	
Tetracycline Hydrochlorid	de				
Tetracycline Phosphate Complex					
Tetroxoprim					
Thallium Acetate					
Thallous Chloride					
Thiabendazo	ole				
Thiambutosi	ne				
Thiethylpera Malate	zine				
Thiethylpera Maleate	zine				
Thiocarlide					
Thioguanine	1				
Thiopentone Sodium	:				
Thiopropaza Hydrochlorid					
Thiopropera: Mesylate	zine				
Thioridazine	;				
Thioridazine Hydrochlorid					
Thiosinamin	e				
Thiotepa					
Thiothixene					
Thiouracil					
Thymoxamii Hydrochlorid					
Thyroid					
			140		

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratic use or pharmaceutic form	Column 4 Treatment limitations on,	Column 5 Maximum quantity		
Thyrotrophir	l					
Thyroxine Sodium						
Tiamulin Fumarate						
Tiaprofenic Acid						
Tibolone						
Ticarcillin Sodium						
[ <sup>F214</sup> Ticlopidi Hydrochloric						
Tigloidine Hydrobromio	le					
[ <sup>F214</sup> Tiludrona Disodium]	ate					
Timolol Maleate						
Tinidazole						
Tinzaparin						
Tioconazole	(1) 2.0 per cent	(1) External (except vaginal)				
		(2) Vaginal for treatment of vaginal candidiasis				
[ <sup>F200</sup> Tizanidir Hydrochloric						
Tobramycin						
Tobramycin Sulphate						
Tocainide Hydrochlorid	le					

	Exemptions from the restrictions on the sale and supply of prescription only medicines				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administratio use or pharmaceuti form		Column 5 Maximum quantity	
Tofenacin Hydrochlorid	e				
Tolazamide					
Tolazoline Hydrochloride	e	External			
Tolbutamide					
Tolbutamide Sodium					
Tolfenamic Acid					
Tolmetin Sodium					
[ <sup>F199</sup> Topirama	te]				
[ <sup>F224</sup> Torasemic	de]				
[ <sup>F214</sup> Toremifer	ne]				
Tramadol Hydrochlorid	e				
Trandolapril					
Tranexamic Acid					
Tranylcyprom Sulphate	nine				
Trazodone Hydrochlorid	e				
Treosulfan					
Tretinoin					
Triamcinolon	e				
Triamcinolon Acetonide	<b>q</b> <sup>F273</sup> (1)] 0.1 per cent	[ <sup>F273</sup> (1)] For the treatment of common mouth ulcers		[ <sup>F273</sup> (1)] Container or package containing not more than 5g of	

	<i>Exemptions from the restrictions on the sale and supply of prescription only medicines</i>				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity	
				medicinal product	
Triamcinolona Diacetate Triamcinolona Hexacetonide Triamterene Tribavirin Triclofos Sodium Trientine Dihydrochlori Trifluoperazin Trifluoperazin Trifluperidol Trifluperidol Trifluperidol Trifluperidol Trifluperidol	de le le	[ <sup>F274</sup> (2) In the form of a non- pressurised nasal spray, for the treatment of symptoms of seasonal allergic rhinitis in persons aged 18 years and over]	[ <sup>F274</sup> (2) 110mcg per nostril (MD) 110mcg per nostril (MDD) For a maximum period of 3 months]	[ <sup>F274</sup> Container or package containing not more than 3.575mg of Triamcinolone Acetonide]	
Trimeprazine Trimeprazine Tartrate					

Column 1 Substance Trimetaphan Camsylate Trimetazidine	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity	
Camsylate					
Trimetazidine	:				
	:				
Trimetazidine Hydrochloride					
Trimethoprim					
Trimipramine Maleate					
Trimipramine Mesylate					
Tropicamide					
Tropisetron Hydrochloride	:				
Troxidone					
L-		(1) Oral			
Tryptophan		Dietary supplementat	tion		
		(2) External			
Tubocurarine Chloride					
Tulobuterol					
Tulobuterol Hydrochloride	:				
Tyrothricin		Throat lozenges or throat pastilles			
Uramustine					
Urea Stibamine					
Urethane					
Uridine 5'- triphosphate					
Urofollitrophi	n				

	Exemptions from the restrictions on the sale and supply of prescription only medicines				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity	
Urokinase					
Ursodeoxych Acid	noic				
Vaccine: Bacillus Salmonella Typhi					
Vaccine: Poliomyelitis (Oral)	3				
[ <sup>F200</sup> Valaciclo Hydrochloric					
Valproic Acid					
[ <sup>F203</sup> Valsartar	1]				
Vancomycin Hydrochloric	le				
Vasopressin					
Vasopressin Tannate					
Vecuronium Bromide					
[ <sup>F200</sup> Venlafax Hydrochloric					
Verapamil Hydrochloric	le				
Veratrine					
Veratrum, Green					
Veratrum, White					
Vidarabine					
Vigabatrin					
Viloxazine Hydrochloric	le				

	<i>Exemptions from the restrictions on the sale and supply of prescription only medicines</i>				
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form	Column 4 Treatment limitations ion,	Column 5 Maximum quantity	
Vinblastine Sulphate					
Vincristine Sulphate					
Vindesine Sulphate					
Viomycin Pantothenate					
Viomycin Sulphate					
Vitamin A		(1) Internal	(1) 7,500iu (2,250mcg Retinol equivalent) (MDD)		
		(2) External			
Vitamin A Acetate		(1) Internal	<ul><li>(1) Equivalent to 7,500iu</li><li>Vitamin A (2,250mcg</li><li>Retinol equivalent)</li><li>(MDD)</li></ul>		
		(2) External			
Vitamin A Palmitate		(1) Internal	<ul><li>(1) Equivalent to 7,500iu</li><li>Vitamin A (2,250mcg</li><li>Retinol equivalent)</li><li>(MDD)</li></ul>		
		(2) External			
Warfarin					
Warfarin Sodium					
Xamoterol Fumarate					
Xipamide					
Yohimbine Hydrochloride	e				
[ <sup>F200</sup> Zalcitabin	e]				
Zidovudine					

	Exemptions from the restrictions on the sale and supply of prescription only medicines					
Column 1 Substance	Column 2 Maximum strength	Column 3 Route of administrati use or pharmaceut form		Column 5 Maximum quantity		
Zimeldine Hydrochlorid	de					
Zolpidem Tartrate						
Zomepirac Sodium						
Zopiclone						
Zuclopenthiz Acetate	kol					
Zuclopenthiz Decanoate	kol					
Zuclopenthiz Hydrochlorid						

<b>`extu</b> a	l Amendments
F199	Words in Sch. 1 inserted (9.8.2000) by The Prescription Only Medicines (Human Use) Amendment
	Order 2000 (S.I. 2000/1917), arts. 1(1), 3(c)
F200	Words in Sch. 1 inserted (19.1.2000) by The Prescription Only Medicines (Human Use) Amendment
	(No. 2) Order 1999 (S.I. 1999/3463), arts. 1(1), 2(c)
F201	Words in Sch. 1 inserted (24.8.2001) by The Prescription Only Medicines (Human Use) Amendment
	Order 2001 (S.I. 2001/2777), arts. 1(1), 3(a)
F202	Words in Sch. 1 inserted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment
	(No. 3) Order 1998 (S.I. 1998/2081), art. 1(1), Sch.
F203	Sch. 1 entries inserted (24.8.2001) by The Prescription Only Medicines (Human Use) Amendment Ord
	2001 (S.I. 2001/2777), arts. 1(1), <b>3(g)</b>
F204	Words in Sch. 1 inserted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendme
	Order 1997 (S.I. 1997/2044), art. 1(1), Sch. 1
F205	Words in Sch. 1 inserted (22.4.1999) by The Prescription Only Medicines (Human Use) Amendme
	Order 1999 (S.I. 1999/1044), arts. 1(1), <b>2(a)</b>
F206	Words in Sch. 1 substituted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendme
	(No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), <b>3(a)</b>
F207	Sch. 1: entries renumbered (22.4.1999) by The Prescription Only Medicines (Human Use) Amendme
	Order 1999 (S.I. 1999/1044), arts. 1(1), <b>2(a)</b>
F208	Words in Sch. 1 omitted (16.9.1998) by virtue of The Prescription Only Medicines (Human Us
	Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), <b>3(b)</b>
F209	Words in Sch. 1 inserted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendme
	(No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(a)
F210	Words in Sch. 1 substituted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendme
	(No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(a)

- F211 Word in Sch. 1 substituted (22.4.1999) by The Prescription Only Medicines (Human Use) Amendment Order 1999 (S.I. 1999/1044), arts. 1(1), 2(b)
- **F212** Words in Sch. 1 inserted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), **3(c)(ii)**
- **F213** Words in Sch. 1 substituted (16.9.1998) by virtue of The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), **3(c)(i)**
- **F214** Words in Sch. 1 inserted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), **4(h)**
- F215 Words in Sch. 1 inserted (22.4.1999) by The Prescription Only Medicines (Human Use) Amendment Order 1999 (S.I. 1999/1044), arts. 1(1), 2(e)
- F216 Word in Sch. 1 substituted (16.9.1997) by The Prescription Only Medicines (Human Use) Amendment Order 1997 (S.I. 1997/2044), arts. 1(1), 2(a)
- F217 Words in Sch. 1 revoked (31.12.2001) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2001 (S.I. 2001/3942), arts. 1(1), 2(a)
- F218 Sch. 1 entries inserted (24.8.2001) by The Prescription Only Medicines (Human Use) Amendment Order 2001 (S.I. 2001/2777), arts. 1(1), 3(f)
- F219 Sch. 1 entries inserted (31.12.2001) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2001 (S.I. 2001/3942), arts. 1(1), 2(e)
- F220 Sch. 1 entries inserted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(b)
- F221 Words in Sch. 1 inserted (1.6.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 1998 (S.I. 1998/1178), arts. 1(1), 2(a)
- F222 Words in Sch. 1 substituted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), 3(d)
- F223 Words in Sch. 1 substituted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(c)
- F224 Words in Sch. 1 inserted (13.2.1998) by The Prescription Only Medicines (Human Use) Amendment Order 1998 (S.I. 1998/108), arts. 1, 3(e)(i)
- F225 Word in Sch. 1 substituted (19.1.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 1999 (S.I. 1999/3463), arts. 1(1), 2(a)
- F226 Words in Sch. 1 substituted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), 3(e)
- F227 Words in Sch. 1 inserted (24.8.2001) by The Prescription Only Medicines (Human Use) Amendment Order 2001 (S.I. 2001/2777), arts. 1(1), **3(b)**
- **F228** Words in Sch. 1 substituted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), **3(f)(i)**
- **F229** Words in Sch. 1 inserted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), **3(f)(ii)**
- **F230** Words in Sch. 1 omitted (16.9.1998) by virtue of The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), **3(f)(iii)**
- F231 Words in Sch. 1 inserted (31.12.2001) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2001 (S.I. 2001/3942), arts. 1(1), 2(b)(i)
- **F232** Words in Sch. 1 inserted (31.12.2001) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2001 (S.I. 2001/3942), arts. 1(1), **2(b)(ii)**
- F233 Words in Sch. 1 inserted (31.12.2001) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2001 (S.I. 2001/3942), arts. 1(1), 2(b)(iii)
- **F234** Words in Sch. 1 inserted (31.12.2001) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2001 (S.I. 2001/3942), arts. 1(1), **2(b)(iv)**
- F235 Words in Sch. 1 inserted (22.4.1999) by The Prescription Only Medicines (Human Use) Amendment Order 1999 (S.I. 1999/1044), arts. 1(1), 2(c)
- **F236** Sch. 1: entries renumbered (22.4.1999) by The Prescription Only Medicines (Human Use) Amendment Order 1999 (S.I. 1999/1044), arts. 1(1), **2(c)**

- F237 Words in Sch. 1 inserted (13.2.1998) by The Prescription Only Medicines (Human Use) Amendment Order 1998 (S.I. 1998/108), arts. 1, 3(a)
- **F238** Words in Sch. 1 inserted (24.8.2001) by The Prescription Only Medicines (Human Use) Amendment Order 2001 (S.I. 2001/2777), arts. 1(1), **3(c)**
- F239 Words in Sch. 1 inserted (9.8.2000) by The Prescription Only Medicines (Human Use) Amendment Order 2000 (S.I. 2000/1917), arts. 1(1), 3(a)
- F240 Words in Sch. 1 substituted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(d)
- F241 Word in Sch. 1 inserted (13.8.1998) by The Prescription Only Medicines (Human Use) Amendment Order 1998 (S.I. 1998/108), arts. 1, 3(e)(ii)
- **F242** Word in Sch. 1 inserted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), **3(g)(ii)**
- **F243** Words in Sch. 1 inserted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), **3(g)(i)**
- F244 Words in Sch. 1 inserted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), 3(g)(iii)
- F245 Words in Sch. 1 inserted (19.1.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 1999 (S.I. 1999/3463), arts. 1(1), 2(b)
- F246 Sch. 1 entries inserted (1.1.2001) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 2000 (S.I. 2000/3231), arts. 1(1), 2
- F247 Words in Sch. 1 inserted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(e)(i)
- F248 Words in Sch. 1 inserted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(e)(ii)
- F249 Words in Sch. 1 revoked (31.12.2001) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2001 (S.I. 2001/3942), arts. 1(1), 2(c)
- **F250** Words in Sch. 1 substituted (13.2.1998) by The Prescription Only Medicines (Human Use) Amendment Order 1998 (S.I. 1998/108), arts. 1, **3(b)**
- F251 Words in Sch. 1 substituted (1.6.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 1998 (S.I. 1998/1178), arts. 1(1), 2(b)
- **F252** Words in Sch. 1 inserted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), **3(h)**
- **F253** Words in Sch. 1 inserted (1.6.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 1998 (S.I. 1998/1178), arts. 1(1), **2(c)(i)**
- **F254** Words in Sch. 1 inserted (1.6.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 1998 (S.I. 1998/1178), arts. 1(1), **2(c)(ii)**
- F255 Words in Sch. 1 substituted (1.6.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 1998 (S.I. 1998/1178), arts. 1(1), 2(c)(iii)
- F256 Words in Sch. 1 inserted (22.4.1999) by The Prescription Only Medicines (Human Use) Amendment Order 1999 (S.I. 1999/1044), arts. 1(1), 2(d)
- F257 Words in Sch. 1 substituted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(f)(i)
- **F258** Words in Sch. 1 inserted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(f)(ii)
- **F259** Words in Sch. 1 inserted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(f)(iii)
- **F260** Words in Sch. 1 inserted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 1998 (S.I. 1998/1178), arts. 1(1), **2(d)(ii)**
- **F261** Words in Sch. 1 inserted (1.6.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 1998 (S.I. 1998/1178), arts. 1(1), **2(d)(i)**
- F262 Words in Sch. 1 added (13.2.1998) by The Prescription Only Medicines (Human Use) Amendment Order 1998 (S.I. 1998/108), arts. 1, 3(c)

- F263 Words in Sch. 1 inserted (13.2.1998) by The Prescription Only Medicines (Human Use) Amendment Order 1998 (S.I. 1998/108), arts. 1, 3(d)
- F264 Sch. 1: entries renumbered (24.8.2001) by The Prescription Only Medicines (Human Use) Amendment Order 2001 (S.I. 2001/2777), arts. 1(1), **3(d)**
- F265 Words in Sch. 1 inserted (31.12.2001) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2001 (S.I. 2001/3942), arts. 1(1), 2(d)(i)
- F266 Words in Sch. 1 inserted (31.12.2001) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2001 (S.I. 2001/3942), arts. 1(1), 2(d)(ii)
- F267 Words in Sch. 1 inserted (31.12.2001) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2001 (S.I. 2001/3942), arts. 1(1), 2(d)(iii)
- **F268** Words in Sch. 1 inserted (9.8.2000) by The Prescription Only Medicines (Human Use) Amendment Order 2000 (S.I. 2000/1917), arts. 1(1), **3(b)**
- **F269** Sch. 1: entries renumbered (24.8.2001) by The Prescription Only Medicines (Human Use) Amendment Order 2001 (S.I. 2001/2777), arts. 1(1), **3(e)**
- **F270** Words in Sch. 1 inserted (24.8.2001) by The Prescription Only Medicines (Human Use) Amendment Order 2001 (S.I. 2001/2777), arts. 1(1), **3(e)**
- F271 Sch. 1 entries inserted (24.8.2001) by The Prescription Only Medicines (Human Use) Amendment Order 2001 (S.I. 2001/2777), arts. 1(1), 3(e)
- F272 Words in Sch. 1 omitted (16.9.1997) by virtue of The Prescription Only Medicines (Human Use) Amendment Order 1997 (S.I. 1997/2044), arts. 1(1), 2(b)
- **F273** Sch. 1 entries re-numbered (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(g)
- F274 Sch. 1 entries inserted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), 4(g)

### [<sup>F275</sup>SCHEDULE 2

Article 10(1)

### SUBSTANCES WHICH MAY BE EXCLUDED FROM THE CLASS OF PRESCRIPTION ONLY MEDICINES AT HIGH DILUTION

### **Textual Amendments**

F275 Sch. 2 substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), 12

Codeine and its salts

Dihydrocodeine and its salts

Ethylmorphine and its salts

Morphine and its salts

Medicinal Opium

Pholcodine and its salts]

### SCHEDULE 3

Article 2(b)

# DESCRIPTIONS AND CLASSES OF PRESCRIPTION ONLY MEDICINES IN RELATION TO WHICH [<sup>F276</sup>community practitioner nurse prescribers] ARE APPROPRIATE PRACTITIONERS

**Textual Amendments** 

F276 Words in Sch. 3 heading substituted (1.5.2006) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2006 (S.I. 2006/915), arts. 1(1), 13

[<sup>F277</sup>Co-danthramer Capsules NPF]

[<sup>F277</sup>Co-danthramer Capsules, Strong NPF]

Co-danthramer-Oral Suspension NPF

Co-danthramer-Oral Suspension Strong NPF

Co-danthrusate Capsules

[<sup>F277</sup>Co-danthrusate Oral Suspension NPF]

Mebendazole Tablets NPF

Mebendazole Oral Suspension NPF

Miconazole Oral Gel NPF

Nystatin Oral Suspension

Nystatin Pastilles NPF

Streptokinase and Streptodornase Topical Powder NPF

[F278Water for Injections]

### **Textual Amendments**

- F277 Words in Sch. 3 inserted (13.2.1998) by The Prescription Only Medicines (Human Use) Amendment Order 1998 (S.I. 1998/108), arts. 1, 4
- F278 Words in Sch. 3 inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 14

### F279SCHEDULE 3A

Article 3A

### CONTROLLED DRUGS WHICH MAY BE PRESCRIBED, ADMINISTERED OR DIRECTED FOR ADMINISTRATION BY NURSE INDEPENDENT PRESCRIBERS AND CONDITIONS FOR SUCH PRESCRIPTION OR ADMINISTRATION

### **Textual Amendments**

F279 Sch. 3A omitted (1.4.2008) by virtue of The Prescription Only Medicines (Human Use) Amendment Order 2008 (S.I. 2008/464), arts. 1(1), 10

<sup>F280</sup>SCHEDULE 3B

Regulation 3B(3)(a)(iii)

### PARTICULARS FOR CLINICAL MANAGEMENT PLANS

#### **Textual Amendments**

F280 Sch. 3B inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 16

A clinical management plan shall contain the following particulars-

- (a) the name of the patient to whom the plan relates;
- (b) the illnesses or conditions which may be treated by the supplementary prescriber;
- (c) the date on which the plan is to take effect and when it is to be reviewed by the doctor or dentist who is a party to the plan;
- (d) reference to the class or description of medicinal product which may be prescribed or administered under the plan;
- (e) any restrictions or limitations as to the strength or dose of any product which may be prescribed or administered under the plan, and any period of administration or use of any medicinal product which may be prescribed or administered under the plan;
- (f) relevant warnings about the known sensitivities of the patient to, or known difficulties of the patient with, particular medicinal products;
- (g) the arrangements for notification of-
  - (i) suspected or known adverse reactions to any medicinal product which may be prescribed or administered under the plan, and
  - (ii) suspected or known adverse reactions to any other medicinal product taken at the same time as any medicinal product prescribed or administered under the plan; and
- (h) the circumstances in which the supplementary prescriber should refer to, or seek the advice of, the doctor or dentist who is a party to the plan.]

#### SCHEDULE 4

Article 8(4)(c)

### SUBSTANCES NOT TO BE CONTAINED IN A PRESCRIPTION ONLY MEDICINE SOLD OR SUPPLIED UNDER THE EXEMPTION CONFERRED BY ARTICLE 8(3)

- Calcium Bromide
- Calcium Bromidolactobionate

Embutramide

Fencamfamin Hydrochloride

Fluanisone

Hexobarbitone

Hexobarbitone Sodium

Hydrobromic Acid

Meclofenoxate Hydrochloride

Ammonium Bromide

Methohexitone Sodium Pemoline Piracetam Potassium Bromide Prolintane Hydrochloride Sodium Bromide Strychnine Hydrochloride Tacrine Hydrochloride Thiopentone Sodium

### SCHEDULE 5

Article 11(1)(a)

### EXEMPTION FOR CERTAIN PERSONS FROM SECTION 58(2) OF THE ACT

# PART I

### EXEMPTION FROM RESTRICTIONS ON SALE OR SUPPLY

Column 1	Column 2		olumn 3
Persons exempted	Prescription only medicines to which the exemption applies	Со	onditions
1. Persons selling or supplying prescription only medicines to universities, other institutions concerned with higher education or institutions concerned with research.	1. All prescription only medicines	1.	The sale or supply shall be- (a) subject to the presentation of an order signed by the principal of the institution concerned with education or research or the appropriate head of department in charge of a specified course of research stating- (i) the name of the institution for which the prescription only medicine is required, (ii) the purpose for which the prescription only medicine
	153		

	lumn I rsons e	l exempted	Column 2 Prescription only medicines to which the exemption applies	Column . Condition	ns
				(b)	is required, and (iii) the total quantity required, and for the purposes of the education or research with which the institution is concerned.
2.	supp only	ons selling or blying prescription medicines to any of following- a public analyst appointed under section 27 of the Food Safety Act 1990(15) or article 36 of the Food (Northern Ireland) Order 1989(16), an authorized officer within the meaning of section 5(6) of the Food Safety Act 1990,	2. All prescription only medicines.	subject to an order s of any per 1 of this p status of t and the ar only medi shall be o with the e	e or supply shall be the presentation of igned by or on behalf roon listed in column paragraph stating the he person signing it nount of prescription ic ine required, and nly in connection xercise by those f their statutory
	(3)	a sampling officer within the meaning of article 38(1) of the Food (Northern Ireland) Order 1989,			
	(4)	a person duly authorized by an enforcement authority under sections 111 and 112,			
	(5)	a sampling officer within the meaning of Schedule 3 to the Act.			

<sup>(15) 1990</sup> c. 16. (16) S.I. 1989/846 (N.I. 6).

Column 1 Persons exempted	Column 2 Prescription only medicines to which the exemption applies	Column 3 Conditions
3. Persons selling or supplying prescription only medicines to any person employed or engaged in connection with a scheme for testing the quality and checking the amount of the drugs and appliances supplied under the National Health Service Act 1977(17), the National Health Service (Scotland) Act 1978(18) and the Health and Personal Social Services (Northern Ireland) Order 1972(19), or under any subordinate legislation made under those Acts or that Order.	3. All prescription only medicines.	<ul> <li>3. The sale or supply shall be- <ul> <li>(a) subject to the presentation of an order signed by or on behalf of the person so employed or engaged stating the status of the person signing it and the amount of prescription only medicine required, and</li> <li>(b) for the purposes of a scheme referred to in column 1 in this paragraph.</li> </ul> </li> </ul>
4. Registered midwives.	4. [ <sup>F281</sup> Prescription only medicines containing any of the following substances— Diclofenac F282  Hydrocortizone Acetate F282  F282  Miconazole Nystatin Phytomenadione.]	4. The sale or supply shall be only in the course of their professional practice F283 
5. Persons lawfully conducting a retail pharmacy business within the meaning of section 69.	<ul> <li>5. [<sup>F284</sup>Items which are— <ul> <li>(a) prescription</li> <li>only medicines</li> <li>which are not</li> <li>for parenteral</li> <li>administration and</li> <li>which— <ul> <li>(i) are eye drops</li> <li>and are</li> <li>prescription</li> <li>only</li> <li>medicines</li> </ul> </li> </ul></li></ul>	<ul> <li>5. [<sup>F285</sup>The sale or supply shall be subject to the presentation of an order signed by— <ul> <li>(a) a registered</li> <li>optometrist for a medicine listed under item 5(a) in column 2;</li> <li>(b) a registered chiropodist or podiatrist for a</li> </ul> </li> </ul>

<sup>(17) 1977</sup> c. 49. (18) 1978 c. 29. (19) S.I. 1972/1265 (N.I. 14).

Column 1 Persons exempted	Column 2 Prescription only m to which the exempt applies	
	onl com mo per Chi or (ii) are oin and pre onl mea by onl com mo 1.0 Chi or (iii) are pre onl mo 1.0 Chi or (iii) are pre onl mo 1.0 Chi or (iii) are pre onl mo 1.0 Chi or (iii) are pre onl mo 1.0 Chi or (iii) are pre onl mea by onl com mo 1.0 Chi or (iii) are pre onl mea by (iii) Are the of (iii) Are the (iii) Are (iii) Are the (iii) Are (iii) Are	dicines reason y that they tain not re than per cent oramphenicol, scription y dicines reason y that they tain any of following stances— ) Cyclopentolate hydrochloride; ) Fusidic Acid; ) Tropicamide; wing ton only s— torolfine lrochloride am where maximum ength

Column 1	Column 2	Column 3
Persons exempted	Prescription only medicines to which the exemption applies	Conditions
	<ul> <li>weight in weight;</li> <li>(ii) Amorolfine hydrochloride lacquer where the maximum strength of Amorolfine in lacquer does not exceed 5 per cent by weight in volume;</li> <li>(iii) Amoxicillin;</li> <li>(iv) Co-Codamol;</li> <li>(v) Co-Codamol;</li> <li>(v) Co-Codamol;</li> <li>(v) Co-dydramol 10/500 tablets;</li> <li>(vi) Codeine Phosphate;</li> <li>(vii) Erythromycin</li> <li>(viii) Flucloxacillini</li> <li>(ix) Silver Sulfadiazine;</li> <li>(x) Tioconazole 28%;</li> <li>(xi) Topical hydrocortison where the maximum strength of hydrocortison in the medicinal product does not exceed l per cent by weight in weight.]</li> </ul>	e
6. [ <sup>F286</sup> Registered optometrists]	<ul><li>6. Prescription only medicines</li><li>listed in column 2 of paragraph</li><li>5.</li></ul>	
[ <sup>F287</sup> 6A Persons lawfully	Homotropine hydrobromide	The sale or supply shall be subject to the presentation
conducting a retail pharmacy	Ketotifen	subject to the presentation

Column 1	Column 2	Column 3			
Persons exempted	Prescription only medicines to which the exemption applies	Conditions			
business within the meaning of section 69.		of an order signed by an additional supply optometrist.			
	Lodoxamide				
	Nedocromil sodium				
	Olopatadine				
	Pilocarpine hydrochloride				
	[ <sup>F288</sup> Pilocarpine nitrate]				
	Polymyxin B/bacitracin				
	Polymyxin B/trimethoprim				
	Sodium cromoglycate.				
<b>6B</b> Additional supply optometrists.	Prescription only medicines specified in column 2 of paragraph 6A.	The sale or supply shall be only— (a) in the course of their			
		(b) in an emergency.]			
7. Persons selling or supplying prescription only medicines to the British Standards Institution.	7. All prescription only medicines.	<ul> <li>7. The sale or supply shall be- <ul> <li>(a) subject to the presentation of an order signed on behalf of the British Standards Institution stating the status of the person signing it and the amount of the prescription only medicine required, and</li> <li>(b) only for the purpose of testing containers of medicinal products or determining the standards for such containers.</li> </ul> </li> </ul>			
8. Holders of marketing authorizations, product licences or manufacturer's licences	8. Prescription only medicines referred to in the authorizations or licences.	<ul> <li>8. The sale or supply shall be only–</li> <li>(a) to a pharmacist,</li> </ul>			

licences.

Column 1 Persons exempted	Column 2 Prescription on to which the ex applies	nly medicines (	Column 3 Conditions	S
			(b) (c)	so as to enable that pharmacist to prepare an entry relating to the prescription only medicine in question in a tablet or capsule identification guide or similar publication, and of no greater quantity than is reasonably necessary for that purpose.
9. Pharmacists selling or supplying to persons to whom cyanide salts may be sold by virtue of section 3 (regulation of poisons) or section 4 (exclusion of sales by wholesale and certain other sales) of the Poisons Act 1972( <b>20</b> ) or by virtue of article 5 (prohibitions and regulations with respect to sale of poisons) or article 6 (exemption with respect to certain sales) of the Poisons (Northern Ireland) Order 1976( <b>21</b> ).	9. Amyl nitrite.	oi to av	nly be so	or supply shall far as is necessary n antidote to be persons at risk of isoning.
[ <sup>F289</sup> [ <sup>F290</sup> 10. Registered chiropodists [ <sup>F291</sup> or podiatrists] against whose names are recorded in the relevant register annotations signifying that they are qualified to use the medicines specified in column 2.]	hydr crea max of th in th not e cent weig (b) Amo	on only be proof on only pro- proof or or or of the model of the pro- model of the pr	e only in	sale or supply shall the course of their al practice <sup>F293</sup> ]

<sup>(20) 1972</sup> c. 66. (21) S.I. 1976/1214 (N.I. 23).

Column 1	Column 2	?	Column 3	
Persons exempted		ion only medicines the exemption	Conditions	
	(c) (d) (e) (f) (g) (h) (i) (j) (k)	lacquer where the maximum strength of Amorolfine in lacquer does not exceed 5 per cent by weight in volume; Amoxicillin; Co-Codamol; Co-dydramol 10/500 tablets; Codeine Phosphate; Erythromycin; Flucloxacillin; Silver Sulfadiazine; Tioconazole 28%; Topical hydrocortisone where the maximum strength of hydrocortisone in the medicinal product does not exceed 1 per cent by weight in weight.]		

	l Amendments
F281	Words in Sch. 5 Pt. 1 para. 4 substituted (1.6.2010) by The Medicines for Human Use (Miscellaneous
	Amendments) Order 2010 (S.I. 2010/1136), arts. 1(1), 3(3)(a)
F282	Words in Sch. 5 Pt. 1 para. 4 omitted (1.7.2011) by virtue of The Medicines (Miscellaneous Amendments)
	Order 2011 (S.I. 2011/1327), arts. 1(1), <b>3(2)(a)(i)</b>
F283	Words in Sch. 5 Pt. 1 para. 4 omitted (1.7.2011) by virtue of The Medicines (Miscellaneous Amendments)
	Order 2011 (S.I. 2011/1327), arts. 1(1), <b>3(2)(a)(ii)</b>
F284	Words in Sch. 5 Pt. 1 para. 5 substituted (1.7.2011) by The Medicines (Miscellaneous Amendments)
	Order 2011 (S.I. 2011/1327), arts. 1(1), <b>3(2)(b)(i)</b>
F285	Words in Sch. 5 Pt. 1 para. 5 substituted (1.7.2011) by The Medicines (Miscellaneous Amendments)
	Order 2011 (S.I. 2011/1327), arts. 1(1), <b>3(2)(b)(ii)</b>
F286	Words in Order substituted (30.6.2005 as notified in the London Gazette dated 3.6.2005) by The
	Opticians Act 1989 (Amendment) Order 2005 (S.I. 2005/848), Sch. 1 para. 27(b)
F287	Sch. 5 Pt. 1 paras. 6A, 6B inserted (30.6.2005) by The Medicines for Human Use (Prescribing)
	(Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), 7(1)(b)
F288	Words in Sch. 5 Pt. 1 Table inserted (6.1.2006) by The Medicines for Human Use (Prescribing)
	(Miscellaneous Amendments) (No. 2) Order 2005 (S.I. 2005/3324), arts. 1(1), 3
F289	Words in Sch. 5 Pt. 1 para. 10 substituted (17.11.2006) by The Medicines for Human Use (Administration
	and Sale or Supply) (Miscellaneous Amendments) Order 2006 (S.I. 2006/2807), arts. 1(1), 4(2)(a)

- F290 Sch. 5 Pt. 1 para. 10 inserted (13.2.1998) by The Prescription Only Medicines (Human Use) Amendment Order 1998 (S.I. 1998/108), art. 1, Sch.
- **F291** Words in Sch. 5 Pt. 1 para. 10 inserted (1.7.2011) by The Medicines (Miscellaneous Amendments) Order 2011 (S.I. 2011/1327), arts. 1(1), **3(2)(c)(i)**
- **F292** Words in Sch. 5 Pt. 1 para. 10 substituted (1.7.2011) by The Medicines (Miscellaneous Amendments) Order 2011 (S.I. 2011/1327), arts. 1(1), **3(2)(c)(ii)**
- **F293** Words in Sch. 5 Pt. 1 para. 10 omitted (1.7.2011) by virtue of The Medicines (Miscellaneous Amendments) Order 2011 (S.I. 2011/1327), arts. 1(1), **3(2)(c)(iii)**

Article 11(1)(b)

# PART II

## EXEMPTIONS FROM THE RESTRICTION ON SUPPLY

Column 1	Column 2	Column 3
Persons exempted	Prescription only medicines to which the exemption applies	Conditions
1. Royal National Lifeboat Institution and certified first aiders of the Institution.	1. All prescription only medicines.	1. The supply shall be only so far as is necessary for the the treatment of sick or injured persons in the exercise of the functions of the Institution.
2. The owner or the master of a ship which does not carry a doctor on board as part of her complement.	2. All prescription only medicines.	2. The supply shall be only so far as is necessary for the treatment of persons on the ship.
3. Persons authorized by licences granted under regulation 5 of the Misuse of Drugs Regulations to supply a controlled drug.	3. Such prescription only medicines, being controlled drugs, as are specified in the licence.	3. The supply shall be subject to such conditions and in such circumstances and to such an extent as may be specified in the licence.
[ <sup>F294</sup> 3A. Persons employed or engaged in the provision of lawful drug treatment services.	3A. Ampoules of sterile water for injection containing not more than 2 ml of sterile water.	3A. The supply shall be only in the course of provision of lawful drug treatment services.]
4. Persons requiring prescription only medicines for the purpose of enabling them, in the course of any business carried on by them, to comply with any requirements made by or in pursuance of any enactment with respect to the medical treatment of their employees.	4. Such prescription only medicines as may be specified in the relevant enactment.	<ul> <li>4. The supply shall be–</li> <li>(a) for the purpose of enabling them to comply with any requirements made by or in pursuance of any such enactment, and</li> <li>(b) subject to such conditions and in such circumstances</li> </ul>

as may be specified

Column 1 Persons exempted	Column 2 Prescription only medicines to which the exemption applies	Column 3 Conditions in the relevant
		enactment.
5. Persons operating an occupational health scheme.	5. Prescription only medicines sold or supplied to a person operating an occupational health scheme in response to an order in writing signed by a	5. — (1) The supply shall be in the course of an occupational health scheme.
	doctor or a registered nurse.	<ul> <li>(2) The individual supplying the prescription only medicine, if not a doctor, shall be a registered nurse acting in accordance with the written instructions of a doctor as to the circumstances in which prescription only medicines of the description in question are to be used in the course of the occupational health scheme.</li> </ul>
6. The operator or commander of an aircraft.	6. Prescription only medicines which are not for parenteral administration and which have been sold or supplied to the operator or commander of the aircraft in response to an order in writing signed by a doctor.	6. The supply shall be only so far as is necessary for the immediate treatment of sick or injured persons on the aircraft and shall be in accordance with the written instructions of a doctor as to the circumstances in which prescription only medicines of the description in question are to be used on the aircraft.
7. Persons employed as qualified first-aid personnel on offshore installations.	7. All prescription only medicines.	7. The supply shall be only so far as is necessary for the treatment of persons on the installation.
[ <sup>F295</sup> 8. Persons who hold a certificate in first aid from the Mountain Rescue Council of England and Wales, or from the Northern Ireland Mountain	8 Prescription only medicines supplied to a person specified in column 1 in response to an order in writing signed by a doctor.	8 The supply shall be only so far as is necessary for the treatment of sick or injured persons in the course of providing mountain rescue services.]

Column 1	Column 2	Column 3
Persons exempted	Prescription only medicines to which the exemption applies	Conditions
Rescue Co-ordinating Committee.		
[ <sup>F296</sup> 9. Persons ("P") who are members of Her Majesty's armed forces.	9. All prescription only medicines.	<ul> <li>9. The supply shall be—</li> <li>(a) in the course of P undertaking any function as a member of Her Majesty's armed forces; and</li> </ul>
		(b) where P is satisfied that it is not practicable for another person who is legally entitled to supply a prescription only medicine to do so; and
		<ul> <li>(c) only in so far as is necessary— <ul> <li>(i) for the treatment of a sick or injured person in a medica emergency, or</li> <li>(ii) to prevent ill-health where there is a risk that a person would suffe ill-health if the prescription only medicine is not supplied.]</li> </ul> </li> </ul>

#### **Textual Amendments**

- F294 Sch. 5 Pt. 2 para. 3A inserted (30.6.2005) by The Medicines for Human Use (Prescribing) (Miscellaneous Amendments) Order 2005 (S.I. 2005/1507), arts. 1(1), 7(2)
- F295 Sch. 5 Pt. 2 para. 8 inserted (17.11.2006) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2006 (S.I. 2006/2807), arts. 1(1), 4(3)
- **F296** Sch. 5 Pt. 3 para. 10 added (21.12.2009) by The Medicines (Exemptions and Miscellaneous Amendments) Order 2009 (S.I. 2009/3062), arts. 1(1), **3(2)**

Article 11(2)

# PART III

# EXEMPTIONS FROM RESTRICTION ON ADMINISTRATION Column 1

Column 1 Persons exempted	Column 2 Prescription only medicines to which the exemption applies	Column 3 Conditions
[ <sup>F297</sup> 1. Registered chiropodists [ <sup>F298</sup> or podiatrists] against whose names are recorded in the relevant register annotations signifying that they are qualified to use the medicines specified in column 2]	<ol> <li>Prescription only medicines for parenteral administration that contain<sup>F299</sup></li> <li>[<sup>F300</sup>Adrenaline]</li> <li>[<sup>F301</sup>Bupivacaine hydrochloride Bupivacaine hydrochloride with adrenaline where the maximum strength of the adrenaline does not exceed 1 mg in 200 ml of bupivacaine hydrochloride</li> <li>[<sup>F300</sup>Levobupivacaine hydrochloride]</li> <li>Lignocaine hydrochloride</li> <li>Lignocaine hydrochloride with adrenaline where the maximum strength of the adrenaline does not exceed 1 mg in 200 ml of lignocaine hydrochloride</li> <li>Lignocaine hydrochloride with adrenaline does not exceed 1 mg in 200 ml of lignocaine hydrochloride</li> <li>[<sup>F300</sup>Methylpredniso Prilocaine hydrochloride.]</li> <li>[<sup>F300</sup>Ropivacaine hydrochloride.]</li> </ol>	16
2. Registered midwives [ <sup>F304</sup> and student midwives].	2. Prescription only medicines for parenteral administration containing any of the following substances but no other substance specified in	<ul> <li>[<sup>F308</sup>The medicine shall—         <ul> <li>(a) in the case of</li> <li>Lignocaine,</li> <li>Lignocaine</li> <li>hydrochloride</li> <li>and Promazine</li> <li>hydrochloride, be</li> </ul> </li> </ul>

Column 1	Column 2	Column 3
Persons exempted	Prescription only medicines to which the exemption applies	Conditions
	column 1 of Schedule 1 to this Order– [ <sup>F305</sup> Adrenaline Anti-D immunoglobulin Carboprost [ <sup>F306</sup> cyclizine lactate] Diamorphine Ergometrine maleate Gelofusine <sup>F307</sup> Hartmann's solution	administered only while attending on a woman in childbirth, and (b) where administration is— (i) by a registered midwife, be administered in the course of their professional practice; (ii) by a student
	Hepatitis B vaccine Hepatitis immunoglobulin Lidocaine Lidocaine hydrochloride Morphine Naloxone hydrochloride Oxytocins, natural and synthetic Pethidine hydrochloride Phytomenadione Prochloperazine Sodium chloride 0.9%.]	midwife— (aa) be administered under the direct supervision of a registered midwife; and (bb) not include Diamorphine or Pethidine hydrochloride
3. Persons who are authorized as members of a group by a group authority granted under regulations 8(3) or 9(3) of the Misuse of Drugs Regulations to supply a controlled drug by way of administration only.	3. Prescription only medicines that are specified in the group authority.	3. The administration shall be subject to such conditions and in such circumstances and to such extent as may be specified in the group authority.
4. The owner or master of a ship which does not carry a doctor on board as part of her complement.	4. All prescription only medicines that are for parenteral administration.	4. The administration shall be only so far as is necessary for the treatment of persons on the ship.
5. Persons operating an occupational health scheme.	5. Prescription only medicines for parenteral administration sold or supplied to the person	5. — (1) The administration shall be in the

Column 1 Persons exempted	Column 2 Prescription only medicines to which the exemption applies	Column 3 Conditions
	operating an occupational health scheme in response to an order in writing signed by a doctor or a registered nurse.	<ul> <li>course of an occupational health scheme.</li> <li>(2) The individual administering the prescription only medicine, if neither a doctor nor acting in accordance with the directions of a doctor, shall be a registered nurse acting in accordance with the written instructions of a doctor as to the circumstances in which prescription only medicines of the description in question are to be used in the course of the occupational health scheme.</li> </ul>
6. The operator or commander of an aircraft.	6. Prescription only medicines for parenteral administration which have been sold or supplied to the operator or commander of the aircraft in response to an order in writing signed by a doctor.	6. The administration shall be only so far as is necessary for the immediate treatment of sick or injured persons on the aircraft and shall be in accordance with the written instructions of a doctor as to the circumstances in which prescription only medicines of the description in question are to be used on the aircraft.
7. Persons who are, and at 11th February 1982 were, persons customarily administering medicinal products to human beings by parenteral administration in the course of a business in the field of osteopathy, naturopathy, acupuncture or other similar field except chiropody.	7. Medicinal products that are prescription only medicines by reason only that they fall within the class specified in article 3(c) (products for parenteral administration).	7. The person administering the prescription only medicine shall have been requested by or on behalf of the person to whom it is administered and in that person's presence to use his own judgement as to the treatment required.

Column 1 Persons exempted	Column 2 Prescription only medicines to which the exemption applies	Column 3 Conditions
8. Persons employed as qualified first-aid personnel on offshore installations.	8. All prescription only medicines that are for parenteral administration.	8. The administration shall be only so far as is necessary for the treatment of persons on the installation.
9. Persons who hold a certificate of proficiency in ambulance paramedic skills issued by, or with the approval of, the Secretary of State [ <sup>F309</sup> or persons who are [ <sup>F310</sup> registered] paramedics].	<ul> <li>9. The following prescription only medicines for parenteral administration– <ul> <li>(a) Diazepam 5 mg per ml emulsion for injection;</li> <li>(b) Succinylated Modified Fluid Gelatin 4 per cent intravenous infusion;</li> <li>(bb) [<sup>F311</sup>medicines containing the substances Ergometrine Maleate 500mcg per ml with Oxytocin 5 iu per ml, but no other active ingredient]</li> <li>(d) prescription only medicines containing one or more of the following substances, but no active ingredient– Adrenaline Acid Tartrate [<sup>F312</sup>Adrenaline Hydrochloride [<sup>F313</sup>Amiodaron Anhydrous Glucose [<sup>F314</sup>Benzylpen [<sup>F315</sup>Bretylium Tosylate] Compound Sodium Lactate Intravenous Infusion</li> </ul></li></ul>	] ne]

Column 1	Column 2	Column 3
Persons exempted	Prescription only med to which the exemptio applies	
	(Hart Solut Ergor Malea [ <sup>F314</sup> F Glucc Hepa Sodiu Lignc Hydra [ <sup>F314</sup> M Sulph Nalbu Hydra Naloy Hydra Polyg [ <sup>F316</sup> R ] Sodiu Bicar Sodiu Chlor	netrine tte rusemide] se rusemide] se rin m caine ochloride letoclopramide] lorphine ate] phine ochloride eline eteplase m ponate m
[ <sup>F296</sup> 10 . Persons ("P") who are members of Her Majesty's armed forces.	10. All prescription on medicines.	<ul> <li>ly 10. The administration shall be— <ul> <li>(a) in the course of P</li> <li>undertaking any function as a member of Her</li> <li>Majesty's armed forces; and</li> </ul> </li> <li>(b) where P is satisfied that it is not practicable for another person who is legally entitled to administer a prescription only medicine to do so; and</li> <li>(c) only in so far as is necessary— <ul> <li>(i) for the treatment of a sick or injured person in a medical emergency, or</li> </ul> </li> </ul>

	1 I	Column 2	Column	3
Person	s exempted	Prescription only medic to which the exemption applies	cines Conditio	ns
			(ii)	to prevent ill- health where there is a risk that a person would suffe ill-health if the prescription only medicine is not administered.]
F297		para. 1 substituted (17.11.2006) by T		
F298	Words in Sch. 5 Pt. 3	Miscellaneous Amendments) Order para. 1 inserted (1.7.2011) by The M		
F299	Words in Sch. 5 Pt. 3 p	), arts. 1(1), <b>3(3)(a)(i)</b> para. 1 omitted (1.7.2011) by virtue of 1/1327), arts. 1(1), <b>3(3)(a)(ii)</b>	f The Medicines (Mise	cellaneous Amendments
F300	Words in Sch. 5 Pt. 3	b para. 1 inserted (17.11.2006) by Tl Miscellaneous Amendments) Order		
F301	Words in Sch. 5 Pt.	3 substituted (13.2.1998) by The		
F302	Amendment Order 1998 (S.I. 1998/108), arts. 1, <b>5(2)</b> Words in Sch. 5 Pt. 3 para. 1 inserted (16.9.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 3) Order 1998 (S.I. 1998/2081), arts. 1(1), <b>4(b)</b>			
		O(1001 1770 (0.1, 1770/2001), d(0.1))		
		para. 1 inserted (1.7.2011) by The M	Iedicines (Miscellane	ous Amendments) Orde
F304	2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3			
F304 F305	2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3	para. 1 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(a)(iii)</b> para. 2 inserted (1.7.2011) by The M	fedicines (Miscellane) The Medicines for Hu	ous Amendments) Orde
F304 F305 F306	2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 Amendments) Order Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011	para. 1 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(a)(iii)</b> para. 2 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(b)(i)</b> g para. 2 substituted (1.6.2010) by T 2010 (S.I. 2010/1136), arts. 1(1), <b>3(3)</b> g para. 2 substituted (1.7.2011) by T 1/1327), arts. 1(1), <b>3(3)(b)(ii)(aa)</b>	Medicines (Miscellane The Medicines for Hu <b>3)(b)</b> The Medicines (Misc	ous Amendments) Orde man Use (Miscellaneou cellaneous Amendments
F304 F305 F306 F307	2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 Amendments) Order Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Word in Sch. 5 Pt. 3 p Order 2011 (S.I. 2011)	para. 1 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(a)(iii)</b> para. 2 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(b)(i)</b> gara. 2 substituted (1.6.2010) by T 2010 (S.I. 2010/1136), arts. 1(1), <b>3(</b> 3 para. 2 substituted (1.7.2011) by T 1/1327), arts. 1(1), <b>3(3)(b)(ii)(aa)</b> ara. 2 omitted (1.7.2011) by virtue of 1/1327), arts. 1(1), <b>3(3)(b)(ii)(bb)</b>	Aedicines (Miscellane The Medicines for Hu <b>3)(b)</b> The Medicines (Misc The Medicines (Misc	ous Amendments) Orde man Use (Miscellaneou cellaneous Amendments cellaneous Amendments
F304 F305 F306 F307 F308	2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 Amendments) Order Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Word in Sch. 5 Pt. 3 p Order 2011 (S.I. 2011) Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011)	para. 1 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(a)(iii)</b> para. 2 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(b)(i)</b> 8 para. 2 substituted (1.6.2010) by T 2010 (S.I. 2010/1136), arts. 1(1), <b>3(3)</b> 9 para. 2 substituted (1.7.2011) by T 1/1327), arts. 1(1), <b>3(3)(b)(ii)(a)</b> ara. 2 omitted (1.7.2011) by virtue of 1/1327), arts. 1(1), <b>3(3)(b)(ii)(bb)</b> 3 para. 2 substituted (1.7.2011) by T 1/1327), arts. 1(1), <b>3(3)(b)(ii)(bb)</b> 3 para. 2 substituted (1.7.2011) by T	Aedicines (Miscellane The Medicines for Hu <b>3)(b)</b> The Medicines (Misc The Medicines (Misc The Medicines (Misc	ous Amendments) Orde man Use (Miscellaneou cellaneous Amendments cellaneous Amendments
F304 F305 F306 F307 F308 F309	2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 Amendments) Order Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Word in Sch. 5 Pt. 3 p Order 2011 (S.I. 2011) Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Words in Sch. 5 Pt. 3 Amendment (No. 2) (	para. 1 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(a)(iii)</b> para. 2 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(b)(i)</b> gara. 2 substituted (1.6.2010) by T 2010 (S.I. 2010/1136), arts. 1(1), <b>3(</b> 3 para. 2 substituted (1.7.2011) by T 1/1327), arts. 1(1), <b>3(3)(b)(ii)(aa)</b> ara. 2 omitted (1.7.2011) by virtue of 1/1327), arts. 1(1), <b>3(3)(b)(ii)(bb)</b> 3 para. 2 substituted (1.7.2011) by T 1/1327), arts. 1(1), <b>3(3)(b)(ii)(bb)</b> 3 para. 2 substituted (1.7.2011) by T 1/1327), arts. 1(1), <b>3(3)(b)(iii)</b> 3 para. 9 added (16.11.2000) by Th Order 2000 (S.I. 2000/2899), arts. 1(1)	Aedicines (Miscellane the Medicines for Hu <b>3)(b)</b> The Medicines (Misc The Medicines (Misc The Medicines (Misc the Prescription Only (1), <b>5(a)</b>	ous Amendments) Orde man Use (Miscellaneou cellaneous Amendments cellaneous Amendments cellaneous Amendments Medicines (Human Use
F304 F305 F306 F307 F308 F309 F310	2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 Amendments) Order Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Word in Sch. 5 Pt. 3 p Order 2011 (S.I. 2011) Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Words in Sch. 5 Pt. 3 Amendment (No. 2) ( Word in Sch. 5 Pt. 3 (Consequential Amer	para. 1 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(a)(iii)</b> para. 2 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(b)(i)</b> gara. 2 substituted (1.6.2010) by T 2010 (S.I. 2010/1136), arts. 1(1), <b>3(</b> 3 para. 2 substituted (1.7.2011) by 1/1327), arts. 1(1), <b>3(3)(b)(ii)(aa)</b> ara. 2 omitted (1.7.2011) by virtue of 1/1327), arts. 1(1), <b>3(3)(b)(ii)(bb)</b> 3 para. 2 substituted (1.7.2011) by 1/1327), arts. 1(1), <b>3(3)(b)(ii)(bb)</b> 3 para. 2 substituted (1.7.2011) by 1/1327), arts. 1(1), <b>3(3)(b)(ii)</b> 3 para. 9 added (16.11.2000) by Th Order 2000 (S.I. 2000/2899), arts. 1( para. 9 substituted for words (9.7.2 adments) Order 2003 (S.I. 2003/1590	Aedicines (Miscellane The Medicines for Hu <b>3)(b)</b> The Medicines (Misc The Medicines (Misc The Medicines (Misc the Prescription Only (1), <b>5(a)</b> 2003) by The Health 0), art. 1, <b>Sch. para.</b>	ous Amendments) Orde man Use (Miscellaneou cellaneous Amendments cellaneous Amendments cellaneous Amendments Medicines (Human Use Professions Order 200 <b>21(3)(b)(ii)</b>
F304 F305 F306 F307 F308 F309 F310 F311	2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 Amendments) Order Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Word in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Words in Sch. 5 Pt. 3 Amendment (No. 2) ( Word in Sch. 5 Pt. 3 Amendment (No. 2) (	para. 1 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(a)(iii)</b> para. 2 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(b)(i)</b> gara. 2 substituted (1.6.2010) by T 2010 (S.I. 2010/1136), arts. 1(1), <b>3(3)</b> gara. 2 substituted (1.7.2011) by ' 1/1327), arts. 1(1), <b>3(3)(b)(ii)(aa)</b> ara. 2 omitted (1.7.2011) by virtue of 1/1327), arts. 1(1), <b>3(3)(b)(ii)(bb)</b> 3 para. 2 substituted (1.7.2011) by ' 1/1327), arts. 1(1), <b>3(3)(b)(ii)(bb)</b> 3 para. 2 substituted (1.7.2011) by ' 1/1327), arts. 1(1), <b>3(3)(b)(iii)(bb)</b> 3 para. 9 added (16.11.2000) by Th Order 2000 (S.I. 2000/2899), arts. 1( para. 9 inserted (16.11.2000) by Th Order 2000 (S.I. 2000/2899), arts. 1( Order 2000 (S.I. 2000/2899), arts. 1(	Aedicines (Miscellane The Medicines for Hu <b>3)(b)</b> The Medicines (Misc The Medicines (Misc The Medicines (Misc The Medicines (Misc the Prescription Only (1), <b>5(a)</b> 2003) by The Health 0), art. 1, <b>Sch. para.</b> 7 the Prescription Only (1), <b>5(b)</b>	ous Amendments) Orde man Use (Miscellaneou cellaneous Amendments cellaneous Amendments cellaneous Amendments Medicines (Human Use Professions Order 200 21(3)(b)(ii) Medicines (Human Use
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F304 F305 F306 F307 F308 F309 F310 F311 F312 F313	2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 2011 (S.I. 2011/1327) Words in Sch. 5 Pt. 3 Amendments) Order Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Word in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Words in Sch. 5 Pt. 3 Order 2011 (S.I. 2011) Words in Sch. 5 Pt. 3 Amendment (No. 2) ( Word in Sch. 5 Pt. 3 Amendment (No. 2) ( Words in Sch. 5 Pt. 3 Amendment Order 20 Word in Sch. 5 Pt. 3 Amendment (No. 3) (	para. 1 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(a)(iii)</b> para. 2 inserted (1.7.2011) by The M ), arts. 1(1), <b>3(3)(b)(i)</b> 8 para. 2 substituted (1.6.2010) by T 2010 (S.I. 2010/1136), arts. 1(1), <b>3(</b> 3 para. 2 substituted (1.7.2011) by T 1/1327), arts. 1(1), <b>3(3)(b)(ii)(aa)</b> ara. 2 omitted (1.7.2011) by virtue of 1/1327), arts. 1(1), <b>3(3)(b)(ii)(bb)</b> 3 para. 2 substituted (1.7.2011) by T 1/1327), arts. 1(1), <b>3(3)(b)(ii)(bb)</b> 3 para. 2 substituted (1.7.2011) by T 1/1327), arts. 1(1), <b>3(3)(b)(iii)(bb)</b> 3 para. 9 added (16.11.2000) by Th Order 2000 (S.I. 2000/2899), arts. 1(1) adments) Order 2003 (S.I. 2003/1590) 6 para. 9 inserted (16.11.2000) by T Order 2000 (S.I. 2000/2899), arts. 1(1) 3 para. 9 added (17.1.2011) by Th	Aedicines (Miscellane The Medicines for Hu <b>3)(b)</b> The Medicines (Misc The Medicines (Misc The Medicines (Misc The Medicines (Misc the Prescription Only (1), <b>5(a)</b> 2003) by The Health 0), art. 1, <b>Sch. para.</b> The Prescription Only (1), <b>5(b)</b> the Prescription Only The Prescription Only (1), (1), <b>6</b>	ous Amendments) Orde man Use (Miscellaneou cellaneous Amendments cellaneous Amendments cellaneous Amendments Medicines (Human Use Medicines (Human Use Medicines (Human Use Medicines (Human Use

F315 Words in Sch. 5 Pt. 3 para. 9 inserted (1.6.1998) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 1998 (S.I. 1998/1178), arts. 1(1), 3

F316 Words in Sch. 5 Pt. 3 para. 9 inserted (18.5.2004) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2004 (S.I. 2004/1189), arts. 1(1), 4

### SCHEDULE 6

Article 16(1)

### ORDERS REVOKED

Column 1	Column 2
Orders	References
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Order 1983	S.I. 1983/1212
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment Order 1984	S.I. 1984/756
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment Order 1986	S.I. 1986/586
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment Order 1987	S.I. 1987/674
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment (No. 2) Order 1987	S.I. 1987/1250
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment Order 1988	S.I. 1988/2017
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment Order 1991	S.I. 1991/962
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment Order 1992	S.I. 1992/1534
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment (No. 2) Order 1992	S.I. 1992/2937
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment Order 1993	S.I. 1993/1890
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment (No. 2) Order 1993	S.I. 1993/3256

Column 1	Column 2
Orders	References
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment Order 1994	S.I. 1994/558
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment (No. 2) Order 1994	S.I. 1994/3016
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment (No. 3) Order 1994	S.I. 1994/3050
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment Order 1995	S.I. 1995/1384
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment (No. 2) Order 1995	S.I. 1995/3174
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment Order 1996	S.I. 1996/1514
The Medicines (Products other than Veterinary Drugs) (Prescription Only) Amendment (No. 2) Order 1996	S.I. 1996/3193

### [<sup>F317</sup>SCHEDULE 7

Articles 12A to 12C

### **Textual Amendments**

F317 Sch. 7 inserted (9.8.2000) by The Prescription Only Medicines (Human Use) Amendment Order 2000 (S.I. 2000/1917), arts. 1(1), 2(e)

# PART I

### PARTICULARS TO BE INCLUDED IN A PATIENT GROUP DIRECTION

- (a) the period during which the Direction shall have effect;
- (b) the description or class of prescription only medicine to which the Direction relates;
- (c) whether there are any restrictions on the quantity of medicine which may be [<sup>F318</sup>sold or] supplied on any one occasion, and, if so, what restrictions;
- (d) the clinical situations which prescription only medicines of that description or class may be used to treat;
- (e) the clinical criteria under which a person shall be eligible for treatment;

- (f) whether any class of person is excluded from treatment under the Direction and, if so, what class of person;
- (g) whether there are circumstances in which further advice should be sought from a doctor or dentist and, if so, what circumstances;
- (h) the pharmaceutical form or forms in which prescription only medicines of that description or class are to be administered;
- (i) the strength, or maximum strength, at which prescription only medicines of that description or class are to be administered;
- (j) the applicable dosage or maximum dosage;
- (k) the route of administration;
- (l) the frequency of administration;
- (m) any minimum or maximum period of administration applicable to prescription only medicines of that description or class;
- (n) whether there are any relevant warnings to note, and, if so, what warnings;
- (o) whether there is any follow up action to be taken in any circumstances, and, if so, what action and in what circumstances;
- (p) arrangements for referral for medical advice;
- (q) details of the records to be kept of the supply, or the administration, of medicines under the Direction.

#### **Textual Amendments**

F318 Words in Sch. 7 Pt. 1 inserted (1.9.2007) by The Medicines for Human Use (Administration and Sale or Supply) (Miscellaneous Amendments) Order 2007 (S.I. 2007/2178), arts. 1(1), 5

# PART II

### PERSONS ON WHOSE BEHALF A PATIENT GROUP DIRECTION MUST BE SIGNED

Column 1	Column 2
Class of person by whom a prescription only medicine is supplied or administered	Person on whose behalf the Direction must be signed
Common Services Agency	The Agency
[ <sup>F319</sup> Strategic Health Authority]	[ <sup>F319</sup> The Strategic Health Authority]
Health Authority	The Health Authority
Special Health Authority	The Special Health Authority
NHS trust [F320 or NHS foundation trust]	The trust
Primary Care Trust	The Trust
A person who supplies or administers a prescription only medicine pursuant to an arrangement made with the Common Services Agency, a Health Authority, a Special Health	The Common Services Agency, where the arrangement has been made with the Agency; or the Health Authority, Special Health Authority, NHS trust [ <sup>F322</sup> or NHS foundation

Column 1	Column 2
Class of person by whom a prescription only medicine is supplied or administered	Person on whose behalf the Direction must be signed
Authority, an NHS trust, [ <sup>F321</sup> , an NHS foundation trust] or a Primary Care Trust	trust] or Primary Care Trust with which the arrangement has been made

### **Textual Amendments**

- F319 Words in Sch. 7 Pt. 2 inserted (1.10.2002) by The National Health Service Reform and Health Care Professions Act 2002 (Supplementary, Consequential etc. Provisions) Regulations 2002 (S.I. 2002/2469), reg. 1, Sch. 1 para. 73(4)
- **F320** Words in Sch. 7 Pt. 2 inserted (1.4.2004) by The Health and Social Care (Community Health and Standards) Act 2003 (Supplementary and Consequential Provision) (NHS Foundation Trusts) Order 2004 (S.I. 2004/696), arts. 1(1)(b), 3(2), Sch. 2
- **F321** Words in Sch. 7 Pt. 2 inserted (1.4.2004) by The Health and Social Care (Community Health and Standards) Act 2003 (Supplementary and Consequential Provision) (NHS Foundation Trusts) Order 2004 (S.I. 2004/696), arts. 1(1)(b), 3(4), Sch. 4
- **F322** Words in Sch. 7 Pt. 2 inserted (1.4.2004) by The Health and Social Care (Community Health and Standards) Act 2003 (Supplementary and Consequential Provision) (NHS Foundation Trusts) Order 2004 (S.I. 2004/696), arts. 1(1)(b), 3(3), Sch. 3

# [<sup>F323</sup>PART IIA

# PERSONS BY WHOM OR ON WHOSE BEHALF A PATIENT GROUP DIRECTION USED IN THE PROVISION OF HEALTH CARE BY OR ON BEHALF OF THE POLICE, THE PRISON SERVICES OR THE ARMED FORCES MUST BE SIGNED

<ul> <li>Textual Amendments</li> <li>F323 Sch. 7 Pt. 2A inserted (4.4.2003) by The Prescription Only Medicines (Human Use) Amendment Order 2003 (S.I. 2003/696), arts. 1(1), 17</li> </ul>	
Column 1	Column 2
Force or service by whom or on whose behalf the health care is provided	Person by whom or on whose behalf the Direction must be signed
A police force in England or Wales	The chief officer of police for that police force (within the meaning of the Police Act 1996)
A police force in Scotland	The chief constable of that police force (within the meaning of the Police (Scotland) Act 1997)
The Police Service of Northern Ireland	The Chief Constable of the Police Service of Northern Ireland

The prison service in England and Wales

The governor of the prison in relation to which the health care in question is being provided

Column 1	Column 2
Force or service by whom or on whose behalf the health care is provided	Person by whom or on whose behalf the Direction must be signed
The prison service in Scotland	The Scottish Prison Service Management Board
The prison service in Northern Ireland	The Northern Ireland Prison Service Management Board
Her Majesty's Forces	(i) the Surgeon General,
	(ii) a Medical Director General, or
	<ul><li>(iii) a chief executive of an executive agency of the Ministry of Defence]</li></ul>

# PART III

# CLASSES OF INDIVIDUAL [<sup>F324</sup>BY WHOM PRESCRIPTION ONLY MEDICINES MAY BE SUPPLIED OR ADMINISTERED]

### **Textual Amendments**

**F324** Words in Sch. 7 Pt. 3 heading substituted (16.11.2000) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2000 (S.I. 2000/2899), arts. 1(1), **3(2)** 

[<sup>F325</sup>Dental hygienist.

Dental therapist.]

[<sup>F326</sup>Registered paramedics or individuals who hold a certificate of proficiency in ambulance paramedic skills issued by, or with the approval of, the Secretary of State.]

Pharmacists.

F327

Registered midwives.

Registered nurses.

[F328Registered optometrists]

[<sup>F329</sup>Registered] chiropodists.

[<sup>F330</sup>Registered orthoptists.

Registered physiotherapists.

Registered radiographers.]

[F331"Registered dietitians.";]

[<sup>F331</sup>"Registered occupational therapists.";]

[<sup>F331</sup>"Registered orthotists and prosthetists."; and]

[<sup>F331</sup>"Registered speech and language therapists."]]

#### **Textual Amendments**

- F325 Words in Sch. 7 Pt. 3 inserted (1.6.2010) by The Medicines for Human Use (Miscellaneous Amendments) Order 2010 (S.I. 2010/1136), arts. 1(1), 3(4)
- F326 Words in Sch. 7 Pt. 3 substituted (9.7.2003) by The Health Professions Order 2001 (Consequential Amendments) Order 2003 (S.I. 2003/1590), art. 1, Sch. para. 21(4)(a)
- F327 Words in Sch. 7 Pt. 3 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. 39(b)
- F328 Words in Order substituted (30.6.2005 as notified in the London Gazette dated 3.6.2005) by The Opticians Act 1989 (Amendment) Order 2005 (S.I. 2005/848), Sch. 1 para. 27(b)
- F329 Word in Sch. 7 Pt. 3 substituted (9.7.2003) by The Health Professions Order 2001 (Consequential Amendments) Order 2003 (S.I. 2003/1590), art. 1, Sch. para. 21(4)(b)
- F330 Words in Sch. 7 Pt. 3 substituted (9.7.2003) by The Health Professions Order 2001 (Consequential Amendments) Order 2003 (S.I. 2003/1590), art. 1, Sch. para. 21(4)(c)
- F331 Words in Sch. 7 Pt. 3 added (18.5.2004) by The Prescription Only Medicines (Human Use) Amendment (No. 2) Order 2004 (S.I. 2004/1189), arts. 1(1), 5

### **EXPLANATORY NOTE**

### (This note is not part of the Order)

This Order consolidates with amendments the Medicines (Products Other Than Veterinary Drugs) (Prescription Only) Order 1983 as amended. That Order and the Orders amending it ("the 1983 Order as amended") are revoked by article 16 and Schedule 6.

This Order specifies the descriptions and classes of prescription only medicines (i.e. medicinal products which, subject to exemptions, may be sold or supplied by retail only in accordance with a prescription given by an appropriate practitioner and which may be administered only by or in accordance with the directions of such a practitioner). Many medicinal products are included in a class of such medicines by reason of the substances contained in them (*see*Schedule 1) but others are included because of other criteria, such as their method of administration (*see*article 3). In many cases the provisions of the Act apply subject to exemptions (*see*articles 4 and 5 to 13 and Schedule 1).

The principal amendments relate to those medicines in respect of which marketing authorizations have been granted by the European Community. They include as prescription only medicines those medicines in respect of which such an authorization has been granted which classifies a medicine as being subject to medical prescription (article 3(f)). They exclude from the class of prescription only medicines those medicines in respect of which such an authorization provides for supply which is not subject to medical prescription (article 6(3)).

The differences between this Order and the 1983 Order as amended are in the main technical changes concerning the location of provisions such as the division of material in Schedule 1 to the 1983 Order as amended between the new Schedules 1 and 2. But within the new Schedule 1 there are changes which relate to-

(a) the deletion from Column 1 of substances which are no longer used in those medicinal products which are on the market;

- (b) the use of current names for the substances which are specified in that Column where their names have changed;
- (c) the incorporation in that Schedule of provisions from article 4 of, and Part IV of Schedule 1 to, the 1983 Order as amended so that they may be found more easily;
- (d) a change in the legal base for the entries in Columns 2 to 4 so that those entries now form the criteria for exemptions from the sale or supply requirements of section 58(2) of the Medicines Act 1968 instead of the criteria for excluding medicinal products from the class of prescription only medicines (*see also*article 5);
- (e) the introduction of a fifth Column which specifies the maximum pack sizes to which exemptions apply.

As this order will impose no additional costs to business a compliance cost assessment has not been prepared.

# Status:

Point in time view as at 01/08/2012.

### Changes to legislation:

There are currently no known outstanding effects for the The Prescription Only Medicines (Human Use) Order 1997.