

MEDICINES (PRESCRIPTION ONLY) (JERSEY) ORDER 1997

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MEDICINES (PRESCRIPTION ONLY) (JERSEY) ORDER 1997

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MEDICINES (PRESCRIPTION ONLY) (JERSEY) ORDER 1997¹

THE HEALTH AND SOCIAL SERVICES COMMITTEE in pursuance of Articles 57 and 110 of the Medicines (Jersey) Law 1995², after consultation with the Medicines Advisory Council and having otherwise complied with Article 110 of the Law, orders as follows –

Commencement [see endnotes]

1 Interpretation

(1) In this Order, unless the context otherwise requires –

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

"controlled drug" has the same meaning as it has in Article 3 of the Misuse of Drugs (Jersey) Law 1978³;

"cyanogenetic substances" means preparations –

- (a) that are presented for sale or supply under the name of, or as containing, amygdalin, laetrile or vitamin B17; or
- (b) that contain more than 0.1% by weight of any substance having the formula α-cyanobenzyl-6-*O*-β-D-glucopyranosyl-β-D-glucopyranosiduronic acid;

"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; and
- (b) where a medicinal product is not in any such form, the quantity of the product that is used as the unit by reference to which the dose is measured;

"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; but does not mean application by means of a throat spray, throat pastille, throat lozenge, throat tablet, nasal drop, nasal spray, nasal inhalation or teething preparation;

"health prescription" means a prescription described in Article 15(2) of the Health Insurance (Jersey) Law 1967⁴;

"health record" has the same meaning as in the Data Protection (Jersey) Law 2005;

"inhaler" does not include an aerosol;

"Law" means the Medicines (Jersey) Law 19956;

"master" has the same meaning as it has in the Merchant Shipping Act 1894 of the United Kingdom;

"maximum daily dose" or "MDD" means, in relation to a substance contained in the amount of a medicinal product for internal use, the recommended maximum quantity to be taken or administered in a period of 24 hours;

"maximum dose" or "MD" means, in relation to a substance contained in the amount of a medicinal product for internal use, the recommended maximum quantity to be taken or administered at any one time;

"maximum strength" means such of the following as may be specified –

- (a) the maximum quantity of a substance by weight or volume contained in a dosage unit of a medicinal product;
- (b) the maximum number of units of activity contained in a dosage unit or a weight of a medicinal product; and
- (c) the maximum percentage of a substance contained in a medicinal product calculated in terms of weight in weight, weight in volume, volume in weight or volume in volume, as appropriate;

"medicinal product" does not include a veterinary drug;

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

"operator", in relation to an aircraft, means the person for the time being having the management of the aircraft;

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

"prescription only medicine" means a medicinal product that is specified by this Order as a prescription only medicine;

"registered ophthalmic optician" has the same meaning as it has in Article 1(1) of the Opticians (Registration) (Jersey) Law 1962⁷;

"repeatable prescription" means a prescription containing a direction that it shall or may be dispensed more than once;

"soaps" means any compounds of a fatty acid with an alkali or amine;

"state registered paramedic" means a person who is registered in the register established and maintained under section 60 and paragraph 1(a) of Schedule 3 to the Health Act 1999 of the United Kingdom;

"supplementary prescriber" means an appropriate practitioner referred to in Article 5(d), (e) or (f) whose entry on the register established and maintained under article 5 of the Nursing and Midwifery Order 2001 of the United Kingdom, as referred to in the Health Care (Registration) (Prescribed Qualifications) (Jersey) Order 2003⁸, indicates that he or she is, or may act as, only a supplementary prescriber;

"unit preparation" means a preparation (including a mother tincture) that is 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 that diluted form or (where applicable) by impregnating tablets, granules, powders or other inert substances.⁹

(2) In this Order –

- (a) in Schedules 1 and 2, the following abbreviations are used
 - "g" for gram;
 - "iu" for international unit of activity;
 - "mcg" for microgram;
 - "mg" for milligram; and
 - "ml" for millilitre; and
- (b) in Schedule 1
 - (i) entries in any of columns 2, 3 and 4 of Parts 1 and 2 relate only to the substances specified in column 1 against which they appear,
 - (ii) where, in relation to a particular substance specified in column 1, an entry in any of columns 2, 3 and 4 bears a number or letter, that entry relates only to entries in the other of columns 2, 3 and 4 that bear the same number or letter, and
 - (iii) the entries in column 4 of Part 1 shall be read subject to the note at the end of that Part.
- (3) Without prejudice to Article 10 of the Interpretation (Jersey) Law 1954¹⁰, every provision in the Medicines (Jersey) Law 1995¹¹ that relates in any other way to its interpretation shall also apply in the same way to this Order, unless the context otherwise requires.

2 Prescription only medicines

The following descriptions and classes of medicinal products are specified for the purposes of Article 57(1)(a) of the Law, and are accordingly prescription only medicines, namely –

(a) medicinal products that consist of or contain a substance specified in column 1 of Part 1 of Schedule 1 to this Order;

- (b) medicinal products that are controlled drugs;
- (c) medicinal products that are for parenteral administration, whether or not they fall within sub-paragraph (a) or (b) of this paragraph;
- (d) medicinal products that -
 - (i) are not of a description and do not fall within a class specified in any of sub-paragraphs (a), (b) and (c) of this paragraph,
 - (ii) are of a description in respect of which the conditions in Article 58(1) of the Law are fulfilled, and
 - (iii) are products in respect of which a product licence is granted, after the commencement of this Order, containing a provision to the effect that the method of sale or supply of the medicinal product is to be only in accordance with a prescription given by a person who is an appropriate practitioner; and
- (e) cyanogenetic substances, other than preparations for external use. 12

3 Medicinal products that are not prescription only medicines

- (1) Notwithstanding Article 2, a medicinal product shall not be a prescription only medicine by reason that it consists of or contains a substance specified in column 1 of Part 1 of Schedule 1, where
 - (a) in relation to that substance there is an entry in any of columns 2, 3 and 4;
 - (b) the maximum strength in the product of that substance does not exceed the maximum strength (if any) specified in column 2; and
 - (c) the medicinal product is sold or supplied
 - (i) if a pharmaceutical form or a route of administration is specified in column 3, in such pharmaceutical form, and for administration only by such route, as may be so specified,
 - (ii) if a use is specified in column 3, in a container or package labelled (in either case) to show a use so specified to which the medicinal product is to be put but no use not so specified,
 - (iii) if a maximum dose is specified in column 4, in a container or package labelled (in either case) to show a maximum dose not exceeding that specified, and
 - (iv) if a maximum daily dose is specified in column 4, in a container or package labelled (in either case) to show a maximum daily dose not exceeding that specified.
- (2) Notwithstanding Article 2 of this Order, a medicinal product shall not be a prescription only medicine by reason that it is a controlled drug, where
 - (a) it contains not more than one of the substances specified in column 1 of Part 2 of Schedule 1 to this Order and no other controlled drug;
 - (b) it contains that substance at a strength that does not exceed the maximum strength specified in column 2; and

- (c) it is sold or supplied
 - (i) in such pharmaceutical form as may be specified in column 3, and
 - (ii) in or from a container or package labelled (in either case) to show a maximum dose not exceeding that specified in column 4.
- (4) Notwithstanding Article 2, a medicinal product specified in Part 3 or 4 of Schedule 1 shall not be a prescription only medicine.

4 New medicinal products

For the purposes of Article 58(2)(a) of the Law their duration shall be a period of 5 years.

5 Appropriate practitioner¹³

For the purposes of the Law, the following shall be appropriate practitioners –

- (a) a doctor;
- (b) a dentist;
- (c) a veterinary surgeon;
- (d) a midwife prescribing practitioner registered under the Health Care (Registration) (Jersey) Law 1995¹⁴;
- (e) a nurse prescribing practitioner registered under the Health Care (Registration) (Jersey) Law 1995;
- (f) a specialist community public health nurse prescribing practitioner registered under the Health Care (Registration) (Jersey) Law 1995.

6 Conditions for prescriptions relating to sale and supply¹⁵

- (1) For the purposes of Article 57(2)(a) of the Law (read with paragraph (4) of that Article), 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 in paragraph (2) of this Article are fulfilled.
- (2) The conditions to which paragraph (1) refers are
 - (a) the prescription shall be written in ink or otherwise so as to be indelible, unless it is a health prescription that is not for a controlled drug specified in any of Schedules 1, 2 and 3 to the Misuse of Drugs (General Provisions) (Jersey) Order 1989¹⁶, in which case it may be written using carbon paper or similar material:
 - (b) the prescription shall be signed in ink, with his or her own name, by the practitioner giving it;
 - (c) the prescription shall contain the following information –

- (i) the address of the practitioner giving it,
- (ii) the appropriate date,
- (iii) by virtue of which of the paragraphs of Article 5 the practitioner giving it is an appropriate practitioner,
- (iv) where the practitioner giving it is an appropriate practitioner other than a veterinary surgeon, the name and address of the person for whose treatment it is given and (if that person is under 12) his or her age, and
- (v) where the practitioner giving it is a veterinary surgeon, the name and address of the person to whom the prescription only medicine is to be delivered, and a declaration by that veterinary surgeon that the prescription only medicine is prescribed for an animal or herd under his or her care;
- (ca) in the case of a prescription given by a supplementary prescriber, the supplementary prescriber
 - (i) has given the prescription in accordance with the terms of a clinical management plan containing the information specified in Schedule 4, such plan relating to an individual patient and to which the following are parties –
 - (A) the patient,
 - (B) the patient's doctor or dentist, and
 - (C) the supplementary prescriber, and
 - (ii) has access to the health records of the patient to whom the clinical management plan relates to the extent that such records are used by the doctor or dentist who is a party to the plan;
- (d) the prescription 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 or otherwise than in accordance with the direction contained in the repeatable prescription; and
- (e) in the case of a repeatable prescription that does not specify the number of times that it may be dispensed, the prescription shall not be dispensed on more than 2 occasions unless it is a prescription for oral contraceptives, in which case it may be dispensed 6 times before the end of the period of 6 months from the appropriate date.¹⁷
- (3) The restrictions in Article 57(2)(a) of the Law shall not apply to a sale or supply of a prescription only medicine that, by reason only that a condition in paragraph (2) of this Article is not fulfilled, is not in accordance with a prescription given by an appropriate practitioner, where the person selling or supplying the prescription only medicine, having exercised all due diligence, believes on reasonable grounds that that condition is fulfilled in relation to that sale or supply.
- (4) In paragraph (2), the "appropriate date" means
 - (a) in the case of a health prescription –

- (i) the date on which it was signed by the practitioner by whom it was given, or
- (ii) if it also contains a date indicated by him or her as being the date before which it shall not be dispensed, the later of the 2 dates; and
- (b) in every other case, the date on which the prescription was signed by the practitioner by whom it was given.

6A Conditions for prescriptions – administration¹⁸

- (1) For the purposes of Article 57(2)(b) of the Law (read with paragraph (4) of that Article), a prescription only medicine shall not be taken to be administered by a supplementary prescriber or by a person acting in accordance with the directions of a supplementary prescriber unless the conditions in paragraph (2) are met.
- (2) Those conditions are that
 - (a) the supplementary prescriber is acting in accordance with the terms of a clinical management plan containing the information specified in Schedule 4, such plan relating to an individual patient to which the following are parties
 - (i) the patient,
 - (ii) the patient's doctor or dentist, and
 - (iii) the supplementary prescriber; and
 - (b) the supplementary prescriber has access to the health records of the patient to whom the clinical management plan relates to the extent that such records are used by the doctor or dentist who is a party to the plan.

7 Exemption for highly diluted medicinal products

The restrictions in Article 57(2) of the Law shall not apply to the sale, supply or administration of a medicinal product that is not for parenteral administration and only consists of or only contains one or more of the substances specified in column 1 of Part 1 or 2 of Schedule 1 to this Order, where –

- (a) each 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 or her 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).

8 Exemptions for specified categories of persons

(1) The restrictions in Article 57(2)(a) of the Law shall not apply –

- (a) to the sale or supply by a person specified in column 1 of Part 1 of Schedule 2 to this Order; or
- (b) to the supply by a person specified in column 1 of Part 2 of Schedule 2 to this Order,

of a prescription only medicine specified in column 2 of Part 1 or 2 of that Schedule in relation to that person, where the conditions in the corresponding paragraph in column 3 of that Part are fulfilled.

(2) The restriction in Article 57(2)(b) of the Law shall not apply to the administration by a person specified in column 1 of Part 3 of Schedule 2 to this Order of a prescription only medicine for parenteral administration specified in column 2 of that Part in relation to that person, where the conditions in the corresponding paragraph in column 3 of that Part are fulfilled.

9 Exemption for emergency sale or supply

- (1) The restrictions in Article 57(2)(a) of the Law shall not apply to the sale or supply of a prescription only medicine by a person who is lawfully conducting a retail pharmacy business, where the conditions in paragraph (2) of this Article or the alternative conditions in paragraph (3) of this Article are fulfilled.
- (2) The conditions to which paragraph (1) of this Article refers are
 - (a) 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 who, by reason of any emergency, is unable to furnish a prescription immediately;
 - (b) the doctor has undertaken to furnish the person lawfully conducting the retail pharmacy business with a prescription within 72 hours;
 - (c) the prescription only medicine is sold or supplied in accordance with the directions of the doctor requesting it;
 - (d) the prescription only medicine is not a controlled drug specified in any of Schedules 1, 2 and 3 to the Misuse of Drugs (General Provisions) (Jersey) Order 1989;¹⁹ and
 - (e) an entry is made in the register to be kept under Article 3(1) of the Medicines (Sale and Supply) (Miscellaneous Provisions) (Jersey) Order 1997²⁰, within the appropriate time specified in that Article, of the information in paragraph 1 of Schedule 2 to that Order.
- (3) The alternative conditions to which paragraph (1) of this Article refers
 - (a) 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 is satisfied
 - that there is an immediate need for that prescription only medicine to be sold or supplied and that it is impracticable in the circumstances to obtain a prescription without undue delay,

- (ii) that treatment with that prescription only medicine has been prescribed on a previous occasion by a doctor for the person requesting it from the pharmacist, or (as far as the pharmacist is reasonably able to ascertain) has been lawfully prescribed on a previous occasion by a medical practitioner outside Jersey for the person requesting it from the pharmacist, and
- (iii) as to the dose which, in the circumstances, it would be appropriate for that person to take;
- (b) the prescription only medicine
 - (i) will be sold or supplied in no greater quantity than will provide 5 days' treatment,
 - (ii) is a preparation of insulin, an aerosol for the relief of asthma, an ointment or a cream, and has been made up for sale in a container elsewhere than at the place of sale or supply, and is the smallest pack that the pharmacist has available for sale or supply,
 - (iii) is an oral contraceptive and is sufficient, but no more than sufficient, for a full cycle, or
 - (iv) is an antibiotic for oral administration in liquid form, and is the smallest quantity that will provide a full course of treatment;
- (c) the prescription only medicine does not consist of or contain a substance specified in Schedule 3 to this Order and is not a controlled drug specified in any of the Schedules to the Misuse of Drugs (General Provisions) (Jersey) Order 1989²¹;
- (d) an entry is made in the register to be kept under Article 3(1) of the Medicines (Sale and Supply) (Miscellaneous Provisions) (Jersey) Order 1997²² within the appropriate time specified in that Article, of the information in paragraph 3 of Schedule 2 to that Order; and
- (e) 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 (unless 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".23
- (4) The conditions in paragraph (2)(d) of this Article and in paragraph (3)(c) of this Article shall not apply where the prescription only medicine
 - (a) consists of or contains phenobarbitone or phenobarbitone sodium (but no other substance specified in Schedule 3 to this Order or in

any of the Schedules to the Misuse of Drugs (General Provisions) (Jersey) Order 1989²⁴); and

(b) is sold or supplied for use in the treatment of epilepsy.

10 Exemption for sale or supply in hospitals or the prison²⁵

- (1) The restrictions in Article 57(2)(a) of the Law shall not apply to the sale or supply of any prescription only medicine
 - (a) in the course of the business of a hospital; or
 - (b) by a pharmacy to the prison under the terms of a contract to supply medicinal products for the benefit of prisoners,

in accordance with the written directions of an appropriate practitioner, other than a supplementary prescriber, even though those directions do not fulfil the conditions in Article 6(2) of this Order.

(2) In the case of directions given by a supplementary prescriber, paragraph (1) applies except that the condition in Article 6(2)(ca) must be fulfilled as if the references to a prescription in that sub-paragraph were references to the directions given by the supplementary prescriber.

11 Exemption for authorised needle supply services²⁶

The restrictions of Article 57(2)(a) of the Law shall not apply to the supply by a person, for parenteral administration, of ampoules of sterile water, if the supply is made by the person in the course of acting on behalf of a service provided by or on behalf of the States for the purpose of enabling the supply of syringes, and associated articles, so as to reduce the spread of disease.

12 Exemption for sale or supply in cases involving another's default

The restrictions in Article 57(2)(a) of the Law 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, and it is because of an act or default of another person that the product is a product to which that sub-paragraph applies.

13 Exemption in the case of forged prescription

The restrictions in Article 57(2)(a) of the Law 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.

14 Exemption for parenteral administration to human beings

The restriction in Article 57(2)(b) of the Law shall not apply to the administration to human beings of any of the following medicinal products for parenteral administration, namely –

adrenaline injection BP;

atropine sulphate injection;
chlorpheniramine injection;
cobalt edetate injection;
dextrose injection strong B.P.C.;
diphenhydramine injection;
glucagon injection;
hydrocortisone injection;
mepyramine injection;
naloxone injection;
promethazine hydrochloride injection;
snake venom antiserum;
sodium nitrite injection;
sodium thiosulphate injection; and
sterile pralidoxime injection,

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

15 Exemption for non-parenteral administration to human beings

The restriction in Article 57(2)(b) of the Law shall not apply to the administration to human beings of a prescription only medicine that is not for parenteral administration.

16 Citation

This Order may be cited as the Medicines (Prescription Only) (Jersey) Order 1997.

SCHEDULE 1

(Articles 1(2), 2(a), 3(1) and 7)

PART 127

PRESCRIPTION ONLY MEDICINES

[Note -

- (x) indicates that the entry is to be read subject to paragraph 1 of the note at the end of Part 1 of Schedule 1
- (y) indicates that the entry is to be read subject to paragraph 2 of the note at the end of Part 1 of Schedule 1]

Prescription Only	Prescription Only Circumstances In Which Substances Are Not		
Medicine	Prescription Only Medicines		
Column 1	Column 2	Column 3	Column 4
Substance	Maximum strength	Use,	Maximum
		pharmaceutical	dose and
		form or route of	maximum
		administration	daily dose
Acamprosate			
Acarbose			
Acebutolol Hydrochloride			
Aceclofenac			
Acemetacin			
Acetarsol			
Acetazolamide			
Acetazolamide Sodium			
Acetohexamide			
Acetylcholine Chloride	0.2%	External	
Acetylcysteine			
Aciclovir			
Acipimox			
Acitretin			
Aclarubicin			
Hydrochloride			
Aconite	1.3%	External	
Acrivastine			
Acrosoxacin			
Actinomycin C			
Actinomycin D			
Adapalene			
Adenosine			
Adrenaline		(1) By inhaler	
		(2) External	

Prescription Only	Circumstances In Which Substances Are Not		
Medicine	Prescription Only Medicines		
Adrenaline Acid Tartrate	l ·	1) By inhaler 2) External	
Adrenaline Hydrochloride	l ·	1) By inhaler 2) External	
Adrenocortical Extract		,	
Aclofenac			
Albendazole			
Alclometasone			
Dipropionate			
Alcuronium Chloride			
Aldesleukin			
Aldosterone			
Alendronate Sodium			
Alfacalcidol			
Alfuzosin Hydrochloride			
Allergen Extracts			
Allopurinol			
Allyloestrenol			
Aloxiprin			
Alphadolone Acetate			
Alphaxalone			
Alprenolol			
Alprenolol Hydrochloride			
Alprostadil			
Alseroxylon			
Altretamine			
Amantadine			
Hydrochloride			
Ambenonium Chloride			
Ambutonium Bromide			
Amcinonide			
Ametazole Hydrochloride			
Amethocaine	lo	any use (except ocal ophthalmic se)	
Amethocaine Gentisate	A lo	any use (except ocal ophthalmic se)	
Amethocaine	A	any use (except	
Hydrochloride		ocal ophthalmic se)	
Amikacin Sulphate			
Amiloride Hydrochloride			
Aminocaproic Acid			
Aminoglutethimide			
Aminopterin Sodium			

Prescription Only	Circumstances In WI	nich Substances Aı	re Not
Medicine	Prescription Only Mo	edicines	
Amiodarone			
Hydrochloride			
Amiphenazole			
Hydrochloride			
Amisulpride			
Amitriptyline			
Amitriptyline Embonate			
Amitriptyline			
Hydrochloride			
Amlodipine Besylate			
Ammonium Bromide			
Amodiaquine			
Hydrochloride			
Amorolfine			
Hydrochloride			
Amoxapine			
Amoxycillin			
Amoxycillin Sodium			
Amoxycillin Trihydrate			
Amphomycin Calcium			
Amphotericin			
Ampicillin			
Ampicillin Sodium			
Ampicillin Trihydrate			
Amsacrine			
Amygdalin			
Amyl Nitrite			
Amylocaine		Any use (except	
Hydrochloride		local ophthalmic	
		use)	
Anastrazole			
Ancrod			
Androsterone			
Angiotensin Amide			
Anistreplase			
Anterior Pituitary Extract			
Antimony Barium			
Tartrate			
Antimony			
Dimercaptosuccinate			
Antimony Lithium			
Thiomalate			
Antimony Pentasulphide			
Antimony Potassium			
Tartrate Antimony Sodium			
Antimony Sodium Tartrate			
Antimony Sodium Thiogycollate			
imogyconacc	1		

Prescription Only Medicine	Circumstances In Which Substa	ances Are Not
Antimony Sulphate	Prescription Only Medicines	
Antimony Sulphate Antimony Trichloride		
Antimony Trioxide Antimony Trisulphide		
Apiol Apomorphine		
Apomorphine		
Hydrochloride		
Apraclonidine		
Hydrochloride		
Aprotinin		
Arecoline Hydrobromide		
Argipressin		
Aristolochia		
Aristolochia Clematitis		
Aristolochia Contorta		
Aristolochia Debelis		
Aristolochia Fang-chi		
Aristolochia Aristolochia		
Manshuriensis		
Aristolochia Serpentaria		
Arsenic		
Arsenic Triiodide		
Arsenic Trioxide		
Arsphenamine	A nyı form	(avaant
Aspirin	Any form non-effery	
	tablets or	escent
	capsules)	
Astemizole	capsuics)	
Atenolol		
Atorvastatin		
Atorvastatin Calcium		
Atovaquone Atovaquone		
Atracurium Besylate		
· ·	(1) Interna	1.
Atropine	(a) by inha	
	(b) otherw	
	than by in	1 7
	(2) Extern	
	(except loc	
	ophthalmi	
Atropine Methobromide	(1) Interna	
Thopine Memorionide	(a) by inha	
	(b) otherw	
	than by in	

Prescription Only Medicine	Circumstances In Which Substances Are Not Prescription Only Medicines		
		(2) External	
		(except local	
		ophthalmic use)	
Atropine Methonitrate		Internal:	
1		(a) by inhaler	(b) 400 mcg
		(b) otherwise	(MD) 1.3 mg
		than by inhaler	(MDD)(x)
Atropine Oxide		(1) Internal:	
Hydrochloride		(a) by inhaler	(b) 360mcg
		(b) otherwise	(MD) 1.2mg
		than by inhaler	(MDD)(x)
		(2) External	
		(except local	
		opthalmic use)	
Atropine Sulphate		(1) Internal:	
		(a) by inhaler	(b) 360 mcg
		(b) otherwise	(MD) 1.2 mg
		than by inhaler	(MD)(x)
		(2) External	
		(except local	
		ophthalmic use)	
Auranofin			
Azapropazone			
Azathioprine			
Azathioprine Sodium			
Azelaic Acid			
Azelastine Hydrochloride			
Azidocillin Potassium			
Azithromycin			
Azlocillin Sodium			
Aztreonam			
Bacampicillin			
Hydrochloride			
Bacitracin			
Bacitracin Methylene			
Disalicylate			
Bacitracin Zinc			
Baclofen			
Balsalazide Sodium			
Bambuterol			
Hydrochloride			
Barium Carbonate	_		
Barium Chloride			
Barium Sulphide			
Beclamide			
Beclomethasone			
Beclomethasone			
Diproprionate	<u> </u>		

Prescription Only	Circumstances In W	hich Substances A	re Not
Medicine Prescription Only Medicines			
Belladonna Herb		(1) Internal	(1) 1 mg of the
		(2) External	alkaloids (MDD)
Belladonna Root		(1) Internal	(1) 1 mg of the
		(2) External	alkaloids
		,	(MDD)
Bemegride			
Bemegride Sodium			
Benapryzine			
Hydrochloride			
Bendrofluazide			
Benethamine Penicillin			
Benoxaprofen			
Benperidol			
Benserazide			
Benserazide			
Hydrochloride			
Bentiromide			
Benzathine Penicillin			
Benzbromarone			
Benzhexol Hydrochloride			
Benzilonium Bromide			
Benzocaine		Any use (except	
		local ophthalmic use)	
Benzoctamine			
Hydrochloride			
Benzoyl Peroxide	10.0%	External	
N-Benzoyl			
Sulphanilamide			
Benzquinamide			
Benzquinamide			
Hydrochloride			
Benzthiazide			
Benztropine Mesylate			
Benzylpenicillin Calcium			
Benzylpenicillin			
Potassium			
Benzylpenicillin Sodium			
Beractant			
Betahistine			
Hydrochloride			
Betamethasone			
Betamethasone			
Adamantoate			
Betamethasone Benzoate			
Betamethasone			
Dipropionate			

Medicine Prescription Only Medicines Betamethasone Sodium Phosphate Betamethasone Valerate Betaxolol Hydrochloride Bethanchol Chloride Bicalutamide Bicalutamide Biperiden Hydrochloride Biperiden Lactate Bisicalutamide Biperiden Lactate Bisimuth Glycollylarsanilate Bismuth Bismuth Bismuth Bismorpolol Fumarate Bleomycin Sulphate Brimonidine Tartrate Brimonidine Tartrate Brimonidine Tartrate Brimonidine Tartrate Brombexine Brombexine Brombexine Bromocriptine Mesylate Bromocriptine Budesonide Busulphan Budesonide Busulphan Bupivacaine Any use (except local ophthalmic use) Buserelin Acetate Buspirone Hydrochloride Busulphan Butacaine Sulphate Any use (except local ophthalmic use) Butorphenol Tartrate Butorphenol Hydrate Calcipotriol	Prescription Only	Circumstances In Wh	nich Substances A	re Not
Betamethasone Sodium Phosphate Betamethasone Valerate Betaxolol Hydrochloride Bethanechol Chloride Bethanidine Sulphate Bezafibrate Bicalutamide Biperiden Hydrochloride Biperiden Hydrochloride Biperiden Hydrochloride Biperiden Hydrochloride Bismuth Glycollylarsanilate Bisoprolol Fumarate Bleomycin Bleomycin Sulphate Bretylium Tosylate Brimonidine Tartrate Bromhexine Hydrochloride Bromocriptine Mesylate Bromocriptine Mesylate Bromocriptine Mesylate Bromozindia Budesonide Bufexamac Buphenine Hydrochloride Gabergoline Any use (except local ophthalmic use) Buserelin Acetate Buspirone Hydrochloride Buserine Hydrochloride Any use (except local ophthalmic use) Buserelin Acetate Buspirone Hydrochloride Busilphan Butacaine Sulphate Any use (except local ophthalmic use) Butorphenol Tartrate Butorphenol Tartrate Butorphenol Tartrate Butorpiolic Calcipotriol Calcipotriol Hydrate Calcitoriin Hydrothoride Calcipotriol Hydrate Calcitoriin Hydrate Calcipotriol Hydrate Calcipot		Prescription Only Me	edicines	
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Butriptyline Hydrochloride Cabergoline Calcipotriol Calcipotriol Hydrate Calcitonin			use)	
Hydrochloride Cabergoline Calcipotriol Calcipotriol Hydrate Calcitonin	Butorphenol Tartrate			
Cabergoline Calcipotriol Calcipotriol Hydrate Calcitonin				
Calcipotriol Calcipotriol Hydrate Calcitonin	Hydrochloride			
Calcipotriol Hydrate Calcitonin	Cabergoline			
Calcitonin	Calcipotriol			
Calcitonin	Calcipotriol Hydrate			
Calcitriol				
	Calcitriol			

Prescription Only		In Which Substance	es Are Not
Medicine	Prescription O	nty Medicines	
Calcium Amphomycin Calcium			
Benzamidosalicylate			
Calcium Bromide			
Calcium			
Bromidolactobionate			
Calcium Carbimide			
Calcium Folinate			
Calcium Metrizoate			
Calcium Sulphaloxate			
Candesartan Cilexetil			
Candicidin			
Canrenoic Acid	0.04		
Cantharidin	0.01%	External	
Capreomycin Sulphate			
Captopril			
Carbachol			
Carbamazepine			
Carbaryl			
Carbasalate Calcium			
Carbenicillin Sodium			
Carbenoxolone Sodium		(1) Pellet	(1) 5 mg (MD) 25 mg (MDD)
	(2) 2.0%	(2) Gel	
Carbidopa	(2) 2.070	(2) Ger	
Carbimazole			
Carbocisteine			
Carbon Tetrachloride			
Carboplatin			
Carboprost Trometamol			
Carbuterol Hydrochloride			
Carfecillin Sodium			
Carindacillin Sodium			
Carisoprodol			
Carmustine			
Carperidine Carteolol Hydrochloride			
Cefaclor			
Cefadroxil			
Cefazedone Sodium			
Cefdinir			
Cefixime			
Cefodizime Sodium			
Cefotaxime Sodium			
Cefoxitin Sodium			
Cefpodoxime Proxetil			

Prescription Only		s In Which Substances A	re Not
Medicine	Prescription C	only Medicines	
Cefprozil			
Cefsulodin Sodium			
Ceftazidime			
Ceftizoxime Sodium			
Ceftriaxone Sodium			
Cefuroxime Axetil			
Cefuroxime Sodium			
Celiprolol Hydrochloride			
Cephalexin			
Cephalexin Sodium			
Cephaloridine			
Cephalothin Sodium			
Cephamandole Nafate			
Cephazolin Sodium			
Cephradine			
Cerium Oxalate			
Cerivastatin			
Cerivastatin Sodium			
Ceruletide Diethylamine			
Cetirizine			
Chenodeoxycholic Acid			
Chloral Hydrate		External	
Chlorambucil		External	
Chloramphenicol			
Chloramphenicol			
Cinnamate			
Chloramphenicol			
Palmitate			
Chloramphenicol Sodium			
Succinate			
Chlorhexadol			
Chlormadinone Acetate			
Chlormerodrin			
Chlormethiazole			
Chlormethiazole			
Edisylate			
Chlormezanone			
Chloroform	(1) 5.0%	(1) Internal	
Cinorororiii	(1) 5.070	(2) External	
Chloroquine Phosphate		Prophylaxis of	
omoroganio i nospilate		malaria	
Chloroquine Sulphate		Prophylaxis of	
		malaria	
Chlorothiazide			
Chlorotrianisene			
Chlorphenoxamine			
Hydrochloride			
Chlorpromazine			
Cinoipioniazine	1		

Prescription Only	Circumstances In Which Substances Are Not		
Medicine	Prescription Only Me	edicines	1
Chlorpromazine			
Embonate			
Chlorpromazine			
Hydrochloride			
Chlorpropamide			
Chlorprothixene			
Chlorprothixene			
Hydrochloride			
Chlortetracycline			
Chlortetracycline			
Calcium			
Chlortetracycline			
Hydrochloride			
Chlorthalidone			
Chlorzoxazone			
Cholestyramine			
Chorionic Gonadotrophin			
Ciclacillin			
Ciclobendazole			
Cidofovir			
Cilastatin Sodium			
Cilazapril			
Cimetidine			
Cimetidine Hydrochloride			
Cinchocaine	3.0%	Any use (except	
		local ophthalmic	
		use)	
Cinchocaine	Equivalent of 3.0% of	Any use (except	
Hydrochloride	Cinchocaine	local ophthalmic	
		use)	
Cinchophen			
Cinoxacin			
Ciprofibrate			
Ciprofloxacin			
Ciprofloxacin			
Hydrochloride			
Cisapride			
Cisplatin			
Citalopram			
Hydrobromide			
Clarithromycin			
Clavulanic Acid			
Clenbuterol			
Hydrochloride			
Clidinium Bromide			
Clindamycin			

Prescription Only Medicine	Circumstances In V Prescription Only M		re Not
Clindamycin			
Hydrochloride			
Clindamycin Palmitate			
Hydrochloride			
Clindamycin Phosphate			
Clioquinol	(1) 35 mg	(1) Treatment of	(1) 350 mg
1		mouth ulcers (2) External (except treatment of mouth ulcers)	(MDD)
Clobetasol Propionate			
Clobetasone Butyrate			
Clofazimine			
Clofibrate			
Clomiphene Citrate			
Clomipramine			
Clomipramine			
Hydrochloride			
Clomocycline			
Clomocycline Sodium			
Clonidine			
Clonidine Hydrochloride			
Clopamide			
Clopenthixol Decanoate			
Clopenthixol			
Hydrochloride			
Clorexolone			
Clostebol Acetate			
Clotrimazole		External but, in	
		the case of	
		vaginal use, only	
		for the treatment	
		of vaginal	
		candidiasis	
Cloxacillin Benzathine			
Cloxacillin Sodium			
Clozapine			
Cocculus Indicus			
Co-dergocrine Mesylate			
Colaspase			
Colchicine			
Colestipol Hydrochloride			
Colfosceril Palmitate			
Colistin Sulphate			
Colistin Sulphomethate			
Colistin Sulphomethate			
Sodium			
Coniine			
Conium Leaf	7.0%	External	

Prescription Only Medicine Circumstances In Which Substances Are Not Prescription Only Medicines			e Not
Corticotrophin	1100011011	3 111 J 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
Cortisone			
Cortisone Acetate			
Co-tetroxazine			
Co-Trimoxazole			
Copropamide			
Crotethamide			
Croton Oil			
Croton Seed			
Curare			
Cyclofenil			
Cyclopenthiazide			
Cyclopentolate			
Hydrochloride			
Cyclophosphamide			
Cycloserine			
Cyclosporin Cyclothiazide			
Cyproterone Acetate			
Cytarabine			
Cytarabine Hydrochloride			
Dacarbazine			
Dalteparin Sodium			
Danazol			
Danthron			
Dantrolene Sodium			
Dapsone			
Dapsone Ethane Ortho			
Sulphonate			
Daunorubicin			
Hydrochloride			
Deanol Bitartrate			26 mg (MDD)
Debrisoquine Sulphate			
Demecarium Bromide			
Demeclocycline			
Demeclocycline Calcium			
Demeclocycline			
Hydrochloride			
Deoxycortone Acetate			
Deoxycortone Pivalate			
Deptotropine Citrate			
Dequalinium Chloride	(1) 0.25 mg	(1) Internal:	
	_	throat lozenges or	
		throat pastilles	
	(2) 1.0%	(2) External:	
		paint	
Deserpidine			

Prescription Only Medicine	Circumstances In Which Substances Are Not Prescription Only Medicines		
Desferrioxamine	1 rescription Only Wi	dicines	
Mesylate			
Desflurane			
Desfluorotriamcinolone			
Desipramine			
Hydrochloride Deslanoside			
Desmopressin			
Desogestrel			
Desonide			
Desoxymethasone			
Dexamethasone			
Dexamethasone Acetate			
Dexamethasone			
Isonicotinate			
Dexamethasone			
Phenylpropionate			
Dexamethaone Pivalate			
Dexamethasone Sodium			
<i>m</i> -Sulphobenzoate			
Dexamethasone Sodium			
Phosphate			
Dexamethasone			
Troxundate			
Dexfenfluramine			
Hydrochloride		T., 4 1	T., 41
Dextromethorphan		Internal	In the case of
Hydrobromide			a controlled release
			preparation: equivalent of
			30 mg of
			Dextromethor-
			phan (MD)
			equivalent of
			75 mg of
			Dextromethor-
			phan (MDD)
			In any other
			case:
			equivalent of
			15 mg of
			Dextromethor-
			phan (MD)
			equivalent of
			75 mg of
			Dextromethor-
			phan (MDD)
Dextrothyroxine Sodium			

Prescription Only	Circumstances In Which Substances Are Not		
Medicine	Prescription Only Medi	cines	
Diazoxide			
Dibenzepin			
Hydrochloride			
Dichloralphenazone			
Dichlorphenamide			
Diclofenac			
Diethylammonium			
Diclofenac Potassium			
Diclofenac Sodium			
Dicyclomine			10 mg (MD)
Hydrochloride			60 mg (MDD)
Didanosine			
Dienoestrol			
Diethanolamine Fusidate			
Diflucortolone Valerate			
Diflunisal			
Digitalin			
Digitalis Leaf			
Digitalis, Prepared			
Digitoxin			
Digoxin			
Dihydralazine Sulphate			
Dihydroergotamine Dihydroergotamine			
Mesylate			
Dihydrostreptomycin			
Dihydrostreptomycin			
Sulphate			
Diloxanide Furoate			
Diltiazem Hydrochloride			
Dimercaprol			
Dimethisoquin		ny vsa (avaant	
Hydrochloride		any use (except ocal ophthalmic	
Hydrochioride		se)	
Dimethisterone	us	50)	
Dimethothiazine			
Mesylate			
Dimethyl Sulphoxide			
Dimethyltubocurarine Bromide			
Dimethyltubocurarine Chloride			
Dimethyltubocurarine			
lodide			
Dinoprost			
Dinoprost Trometamol			
Dinoprostone			

Prescription Only	Circumstances In Which Substances Are Not		
Medicine	Prescription Only Medicines		
Diphenhydramine		All preparations	
Hydrochloride		except liquid- filled capsules	
Dipivefrin Hydrochloride			
Dipyridamole			
Disodium Etidronate			
Disopyramide			
Disopyramide Phosphate			
Distigmine Bromide			
Disulfiram			
Dithranol	1.00%		
Dobutamine			
Hydrochloride			
Dolasetron Mesilate			
Domperidone			
Domperidone Maleate			
Donepezil			
Donepezil Hydrochloride			
Dopamine Hydrochloride			
Dopexamine			
Hydrochloride			
Dorzolamide			
Hydrochloride			
Dothiepin			
Dothiepin Hydrochloride			
Doxapram Hydrochloride			
Doxazosin Mesylate			
Doxepin Hydrochloride			
Doxorubicin			
Doxorubicin			
Hydrochloride			
Doxycycline			
Doxycycline Calcium			
Chelate			
Doxycycline			
Hydrochloride			
Droperidol			
Drostanolone			
Drostanolone Propionate			
Dydrogesterone			
Dyflos			
Econazole		External, but in	
		the case of	
		vaginal use, only	
		for the treatment	
		of vaginal	
		candidiasis	

Prescription Only	Circumstances In Which Substances Are Not		
Medicine	Prescription Only Me	edicines	
Econazole Nitrate		External, but in the case of vaginal use, only for the treatment	
		of vaginal candidiasis	
Ecothiopate Iodide			
Edrophonium Chloride			
Eflornithine Hydrochloride			
Eformoterol Fumarate			
Embutramide			
Emepronium Bromide			
Emetine	1.0%		
Emetine Bismuth			
Iodide			
Emetine Hydrochloride	Equivalent of 1.0% of Emetine		
Enalapril Maleate			
Encephalitis Virus, Tickborne, Central European			
Enoxacin			
Enoxaparin Sodium			
Enoximone			
Ephedrine		(1) Internal (other than nasal sprays or nasal drops)	(1) 30 mg (MD) 60 mg (MDD)
	(2) 2.0%	(2) Nasal sprays or nasal drops (3) External	
Ephedrine Hydrochloride		(1) Internal (other than nasal sprays or nasal drops)	(1) Equivalent of 30 mg of Ephedrine (MD) Equivalent of 60 mg of Ephedrine (MDD)
	(2) Equivalent of 2.0% of Ephedrine	(2) Nasal sprays or nasal drops (3) External	

Prescription Only	Circumstances In Which Substances Are Not		
Medicine	Prescription Only Medicines		
Ephedrine Sulphate		(1) Internal (other than nasal sprays or nasal drops)	(1) Equivalent of 30 mg of Ephedrine (MD) Equivalent of 60 mg of Ephedrine (MDD)
	(2) Equivalent of	(2) Nasal sprays	
	2.0% of Ephedrine	or nasal drops	
	•	(3) External	
Epicillin			
Epirubicin			
Epirubicin Hydrochloride			
Epithiazide			
Epoetin Alfa			
Epoetin Beta			
Epoprostenol Sodium			
Ergometrine Maleate			
Ergometrine Tartrate			
Ergot, Prepared			
Ergotamine Tartrate			
Erythromycin			
Erythromycin Estolate			
Erythromycin Ethyl			
Carbonate			
Erythromycin Ethyl			
Succinate			
Erythromycin			
Lactobionate			
Erythromycin Phosphate			
Erythromycin Stearate			
Erythromycin			
Thiocyanate			
Esmolol Hydrochloride			
Estramustine Phosphate			
Estramustine Sodium			
Phosphate			
Etafedrine Hydrochloride			
Ethacrynic Acid			
Ethambutol			
Hydrochloride			
Ethamivan			
Ethamsylate			
Ethiazide			
Ethinyl Androstenediol			
Ethinyloestradiol			
Ethionamide			
Ethisterone			

Prescription Only	Circumstances In Which Substances Are Not		
Medicine	Prescription Only Medicines		
Ethoglucid			
Ethoheptazine Citrate			
Ethopropazine			
Hydrochloride			
Ethosuximide			
Ethotoin			
Ethyl Biscoumacetate			
Ethyloestrenol			
Ethynodiol Diacetate			
Etodolac			
Etomidate			
Etomidate Hydrochloride			
Etoposide			
Etretinate			
Exemestane			
Famotidine			
Fazadinium Bromide			
Felbinac			
Felodipine			
Felypressin			
Fenbufen			
Fencamfamin			
Hydrochloride			
Fenclofenac			
Fenfluramine			
Hydrochloride			
Fenofibrate			
Fenoprofen			
Fenoprofen Calcium			
Fenoterol Hydrobromide		4 .	
Fenticonazole Nitrate	External use		
	in the case of		
	vaginal use, o		
	for the treatm		
	of vulvovagii	nal	
	candidiasis)		
Feprazone			
Ferrous Arsenate			
Ferumoxsil			
Fexofenadine			
Hydrochloride			
Filgrastim			
Finasteride			
Flavoxate Hydrochloride			
Flecainide Acetate			
Flosequinan			
Fluanisone			

Prescription Only	Circumstances In W		e Not
Medicine	Prescription Only M	edicines	I
Flubendazole			
Fluclorolone Acetonide			
Flucloxacillin Magnesium			
Flucloxacillin Sodium			
Fluconazole			
Flucylosine			
Fludrocortisone Acetate			
Flufenamic Acid			
Flumazenil			
Flumethasone			
Flumethasone Pivalate			
Flunisolide			
Fluocinolone Acetonide			
Fluocinonide			
Fluocortin Butyl			
Fluocortolone			
Fluocortolone Hexanoate			
Fluocortolone Pivalate			
Fluorescein Dilaurate			
Fluorometholone			
Fluorouracil			
Fluorouracil Trometamol			
Fluoxetine Hydrochloride			
Fluoxymesterone			
Flupenthixol Decanoate			
Flupenthixol			
Hydrochloride			
Fluperolone Acetate			
•			
Fluphenazine Decanoate Fluphenazine Enantate			
Fluphenazine			
Hydrochloride			
Fluprednidene Acetate			
Fluprednisolone			
Fluprostenol Sodium			
Flurandrenolone	0.77		
Flurbiprofen	8.75 mg	throat lozenges	43.75 mg (MDD)
Flurbiprofen Sodium			
Fluspirilene			
Flutamide			
Fluticasone Propionate		Aqueous nasal	
		sprays for the	
		treatment of	
		allergic rhinitis in	
		persons not less	
		than 18 years	
Flutrimazole			

Prescription Only	escription Only Circumstances In Which Substances Are Not		
Medicine	Prescription On	ly Medicines	
Fluvastatin Sodium	•		
Fluvoxamine Maleate			
Folic Acid			500 mcg
			(MDD)
Formestane			
Formocortal			
Formoterol Fumarate			
Foscarnet Sodium			
Fosfestrol Sodium			
Fosfomycin Trometamol			
Fosinopril Sodium			
Framycetin Sulphate			
Frusemide			
Furazolidone			
Fusafungine			
Fusidic Acid			
Gabapentin			
Gadolinium			
Gadoteridol			
Gallamine Triethiodide			
Ganciclovir			
Ganciclovir Sodium			
Gelsemine	0.1%		
Gelsemium	0.170		25 mg (MD)
Gersenham			75 mg (MDD)
Gemeprost			70 mg (1122)
Gemfibrozil			
Gentamicin			
Gentamicin Sulphate			
Gestodene			
Gestrinone			
Gestronol			
Gestronol Hexanoate			
Glibenclamide			
Glibornuride			
Gliclazide			
Glimepiride			
Glipizide			
Gliquidone			
Glisoxepide			
Glucagon			
Glycopyrronium Bromide			1 mg (MD)
			2 mg (MDD)
Glymidine			
Gonadorelin			
Goserelin Acetate			
Gramicidin	0.2%	External	

Prescription Only	Circumstances In W		re Not
Medicine	Prescription Only M	ledicines	
Granisetron			
Hydrochloride			
Griseofulvin			
Growth Hormone			
Guanethidine			
Monosulphate			
Guanfacine			
Hydrochloride			
Guanoclor Sulphate			
Guanoxan Sulphate			
Halcinonide			
Halofantrine			
Hydrochloride			
Haloperidol			
Haloperidol Decanoate			
Heparin		External	
Heparin Calcium		External	
Hexachlorophane		External:	
1	(a) 2.0%	(a) soaps	
	(b) 0.1%	(b) aerosols	
	(c) 0.75%	(c) preparations	
		other than soaps	
		and aerosols	
Hexamine			
Phenylcinchoninate			
Hexobarbitone			
Hexobarbitone Sodium			
Hexoestrol			
Hexoestrol Dipropionate			
L-Histidine		Dietary or	
Hydrochloride		nutritive use	
Homatropine		(1) Internal	(1) 0.15 mg
1			(MD) 0.45 mg
			(MDD)
		(2) External	
		(except local	
		ophthalmic use)	
Homatropine			0.2 mg (MD)
Hydrobromide			0.6 mg
			(MDD)
Homatropine			2 mg (MD)
Methylbromide			6 mg (MDD)
Hydralazine			
Hydrochloride			<u> </u>
Hydrargaphen		Local application to skin	
Hydrobromic Acid			
Hydrochlorothiazide			
Hydrocortisone			
22, 3100011150110	1		1

Prescription Only	Circumstances In	Which Substances A	re Not
Medicine	Prescription Only	Medicines	
Hydrocortisone Acetate			
Hydrocortisone Butyrate			
Hydrocortisone Caprylate			
Hydrocortisone Hydrogen			
Succinate			
Hydrocortisone Sodium			
Phosphate			
Hydrocortisone Sodium			
Succinnate			
Hydrocyanic Acid			
Hydroflumethiazide			
Hydroxychloroquine		Prophylaxis of	
Sulphate		malaria	
Hydroxyprogesterone			
Hydroxyprogesterone			
Enanthate			
Hydroxyprogesterone			
Hexanoate			
Hydroxyurea			
Hydroxyzine Embonate			
Hydroxyzine			
Hydrochloride			
Hyoscine	(1) 0.15%	(1) Internal	
		(2) External	
		(except local	
		ophthalmic use)	
Hyoscine Butylbromide		(1) Internal:	(MDD)(x)
		(a) by inhaler	
		(2) External	
Hyoscine Hydrobromide		(1) Internal:	
		(a) by inhaler	
		(b) otherwise	(b) 300 mcg
		than by inhaler	(MD) 900 mcg
		(2) 7	(MDD)(x)
		(2) External	
		(except local	
TT : 34 d 1 :1		ophthalmic use)	
Hyoscine Methobromide		(1) Internal:	
		(a) by inhaler	(h) 2.5 m c
		(b) otherwise	(b) 2.5 mg
		than by inhaler	(MD) 7.5 mg (MDD)(x)
		(2) External	
Hyoscine Methonitrate		(1) Internal:	
		(a) by inhaler	
		(b) otherwise	(b) 2.5 mg
		than by inhaler	(MD) 7.5 mg
			(MDD)(x)

Prescription Only Medicine	Circumstances In Which Substances An Prescription Only Medicines	re Not
	(2) External	
Hyoscyamine	(1) Internal:	
	(a) by inhaler	
	(b) otherwise	(b) 300 mcg
	than by inhaler	(MD)
		1 mg
		(MDD)(x)
	(2) External	,,,,
	(3) Preparations	
	for the relief of	
	asthma in the	
	form of	
	cigarettes,	
	smoking mixtures	
	or fumigants	
	which contain	
	Hyoscyamine as	
	an alkaloid of	
	Stramonium	
Hyoscyamine	(1) Internal:	
Hydrobromide	(a) by inhaler	
	(b) otherwise	(b) Equiva-
	than by inhaler	lent of 300
		mcg of
		Hyoscyamine
		(MD) Equi-
		valent of 1 mg
		of Hyos-
		cyamine
		(MDD)(x)
Uranaramina Culmbata	(2) External	
Hyoscyamine Sulphate	(1) Internal: (a) by inhaler	
	(b) otherwise	(b) Equiva-
	than by inhaler	lent of 300
		mcg of
		Hyoscyamine
		(MD)
		Equivalent of
		1 mg of
		Hyoscyamine
	(2) F (1	(MDD)(x)
Ihungafan	(2) External	
Ibuprofen		

Prescription Only	Circumstances In Whic	ch Substances Ar	e Not
Medicine	Prescription Only Medi	icines	
Ibuprofen Lysine	R	Cheumatic and	
•	m	nuscular pain,	
	pa	ain of non-	
	se	erious arthritic	
	CO	onditions,	
	ba	ackache,	
	ne	euralgia,	
	m	nigraine,	
	h	eadache, dental	
	pa	ain,	
		ysmenorrhoea,	
		everishness,	
		ymptoms of	
		olds and	
		nfluenza	
	Ir	nternal	(a) In the case
			of a prolonged
			release
			preparation
			600 mg (MD)
			1,200 mg
			(MDD)
			(b) In any
			other case 400
			mg (MD)
			1,200 mg
			(MDD)
Idarubicin Hydrochloride			
Idoxuridine			
Ifosfamide			
Ignatius Bean			
Imidapril Hydrochloride			
Imipenem Hydrochloride			
Imipramine			
Imipramine			
Hydrochloride			
Imipramine Ion Exchange			
Resin Bound Salt or			
Complex			
Indapamide Hemihydrate			
Indinavir			
Indomethacin			
Indomethacin Sodium			
Indoramin Hydrochloride			
Indoprofen			
Inosine Pranobex			
Insulin			
Iodamide			

Prescription Only Medicine	Circumstances In W Prescription Only M		re Not
Iodamide Meglumine			
Iodamide Sodium			
Iohexol			
Iomeprol			
Iopamidol			
Iopentol			
Iothalamic Acid			
Ioversol			
Ioxaglic Acid			
Ipratropium Bromide			
Iprindole Hydrochloride			
Iproniazid Phosphate			
Irbesartan			
Isoaminile			
Isoaminile Citrate			
Isocarboxazid			
Isoconazole Nitrate		External, but in	
13000ma2010 1 mate		the case of	
		vaginal use, only	
		for the treatment	
		of vaginal	
		candidiasis	
Isoetharine			
Isoetharine Hydrochloride			
Isoetharine Mesylate			
Isoniazid			
Isoprenaline			
Hydrochloride			
Isoprenaline Sulphate			
Isopropamide Iodide			Equivalent of
			2.5 mg of
			Isoprop-amide
			ion (MD)
			Equivalent of
			5.0 mg of
			Isoprop-amide
			ion (MDD)
Isotretinoin			
Isradipine			
Itraconazole			
Jaborandi		External	
Kanamycin Acid Sulphate			
Kanamycin Sulphate			
Ketamine Hydrochloride			
Ketoconazole			
Ketoprofen			
Ketorolac Trometamol			
	1	1	1
Ketotifen Pumarate			

Prescription Only	Circumstances In Wh		re Not
Medicine	Prescription Only Me	edicines	
Lachesine Chloride			
Lacidipine			
Lamivudine			
Lamotrigine			
Lanatoside C			
Lanatoside Complex A, B			
and C			
Lanzoprazole			
Latamoxef Disodium			
Latanaprost			
Lercanidipine			
Hydrochloride			
Letrozole			
Levallorphan Tartrate			
Levobunolol			
Hydrochloride			
Levocabastine			
Hydrochloride			
Levocarnitine		For dietary	
Levocarmene		supplementation	
Levodopa		зарринентатіон	
Levofloxacin			
Hemihydrate			
Levonorgestrel			
Lidoflazine			
Lignocaine		Any use (except	
Lighteame		local ophthalmic	
		use)	
Lignocaine		Any use (except	
Hydrochloride		local ophthalmic	
Trydrocmonde		use)	
Lincomycin		use)	
Lincomycin			
Hydrochloride			
Liothyronine Sodium			
Lisinopril			
Lithium Carbonate			Equivalent of
Zimani caroonac			5 mg of
			Lithium (MD)
			Equivalent of
			15 mg of
			Lithium
			(MDD)
Lithium Citrate			, ,
	<u> </u>	1	1

Prescription Only Circumstances In Which Substances Are Not			re Not
Medicine	Prescription Only M	1edicines	
Lithium Sulphate			Equivalent of 5 mg of Lithium (MD) Equivalent of 5 mg of Lithium (MDD)
Lithium Succinate			(====)
Lobeline		(1) Internal	(1) 3 mg (MD) 9 mg (MDD)
		(2) External	
Lobeline Hydrochloride		(1) Internal	(1) Equivalent of 3 mg of Lobeline (MD)
			Equivalent of 9 mg of Lobeline (MDD)
		(2) External	
Lobeline Sulphate		(1) Internal	(1) Equivalent of 3 mg of Lobeline (MD) Equivalent of 9 mg of Lobeline (MDD)
		(2) External	
Lodaximide Trometamol	equivalent of 0.1% Lodoxamide	For the treatment of ocular signs and symptoms of allergic conjunctivitis, in adults and in children aged 4 years and over	
Lofepramine			
Lofepramine Hydrochloride Lofexidine Hydrochloride			
Lomefloxacin Hydrochloride			
Lornoxicam			
Lomustine		T	
Loperamide		Treatment of	
Hydrochloride Loratidine		acute diarrhoea	
Losartan Potassium			

Prescription Only Medicine	Circumstances In W Prescription Only M		re Not
Loxapine Succinate			
Lung Surfactant Porcine			
Luteinising Hormone			
Lymecycline			
Lynoestrenol			
Lypressin			
Lysuride Maleate			
Mafenide			
Mafenide Acetate			
Mafenide Hydrochloride			
Mafenide Propionate	5.0%	Eye drops	
Magnesium Fluoride	3.070	Lyc drops	
Magnesium Metrizoate			
Mandragora Autumnalis			
Mannomustine			
Hydrochloride			
Maprotiline			
Hydrochloride			
Mebanazine			
Mebendazole			
Mebeverine		(a) For the	(a) 135mg
Hydrochloride		symptomatic relief of irritable bowel syndrome (b) For uses other than the symptomatic relief of irritable bowel syndrome	(MD) 405mg (MDD) (b) 100 mg (MD) 300mg (MDD)
Mebeverine Pamoate		oower syndrome	
Mebhydrolin	1		
Mebhydrolin	1		
Napadisylate			
Mecamylamine			
Hydrochloride			
Mecillinam			
Meclofenoxate			
Hydrochloride			
Medigoxin			
Medrogestone			
Medroxyprogesterone			
Acetate			
Mefenamic Acid			
Mefloquine			
Hydrochloride			
Mefruside			
Megestrol			
Megestrol Acetate			
Micgesuol Actidit	1		1

Prescription Only Medicine	Circumstances In Williamstances In Williamstance		re Not
Meglumine	1 rescription only 1/1		
Gadopentetate			
Meglumine lodoxamate			
Meglumine loglycamate			
Meglumine lothalamate			
Meglumine lotroxate			
Meglumine loxaglate			
Meloxicam			
Melphalan			
Melphalan Hydrochloride			
Menotrophin			
			25 mg (MD)
Mepenzolate Bromide			25 mg (MD) 75 mg (MDD)
Mephenesin			
Mephenesin Carbamate			
Mepivacaine		Any use (except	
Hydrochloride		local ophthalmic	
		use)	
Meptazinol		·	
Hydrochloride			
Mequitazine			
Mercaptamine Bitartrate			
Mercaptopurine			
Mersalyl			
Mersalyl Acid			
Mesalazine			
Mesna			
Mesterolone			
Mestranol			
Metaraminol Tartrate			
Metergoline			
Metformin Hydrochloride			
Methacycline Methacycline			
Methacycline Calcium			
Methacycline Methacycline			
Hydrochloride			
Methallenoestril			
Methandienone			
Methicillin Sodium			
Methixene			
Methixene Hydrochloride			
Methocarbamol			
Methocidin		Throat lozonges	
iviculocidili		Throat lozenges and throat pastilles	
Methohexitone Sodium		pasunes	
Methoin			
Methoserpidine		<u> </u>	

Prescription Only		Which Substances A	re Not
Medicine	Prescription Only N	Medicines	
Methotrexate			
Methotrexate Sodium			
Methotrimeprazine			
Methotrimeprazine			
Hydrochloride			
Methotrimeprazine			
Maleate			
Methoxamine	0.25%	Nasal sprays, or	
Hydrochloride		nasal drops, not	
		containing in	
		either case liquid	
		paraffin as a	
N. d		vehicle	
Methsuximide			
Methyclothiazide			
Methyldopa			
Methyldopa			
Hydrochloride			
Methylephedrine			30 mg (MD)
Hydrochloride			60 mg (MDD)
Methylprednisolone			
Methylprednisolone			
Acetate			
Methylprednisolone			
Sodium Succinate			
Methyltestosterone			
Methylthiouracil			
Methysergide Maleate			
Metipranolol			
Metirosine			
Metoclopramide			
Hydrochloride			
Metolazone			
Metoloprolol Tartrate			
Metoprolol Fumarate			
Metoprolol Succinate			
Metronidazole			
Metronidazole Benzoate			
Metyrapone			
Mexiletine Hydrochloride			
Mezlocillin Sodium			
Mianserin Hydrochloride			
Mibefradil			
Dihydrochloride			
Miconazole		External	

Prescription Only	Circumstances In Which Substances Are Not		
Medicine	Prescription O		
Miconazole Nitrate		External, but in	
		the case of	
		vaginal use, only	
		for the treatment	
		of vaginal	
		candidiasis	
Mifepristone			
Miglitol			
Milrinone			
Milrinone Lactate			
Minocycline			
Minocycline			
Hydrochloride			
Minoxidil	(1) 2.0%	External	
	(2) 5.0%	External, for the	
		treatment of	
		alopecia	
		androgenetica in	
		men who have	
		attained the age	
		of 18 years but	
		have not attained	
		the age of 65	
		years	
Mirtazapine			
Misoprostol			
Mitobronitol			
Mitomycin C			
Mitozantrone			
Hydrochloride			
Mivacurium Chloride			
Mizolastine			
Moclobemide			
Modafinil			
Moexipril Hydrochloride			
Molgramostim			
Molindone Hydrochloride			
Mometasone Furoate			
Moracizine			
Hydrochloride			
Morazone Hydrochloride			
Moxisylyte			Ī
Hydrochloride			
Moxonidine			
Mupirocin			
Mupirocin Calcium			
Mustine Hydrochloride			
Mycophenolate Mofetil			
Nabilone			

Prescription Only	Circumstances In W		re Not
Medicine	Prescription Only M	<u>ledicines</u>	1
Nabumetone			
Nadolol			
Nafarelin Acetate			
Naftidrofuryl Oxalate			
Naftifine Hydrochloride			
Nalbuphine			
Hydrochloride			
Nalidixic Acid			
Nalorphine			
Hydrobromide			
Naloxone Hydrochloride			
Naltrexone Hydrochloride			
Nandrolone Decanoate			
Nandrolone Laurate			
Nandrolone			
Phenylpropionate			
Naphazoline	(1) 0.05%	(1) Nasal sprays,	
Hydrochloride	(1) 0.0070	or nasal drops,	
		not containing in	
		either case liquid	
		paraffin as a	
		vehicle	
	(2) 0.015%	(2) Eye drops	
Naphazoline Nitrate	0.05%	Nasal sprays, or	
Tuphazomie Turate	0.0370	nasal drops, not	
		containing in	
		either case liquid	
		paraffin as a	
		vehicle	
Naproxen			
Naproxen Sodium			
Naratriptan			
Hydrochloride			
Natamycin			
Nebivolol Hydrochloride			
Nedocromil Sodium			
Nefazadone Nefazadone			
Hydrochloride			
Nefopam Hydrochloride			
Neomycin			
Neomycin Oleate			
Neomycin Palmitate			
Neomycin Sulphate			
Neomycin Undecanoate			
Neostigmine Bromide			
Neostigmine Mathylaulahata			
Methylsulphate			
Netilmicin Sulphate			

Prescription Only	Circumstances In W		re Not
Medicine	Prescription Only M	edicines	T
Nicardipine			
Hydrochloride			
Nicergoline			
Niceritrol			
Nicotinic Acid		Any use (except for the treatment of hyperlipid- aemia)	600 mg (MDD)
Nicoumalone			
Nifedipine			
Nifenazone			
Nikethamide			
Nilutamide			
Nimodipine			
Niridazole			
Nisoldipine			
Nitrendipine			
Nitrofurantoin			
Nitrofurazone			
Nizatidine			
Nomifensine Maleate			
Noradrenaline			
Noradrenaline Acid			
Tartrate			
Norethandrolone			
Norethisterone			
Norethisterone Acetate			
Norethisterone Norethisterone			
Heptanoate			
Norethynodrel			
Norfloxacin			
Norgestimate			
Norgestrel			
Nortriptyline			
Hydrochloride			
Noscapine			
Noscapine Hydrochloride			
Novobiocin Calcium			
Novobiocin Sodium			
Nux Vomica Seed			
Nystatin			
Octacosactrin			
Octacosactrin			
Octreolide Oestradiol			
Oestradiol Benzoate			
Oestradiol Cypionate			
Oestradiol Dipropionate			
Oestradiol Diundecanoate			

rescription Only Circumstances In Which Substances Are Not		
Medicine	Prescription Only Me	dicines
Oestradiol Enanthate		
Oestradiol		
Phenylpropionate		
Oestradiol Undecanoate		
Oestradiol Valerate		
Oestriol		
Oestriol		
Di-Hemi Succinate		
Oestrogenic Substances,		
Conjugated		
Oestrone		
Ofloxacin		
Olanzapine		
Olsalazine Sodium		
Omeprazole		
Omeprazole Magnesium		
Ondansetron		
Ondansetron		
Hydrochloride		
Orciprenaline Sulphate		
Orphenadrine Citrate		
Orphenadrine Citrate Orphenadrine		
Hydrochloride		
Ouabain		
Ovarian		
Gland, Dried		
Oxamniquine		
Oxandronolone		
Oxantel Pamoate		
Oxaprozin Oxatomide		
Oxedrine Tartrate		
Oxethazaine		
Oxidronate Sodium		
Oxitropium Bromide		
Oxolinic Acid		
Oxpentifylline		
Oxprenolol		Any use (except
Hydrochloride		local ophthalmic
		use)
Oxybuprocaine		
Hydrochloride		
Oxybutynin		
Hydrochloride		
Oxymetholone		
Oxypertine		

Prescription Only	Circumstances In W		re Not
Medicine	Prescription Only M	edicines	
Oxypertine			
Hydrochloride			
Oxyphenbutazone			
Oxyphencyclimine			
Hydrochloride			
Oxyphenonium Bromide			5 mg (MD) 15 mg (MDD)
Oxytetracycline			, , , , , , , , , , , , , , , , , , ,
Oxytetracycline Calcium			
Oxytetracycline			
Dihydrate			
Oxytetracycline			
Hydrochloride			
Oxytocin, Natural			
Oxytocin, Synthetic			
Pamidronate Disodium			
Pancreatin	(1) 21 000 European		(1) Capsules
rancieatiii	(1) 21,000 European Pharmacopoeia units		(1) Capsules
	of lipase per capsule		
	of fipase per capsule		
	(2) 25,000 European		(2) Powder
	Pharmacopoeia units		(2) Fowder
	of lipase per g		
Pancuronium Bromide	of fipase per g		
Pantoprazole			
Pantoprazole Sodium		(1) D : 1 1	
Papaverine		(1) By inhaler	(2) 50
		(2) Otherwise	(2) 50 mg
		than by inhaler	(MD)
			150 mg
D : 11 11 11		(1) D : 1 1	(MDD)
Papaverine Hydrochloride		(1) By inhaler	(2)
		(2) Otherwise	(2)
		than by inhaler	Equivalent of
			50 mg of
			Papaverine
			(MD)
			Equivalent of
			150 mg of
			Papaverine
D 1		A 6 ((MDD)
Paracetamol		Any form (except	
		non-effervescent	
		tablets and	
D 111 1		capsules)	
Paraldehyde			
Paramethadione			
Paramethasone Acetate			
Parathyroid Gland			

rescription Only Circumstances In Which Substances Are Not			re Not
Medicine	Prescription Only Mo	edicines	
Pargyline Hydrochloride			
Paroxetine Hydrochloride			
Pecilocin			
Penamecillin			
Penbutolol Sulphate			
Penciclovir			
Penicillinamine			
Penicillinamine			
Hydrochloride			
Pentamidine Isethionate			
Pentamidronate Disodium			
Penthienate			5 mg (MD)
Methobromide			15 mg (MDD)
Pentolinium Tartrate			
Perfluamine			
Pergolide Mesylate			
Perhexiline Maleate			
Perindopril			
Pericyazine			
Perindopril Erbumine			
Perphenazine			
Phenacetin	0.1%		
Phenazone	01270	External	
Phenazone and Caffeine			
Citrate			
Phenazone Salicylate			
Phenbutrazate			
Hydrochloride			
Phenelzine Sulphate			
Phenethicillin Potassium			
Phenformin			
Hydrochloride			
Phenglutarimide			
Hydrochloride			
Phenindione			
Phenolphthalein			
Phenoxybenzamine			
Hydrochloride			
Phenoxymethylpenicillin			
Phenoxymethylpenicillin			
Calcium			
Phenoxymethylpenicillin			
Potassium			
Phenprocoumon			
Phensuximide			
Phentolamine			
Hydrochloride			

Prescription Only Medicine	Circumstances Prescription On	In Which Substances Analy Medicines	re Not
Phentolamine Mesylate	•		
Phenylbutazone			
Phenylbutazone Sodium			
Phenylpropanolamine		Internal: (a) all	(a) 25 mg
Hydrochloride		preparations	(MD)
Trydrocmorae		(except	100 mg
		controlled release	(MDD)
		capsules, nasal	(11122)
		sprays or nasal	
		drops)	
		(b) controlled	(b) 50 mg
		release capsules	(MD)
		Total out out of	100 mg
			(MDD)
	(c) 2.0%	(c) nasal sprays	(2.22)
	(3) 2.3 / 0	or nasal drops	
Phenytoin		or masur drops	
Phenytoin Sodium			
Phthalylsulphathiazole			
Physostigmine			
Physostigmine			
Aminoxide Salicylate			
Physostigmine Salicylate			
Physostigmine Sulphate			
Phytomenadine		Any use except	
i nytomenadme		the prevention or	
		treatment of	
		haemorrhagic	
		disorders	
Picrotoxin		disorders	
Pilocarpine			
Pilocarpine			
Hydrochloride			
Pilocarpine Nitrate			
Pimozide			
Pindolol			
			5 ma (MD)
Pipenzolate Bromide			5 mg (MD)
Piperacillin Sodium			15 mg (MDD)
*			
Piperazine Oestrone			
Sulphate Piperidolate			50 mg (MD)
			50 mg (MD)
Hydrochloride			150 mg (MDD)
Pipothiazine Palmitate			(MIDD)
Piracetam			
Pirbuterol Acetate			
Pirbuterol Hydrochloride			

Prescription Only		In Which Substances	Are Not
Medicine	Prescription On	lly Medicines	
Pirenzepine			
Dihydrochloride			
Monohydrate			
Pirenzepine			
Hydrochloride			
Piretamide			
Piroxicam			
Piroxicam Beta-			
Cyclodextrin			
Pituitary Gland (Whole		By inhaler	
Dried)			
Pituitary, Powdered		By inhaler	
(Posterior Lobe)			
Pivampicillin			
Hydrochloride			
Pivmecillinam			
Pivmecillinam			
Hydrochloride			
Pizotifen			
Pizotifen Malate			
Plicamycin			
Podophyllotoxin			
Podophyllum			
Podophyllum Indian			
Podophyllum Resin	20.0%	External:	
1 odopnynum Resm	20.070	ointment or	
		impregnated	
		plaster	
Poldine Methylsulphate		praster	2 mg (MDD)
T ordine Wethy Burphace			6 mg (MDD)
Polidexide			o mg (MDD)
Polymyxin B Sulphate			
Polyestradiol Phosphate			
Polythiazide			
Poppy			
Capsule			
Potassium Arsenite	0.0127%		
Potassium Bromide	0.012770		
Potassium Canrenoate			
Potassium Clavulanate			
Potassium Perchlorate			
Practolol			
Pralidoxime Chloride			
Pralidoxime lodide			
Pralidoxime Mesylate			
Pramipexole Dibydrochloride			
Dihydrochloride Drayactatin Sadium			
Pravastatin Sodium			

Prescription Only	Circumstances In Wh	nich Substances Ai	re Not
Medicine	Prescription Only Me		
Prazosin Hydrochloride	•		
Prednisolone			
Prednisolone Acetate			
Prednisolone Butylacetate			
Prednisolone Hexanoate			
Prednisolone Pivalate			
Prednisolone Sodium			
Phosphate			
Prednisolone Sodium m-			
Sulphobenzoate			
Prednisolone 21-Steaglate			
Prednisolone m-			
Sulphobenzoate			
Prednisone			
Prednisone Acetate			
Prenalterol Hydrochloride			
Prenylamine Lactate			
Prilocaine Hydrochloride		Any use (except	
		local ophthalmic	
		use)	
Primidone			
Probenecid			
Probucol			
Procainamide			
Hydrochloride			
Procaine Hydrochloride		Any use (except	
		local ophthalmic	
		use)	
Procaine Penicillin			
Procarbazine			
Hydrochloride			
Prochlorperazine			
Prochlorperazine			
Edisylate			
Prochlorperazine Maleate			
Prochlorperazine			
Mesylate			
Procyclidine			
Hydrochloride			
Progesterone			
Prolactin			
Proligestone			
Prolintane Hydrochloride			
Promazine Embonate			
Promazine Hydrochloride			
Propafenone			
Propafenone			
Hydrochloride			
Propanidid			
- F	l	l	1

Prescription Only Medicine	Circumstances In W Prescription Only M		re Not
Propantheline Bromide			15 mg (MD)
•			45 mg (MDD)
Propiverine			
Hydrochloride			
Propofol			
Propranolol			
Hydrochloride			
Propylthiouracil			
Proquazone			
Protamine Sulphate			
Prothionamide			
Protirelin			
Protriptyline			
Hydrochloride			
Proxymetacaine		Any use (except	
Hydrochloride		local ophthalmic	
		use)	
Pseudoephedrine		Internal	In the case of
Hydrochloride			a controlled
			release
			preparation:
			120 mg
			(MD)
			180 mg
			(MDD)
			In any
			other case:
			60 mg
			(MD)
			180 mg
D 1 1 1			(MDD)
Pseudoephedrine			60 mg (MD)
Sulphate			180 mg
D (1E 1			(MDD)
Pyrantel Embonate			
Pyrantel Tartrate			
Pyrazinamide			
Pyridostigmine Bromide			
Pyrimethamine			
Quetiapine Fumarate			
Quinagolide			
Hydrochloride			
Quinapril			
Quinapril Hydrochloride			
Quinestradol			
Quinestrol			
Quinethazone	_		
Quinidine			

Prescription Only Medicine	Circumstances In Which Substances Are Not Prescription Only Medicines		
Quinidine Bisulphate			
Quinidine			
Polygalacturonate			
Quinidine Sulphate			
Quinine		100 mg (MD)	
		300 mg	
		(MDD)	
Quinine Bisulphate		Equivalent of	
		100 mg of	
		Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine	
		(MDD)	
Quinine Dihydrochloride		Equivalent of	
		100 mg of	
		Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine	
		(MDD)	
Quinine Ethyl Carbonate		Equivalent of	
		100 mg of	
		Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine	
		(MDD)	
Quinine Glycero-		Equivalent of	
phosphate		100 mg of	
		Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine	
0		(MDD)	
Quinine Hydrobromide		Equivalent of	
		100 mg of	
		Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine (MDD)	
Quinine Hydrocholride		Equivalent of	
Quilling right of the		100 mg of	
		Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine	
		(MDD)	
	1	(ממוא)	

Prescription Only	ly Circumstances In Which Substances Are Not		
Medicine	Prescription Only Medicines		
Quinine Iodobismuthate		Equivalent of	
		100 mg of	
		Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine	
		(MDD)	
Quinine		Equivalent of	
Phenylcinchoninate		100 mg of	
		Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine	
		(MDD)	
Quinine Phosphate		Equivalent of	
		100 mg of	
		Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine	
		(MDD)	
Quinine Salicylate		Equivalent of	
		100 mg of	
		Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine	
		(MDD)	
Quinine Sulphate		Equivalent of	
		100 mg of	
		Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine (MDD)	
Quinine Tannate		Equivalent of	
Quinnie Tainiate			
		100 mg of Quinine (MD)	
		Equivalent of	
		300 mg of	
		Quinine	
		(MDD)	
Quinine and Urea	+	(IVIDD)	
Hydrochloride			
Ramipril			
Ranitidine Bismuth			
Citrate			
Ranitidine Hydrochloride			
ramuume rryurocinoride			

Prescription Only	Circumstances In Wh		e Not
Medicine	Prescription Only Me	eaicines	
Rauwolfia Serpentina			
Rauwolfia Vomitoria			
Reboxetine			
Reboxetine Mesilate			
Remoxipride			
Hydrochloride			
Reproterol Hydrochloride			
Rescinnamine			
Reserpine			
Rifabutin			
Rifampicin			
Rifampicin Sodium			
Rifamycin			
Rimexolone			
Rimiterol Hydrobromide			
Risperidone			
Ritodrine Hydrochloride			
Ritonavir			
Rolitetracycline Nitrate			
Ropinorole			
Hydrochloride			
Sabadilla			
Salbutamol			
Salbutamol Sulphate			
Salcatonin			
Salcatonin Hydrated			
Polyacetate			
Salmefamol			
Salmeterol			
Hydroxynaphthoate			
Salsalate			
Saquinavir			
Saralasin Acetate			
Selegiline Hydrochloride			
Sera and Antisera –			
Botulin Antitoxin			
Diphtheria Antitoxin			
Gas-gangrene Antitoxin			
(Oedematiens)			
Gas-gangrene Antitoxin			
(Perfringens)			
Gas-gangrene Antitoxin			
(Septicum)			
Mixed Gas-gangrene			
Antitoxin			
Leptospira Antiserum			
Rabies Antiserum			
Navies Aliuseiulli	l		

Prescription Only Circumstances In Which Substances Are Not			e Not
Medicine	Prescription Only	Medicines	
Scorpion Venom			
Antiserum			
Snake Venom Antiserum			
Tetanus Antitoxin			
Serum Gonadotrophin			
Sermorelin			
Sertindole			
Sertraline Hydrochloride			
Sevoflurane			
Sibutramine			
Hydrochloride			
Silver Sulphadiazine			
Simvastatin			
Sissomicin			
Sissomicin Sulphate			
Snake Venoms			
Sodium Acetrizoate			
Sodium Aminosalicylate			
Sodium			
Antimonylgluconate			
Sodium Arsanilate			
Sodium Arsenate			
Sodium Arsenite	0.013%		
	0.013%		
Sodium Bromide			
Sodium Clodronate		A 1	
Sodium Cromoglycate		Administration through the nose	
Sodium Ethacrynate			
Sodium Fluoride	(1) 0.33% (y)	(1) Dentifrices	
	() () /	(2) Other	
		preparations for	
		use in the	
		prevention of	
		dental of dental	
		caries in the form	
		of –	
		(a)tablets or	(a) 2.2 mg
		drops;	(MDD)
	(b) 0.2%	(b) mouth-washes	, ,
	. '	(other than those	
		for daily use);	
	(c) 0.05%	(c) mouth-washes	
		for daily use	<u> </u>
Sodium Fusidate			
Sodium Metrizoate			
Sodium	1.14% (y)	Dentifrice	
Monofluorophosphate			
Sodium Stibogluconate			

Prescription Only Medicine	Circumstances In Prescription Only	n Which Substances Are Not y Medicines	
Sodium Valproate			
Somatorelin Acetate			
Somatrem			
Somatropin			
Sotalol Hydrochloride			
Sparfloxacin			
Spectinomycin			
Spectinomycin			
Hydrocloride			
Spiramycin			
Spiramycin Adipate			
Spironolactone			
Stannous Fluoride	(1) 0.62% (y)	(1) Dentifrice	
	(2) 0.4%	(2) Dental gels	
		for use in the	
		prevention and	
		treatment of	
		dental caries and	
		decalcification of	
		the teeth	
Stanolone			,
Stanozolol			,
Stavudine			,
Stilboestrol			
Stilboestrol Dipropionate			
Streptodornase		External	
Streptokinase		External	
Streptomycin			,
Streptomycin Sulphate			
Strychnine			
Strychnine Arsenate			,
Strychnine Hydrochloride			,
Strychnine Nitrate			,
Styramate			,
Succinylsulphathiazole			,
Sucralfate			,
Sulbactam			,
Sodium			
Sulbenicillin			
Sulbenicillin Sodium			
Sulbenicillin Tosylate			
Sulconazole Nitrate		External (except vaginal use)	
Sulfacytine			
Sulfadoxine			
Sulfamonomethoxine			
Sulindac			
Sulphabenzamide			

Prescription Only	Circumstances In Wh		e Not
Medicine	Prescription Only Me	dicines	
Sulphacetamide			
Sulphacetamide Sodium			
Sulphadiazine			
Sulphadiazine Sodium			
Sulphadimethoxine			
Sulphadimidine			
Sulphadimidine Sodium			
Sulphafurazole			
Sulphafurazole			
Diethanolamine			
Sulphaguanidine			
Sulphaloxic Acid			
Sulphamerazine			
Sulphamerazine Sodium			
Sulphamethizole			
Sulphamethoxazole			
Sulphamethoxydazine			
Sulphamethoxypyridazine			
Sulphamethoxypyridazine			
Sodium			
Sulphametopyrazine			
Sulphamoxole			
Sulphanilamide			
Sulphaphenazole			
Sulphapyridine			
Sulphapyridine Sodium			
Sulphasalazine			
Sulphathiazole			
Sulphathiazole Sodium			
Sulphaurea Sulphaurea			
Sulphinpyrazone			
Sulpiride			
Sultamicillin			
Sultamicillin Tosylate			
Sulthiame			
Sumatriptan Sussinata			
Sumatriptan Succinate			
Suprofen			
Suxamethonium Bromide			
Suxamethonium Chloride			
Suxethonium Bromide			
Tacalcitol Monohydrate			
Tacrine Hydrochloride			
Talampicillin			
Talampicillin			
Hydrochloride			
Talampicillin Napsylate			

Prescription Only	Circumstances In Wh	nich Substances Ar	e Not
Medicine	Prescription Only Me	edicines	
Tamoxifen			
Tamoxifen Citrate			
Tamsulosin			
Hydrochloride			
Tazarotene			
Tazobactam Sodium			
Teicoplanin			
Temocapril			
Hydrochloride			
Temocillin Sodium			
Tenoxicam			
Terazosan Hydrochloride			
Terbinafine			
Terbinafine			
Hydrochloride			
Terbutaline			
Terbutaline Sulphate			
Terfenadine			
Terodiline Hydrochloride			
Tertipressin			
Testosterone			
Testosterone Acetate			
Testosterone 17B Chloral			
Hemiacetal			
Testosterone Cyclohexyl-			
propionate			
Testosterone Cypionate			
Testosterone Decanoate			
Testosterone Enanthate			
Testosterone Isocaproate			
Testosterone			
Phenylpropionate			
Testosterone Propionate			
Testosterone			
Undecanoate			
Tetrabenazine			
Tetracosactrin			
Tetracosactrin Acetate			
Tetracycline			
Tetracycline			
Hydrochloride			
Tetracycline Phosphate			
Complex			
Tetroxoprim			
Thallium Acetate			
Thallous Chloride			
Thiabendazole			
Thiambutosine			

Medicine Prescription Only Medicines Thiethylperazine Maleate Thiocarlide Thiogaunine Thiopentone Sodium Thiopropazate Hydrochloride Thioridazine Thiothixene Thiothixene Thiothixene Thiothixene Thiothixene Thiotry and the state of the stat	Prescription Only	Circumstances In Wi		e Not
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Tolazoline Hydrochloride External Tolbutamide External				
Tolbutamide			External	
	-			
	Tolbutamide Sodium			

Prescription Only	Circumstances In Wi		re Not
Medicine	Prescription Only Me	edicines	
Tolcapone			
Tolfenamic Acid			
Tolmetin Sodium			
Topiramate			
Torasemide			
Toremifene			
Tramadol Hydrochloride			
Trandolapril			
Tranexamic Acid			
Tranylepromine Sulphate			
Trazodone Hydrochloride			
Treosulfan			
Tretinoin			
Triamcinolone			
Triamcinolone Acetonide			
Triamcinolone Diacetate			
Triamcinolone			
Hexacetonide			
Triamterene			
Tribavirin			
Triclofos Sodium			
Trientine Dihydrochloride			
Trifluoperazine			
Trifluoperazine			
Hydrochloride			
Trifluperidol			
Trifluperidol			
Hydrochloride			
Trilostane			
Trimeprazine			
Trimeprazine Tartrate			
Trimetaphan Camsylate			
Trimetazidine			
Trimetazidine			
Hydrochloride			
Trimethoprim			
Trimipramine Maleate			
Trimipramine Mesylate			
Tropicamide			
Tropisetron			
Hydrochloride			
Troxidone			
L-Tryptophan		(1) Dietary or	
VI I		nutritive use	
		(2) Any external	
		use	
Tubocurarine Chloride			
Tulobuterol			
		1	

Prescription Only	Circumstances In Wh	nich Substances Ar	e Not
Medicine	Prescription Only Me	edicines	
Tulobuterol			
Hydrochloride			
Tyrothricin		Throat lozenges	
		or throat pastilles	
Uramustine			
Urea Stibamine			
Urethane			
Uridine-5-Triphosphoric			
Acid			
Urofollitrophin			
Urokinase			
Ursodeoxychloric Acid			
Vaccines –			
Athrax Vaccine			
(Bacillus) Anthracis)			
Bacillus Calmette-			
Guerin Vaccine			
Bacillus Salmonella			
Typhi Vaccine			
Percutaneous Bacillus			
Calmette- Guerin			
Vaccine			
Cholera Vaccine			
Diphtheria Vaccine			
Adsorbed Diphtheria			
Vaccine			
Diphtheria and Tetanus			
Vaccine			
Adsorbed Diphtheria			
and Tetanus Vaccine			
Diphtheria, Tetanus			
and Pertussis Vaccine			
Adsorbed Diphtheria,			
Tetanus and Pertussis			
Vaccine			
Diphtheria, Tetanus			
and Poliomyelitis			
Vaccine			
Diphtheria, Tetanus,			
Pertussis and			
Poliomyelitis Vaccine			
Eltor Vaccine			
Influenza Vaccine			
Hepatitis B Vaccine			
Measles Vaccine			
(Live Attenuated)			

Prescription Only	Circumstances In Whi	ch Substances Ar	e Not
Medicine	Prescription Only Med	licines	
Meningococcal			
Polysaccharide			
Vaccine			
Mumps Vaccine			
Pertussis Vaccine			
Plague Vaccine			
Pneumococcal Vaccine			
(Bacterial Antigen)			
Poliomyelitis Vaccine			
(Inactivated)			
Poliomyelitis Vaccine			
(Live Oral)			
Rabies Vaccine			
Rubella Vaccine (Live			
Attenuated)			
Rubella, Mumps,			
Measles Vaccine			
Tetanus Vaccine			
Adsorbed Tetanus			
Vaccine			
Tetanus and Pertussis			
Vaccine			
Tuberculin Purified			
Protein Derivative			
Old Tuberculin			
Typhoid Vaccine			
Typhoid – Paratyphoid			
A and B Vaccine			
Typhoid – Paratyphoid			
A and B and Cholera			
Vaccine			
Typhoid – Paratyphoid			
A and B and Tetanus			
Vaccine			
Typhus Vaccine			
Yellow Fever Vaccine			
Valaciclovir			
Valaciclovir			
Hydrochloride			
Valproic Acid			
Valsartan			
Vancomycin			
Hydrochloride			
Vasopressin Injection			
Vasopressin Tannate			
Vecuronium Bromide			
Venlafaxine			
Venlafaxine			
Hydrochloride			

Prescription Only	Circumstances In	1 Which Substances	Are Not
Medicine	Prescription Only	y Medicines	
Verapamil Hydrochloride			
Veratrine			
Veratrum			
(Green and White)			
Vidarabine			
Vigabatrin			
Viloxazine Hydrochloride			
Vinblastine Sulphate			
Vincristine Sulphate			
Vindesine Sulphate			
Viomycin Pantothenate			
Viomycin Sulphate			
Vitamin A		(1) Internal	(1) 7500 iu (2250 mcg Retinol equivalent) (MDD)
		(2) External	
Vitamin A Acetate		(1) Internal	(1) Equivalent to 7500 iu Vitamin A (2250 mcg Retinol equivalent) (MDD)
		(2) External	
Vitamin A Palmitate		(1) Internal	(1) Equivalent to 7500 iu Vitamin A (2250 mcg Retinol equivalent) (MDD)
		(2) External	
Warfarin			
Warfarin Sodium			
Xamoterol Fumarate			
Xipamide			
Yohimbine			
Hydrochloride			
Zalcitabine			
Zidovudine			
Zimeldine Hydrochloride			
Zolmitriptan			
Zolpidem			
Zomepirac Sodium			
Zopiclone			
Zuclopenthixol Acetate			

Prescription Only	Circumstances In Which Substances	s Are Not		
Medicine	Prescription Only Medicines			
Zuclopenthixol				
Decanoate				
Zuclopenthixol				
Hydrochloride				
Note –	1. In relation to a medicinal product th	at contains more		
	than one of the substances Atropine, A			
	Methobromide, Atropine Methonitrate			
	Hydrochloride, Atropine Sulphate, Hydrochloride, Atropine Sulphate, Hydrochloride, Marchael Sulphate, Hydrochloride, Atropine Sulphate, Hydrochloride, Atropine Sulphate, Hydrochloride, Marchael Sulphate, Marchael Sulpha	Hydrochloride, Atropine Sulphate, Hyoscine, Hyoscine		
		Butylbromide, Hyoscine Hydrobromide, Hyoscine		
	Methobromide, Hyoscine Methonitrate	Methobromide, Hyoscine Methonitrate, Hyoscyamine,		
	Hyoscyamine Hydrobromide and Hyoscyamine Sulphate,			
	the maximum daily dose for the purposes of column 4 is			
	1 mg of the total alkaloids contained in the product that			
	are derived from Belladonna, Hyoscyamus, Stramonium			
	or other solanaceous plant, and there is no maximum			
	dose.			
	2. In relation to a medicinal product the			
	than one of the substances Sodium Flu	·		
	Monofluorophosphate and Stannous F			
	in a dentifrice, the maximum strength			
		for the purposes of column 2 shall not exceed 0.15%		
	calculated as Fluorine.			

PART 2

(Articles 1(2), 3(2) and 7)

Controlled Drugs	Circumstances In Which Controlled Drugs Are Not		
	Prescription Only Medicines		
Column 1	Column 2	Column 3	Column 4
Substance	Maximum	Pharmaceutical	Maximum dose
	strength	form	
Codeine; its salts	Equivalent of 1.5%		Equivalent of 20
	of Codeine		mg of Codeine
	Monohydrate		Monohydrate
Dihydrocodeine; its	Equivalent of		Equivalent of 10
salts	1.5% of		mg of Dihydroco-
	Dihydrocodeine		deine
Ethylmorphine; its	Equivalent of 0.2%		Equivalent of 7.5
salts	of Ethylmorphine		mg of Ethyl-
			morphine
Morphine; its salts	(1) Equivalent of	(1) Liquid	(1) Equivalent of 3
	0.02% of anhydrous		mg of anhydrous
	Morphine		Morphine
	(2) Equivalent of	(2) Solid	(2) Equivalent of 3
	0.04% of anhydrous		mg of anhydrous
	Morphine;		Morphine
	equivalent of 300		
	mcg of anhydrous		
	Morphine		
Medicinal Opium	(1) Equivalent of	(1) Liquid	(1) Equivalent of 3
	0.02% of anhydrous		mg of anhydrous
	Morphine		Morphine
	(2) Equivalent of	(2) Solid	(2) Equivalent of 3
	0.04% of anhydrous		mg of anhydrous
	Morphine		Morphine
Pholcodine; its salts	Equivalent of 1.5%		Equivalent of 20
	of Pholcodine		mg of Pholcodine

PART 3

(Article 3(4))

NAMED PRODUCTS THAT ARE NOT PRESCRIPTION ONLY MEDICINES

TABLE A		
Name and product licence number of medicinal products that are not prescription only medicines		
Adcortyl in Orabase for Mouth Ulcers	0034/0321	
Anusol Plus HC Ointment	0018/0223	
Anusol Plus HC Suppositories	0018/0224	
Beechams Hydrocortisone Cream	0079/0203	
Boots Hydrocortisone Ointment	0014/0364	
Calacort Cream	12650/0001	
Canesten Hydrocortisone Cream	0010/0216	
Corlan Pellets	0039/0397	
Cortaid Cream 1%	0032/0126	
Corteze Cream	0001/0107	
Cortiderm	2855/0010	
Cortril Topical Ointment 1% (non-greasy)	0057/0251	
Dermacort Hydrocortisone Cream	8265/0002	
Dioderm Hydrocortisone Cream	0173/0153	
Efcortelan Eczema Cream	10949/0234	
Efcortelan Eczema Ointment	10949/0235	
Eurax HC Cream	0001/5010R	
Hc45 Hydrocortisone Cream	0327/0039	
Herpetad Cold Sore Cream	4986/0007	
Jungle Formula Bite & Sting Relief Cream	2855/0010	
Lanacort Cream	3157/0008	
Lanacort Ointment	3157/0011	
Perinal Spray	0173/0049	
Pharmacort Cream 0.5%	0011/0077	
Proctocream HC	0036/0065	
Soothelip Cold Sore Cream	0142/0426	
Timocort Hydrocortisone Cream	0044/0090	
Timocort Hydrocortisone Cream 1%	0063/0076	
Wasp-Eze Hydrocortisone Cream	8452/0012	
Zaclovir Cold Sore Cream	4986/0007	
Zenoxone Cream	0181/0033	
Zovirax Cold Sore Cream	0003/0304	

TABLE B	
Relevant product licence holder and name and medicinal products that are not prescri	-
Leo Laboratories Limited:	
Hydrocortisone Acetate Cream BP 0.5%	0043/0150
Hydrocortisone Acetate Cream BP 1.0%	0043/0151
Richard Daniel and Son Limited:	
Hydrocortisone Cream BP 1.0%	0842/0011

PART 428

(Article 3(4))

OTHER MEDICINAL PRODUCTS THAT ARE NOT PRESCRIPTION ONLY MEDICINES

- A medicinal product shall not be a prescription only medicine by reason that it contains the substance aciclovir, where
 - (a) the maximum strength of the aciclovir in the medicinal product does not exceed 5%;
 - (b) the medicinal product is sold or supplied in a container, or package, containing not more than 2 g of the medicinal product; and
 - (c) the medicinal product is indicated only for external application for the treatment of herpes simplex virus infections of the lips and face (Herpes labialis).
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance acrivastine, where
 - (a) the medicinal product is sold or supplied in a container, or package, containing not more than 240 mg of acrivastine; and
 - (b) the container or package is labelled to show a maximum daily dose of 24 mg of acrivastine.
- 3 A medicinal product shall not be a prescription only medicine by reason that it contains the substance aloxiprin, where
 - (a) the medicinal product is in the form of non-effervescent tablets or capsules;
 - (b) the maximum strength of the medicinal product in each tablet or capsule does not exceed 620 mg;
 - (c) the quantity (of tablets or capsules, or of any combination of tablets and capsules) that is sold or supplied in one container or package does not exceed 32; and
 - (d) the quantity (of tablets or capsules, or of any combination of tablets and capsules) that is sold or supplied to a person at any one time does not exceed 100.
- 4 A medicinal product shall not be a prescription only medicine by reason that it contains the substance aloxiprin, where it is not in the form of a non-effervescent tablet or capsule.
- 5 A medicinal product shall not be a prescription only medicine by reason that it contains the substance aspirin, where
 - (a) the medicinal product is in the form of non-effervescent tablets or capsules;
 - (b) the maximum strength of the medicinal product in each tablet or capsule does not exceed 75 mg;

- (c) the quantity (of tablets or capsules, or of any combination of tablets and capsules) that is sold or supplied in one container or package does not exceed 100; and
- (d) the quantity (of tablets or capsules, or of any combination of tablets and capsules) that is sold or supplied to a person at any one time does not exceed 100.
- **6** A medicinal product shall not be a prescription only medicine by reason that it contains the substance aspirin, where
 - (a) the medicinal product is in the form of non-effervescent tablets or capsules;
 - (b) the maximum strength of the medicinal product in each tablet or capsule does not exceed 325 mg;
 - (c) the quantity (of tablets or capsules, or of any combination of tablets and capsules) that is sold or supplied in one container or package does not exceed 32; and
 - (d) the quantity (of tablets or capsules, or of any combination of tablets and capsules) that is sold or supplied to a person at any one time does not exceed 100.
- 7 A medicinal product shall not be a prescription only medicine by reason that it contains the substance azelastine hydrochloride, where
 - (a) the medicinal product is in non-aerosol, aqueous form for nasal administration;
 - (b) the medicinal product is sold or supplied in a container, or package, containing not more than 36 doses each of which contains not more than 140 mcg of azelastine hydrochloride;
 - (c) the container or package is labelled to show a maximum dose of 140 mcg per nostril and a maximum daily dose of 280 mcg per nostril of azelastine hydrochloride; and
 - (d) the medicinal product is indicated only for the treatment of seasonal allergic rhinitis or perennial allergic rhinitis, in persons aged not less than 5 years.
- **8** A medicinal product shall not be a prescription only medicine by reason that it contains the substance azelastine hydrochloride, where
 - (a) the medicinal product is in the form of eye drops; and
 - (b) it is indicated only for the treatment of allergic conjunctivitis, in persons aged not less than 12 years.
- 9 A medicinal product shall not be a prescription only medicine by reason that it contains the substance beclomethasone dipropionate, where
 - (a) the medicinal product is in non-aerosol form for nasal administration;
 - (b) the medicinal product is sold or supplied in a container, or package, containing not more than 5,600mcg of beclomethasone diopropionate;

- (c) the container or package is labelled to show a maximum dose of 100 mcg per nostril and a maximum daily dose of 200 mcg per nostril of beclomethasone dipropionate; and
- (d) the medicinal product is indicated only for the prevention of treatment of allergic rhinitis, in persons aged not less than 12 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance budesonide, where
 - (a) the medicinal product is in non-aerosol, aqueous form for nasal administration;
 - (b) the medicinal product is sold or supplied in a container, or package, containing not more than 10 mg of the medicinal product;
 - (c) the container or package is labelled to show a maximum dose, and a maximum daily dose, of 200 mcg per nostril of budesonide; and
 - (d) the medicinal product is indicated only for the prevention of treatment of seasonal allergic rhinitis, in persons aged not less than 12 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance carbenoxolone sodium, where
 - (a) the medicinal product is in the form of granules;
 - (b) the maximum strength of the carbenoxolone sodium in the medicinal product does not exceed 1%, calculated in terms of weight in weight;
 - (c) the medicinal product is sold or supplied in a container, or package, containing not more than 560 mg of carbenoxolone sodium;
 - (d) the container or package is labelled to show a maximum dose of 20 mg and a maximum daily dose of 80 mg of carbenoxolone sodium; and
 - (e) the medicinal product is indicated only for treatment by mouthwash, in persons aged not less than 12 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance cetirizine, where
 - (a) the medicinal product is sold or supplied in a container, or package, containing not more than 100 mg of cetirizine; and
 - (b) the container or package is labelled to show a maximum daily dose of 10mg of cetirizine.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance cimetidine, where
 - (a) the container or package in which the medicinal product is sold or supplied is labelled to show a maximum dose of 200 mg and a maximum daily dose of 800 mg of cimetidine for a maximum period of 14 days; and
 - (b) the medicinal product is indicated for the short-term symptomatic relief of heartburn, dyspepsia and hyperacidity and for the prophylaxsis of meal-induced heartburn.

- A medicinal product shall not be a prescription only medicine by reason that it contains the substance cimetidene, where
 - (a) the medicinal product is for the prophylactic management of nocturnal heartburn; and
 - (b) the container or package in which the medicinal product is sold or supplied is labelled to show a maximum dose of 100 mg of cimetidine to be taken once daily at night for a maximum period of 14 days.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance clobetasone butyrate, where
 - (a) the medicinal product is in the form of a cream;
 - (b) the maximum strength of the clobetasone butyrate in the medicinal product does not exceed 0.05%;
 - (c) the medicinal product is sold or supplied in a container, or package, containing not more than 15 g of the medicinal product; and
 - (d) the medicinal product is indicated only for external application for the short-term treatment of eczema and dermatitis, in persons aged not less than 12 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance diclofenac diethylammonium, where
 - (a) the maximum strength of the diclofenac diethylammonium in the medicinal product does not exceed 1.16%, calculated in terms of weight in weight;
 - (b) the medicinal product is sold or supplied in a container, or package, containing not more than 30 g of the medicinal product;
 - (c) the container or package is labelled to show a maximum period of use of 7 days; and
 - (d) the medicinal product is indicated for external application for the local symptomatic relief of pain and inflammation in trauma of the tendons, ligaments, muscles and joints and in localized forms of soft tissue rheumatism, in persons aged not less than 12 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance domperidone, where
 - (a) the medicinal product is indicated for the relief of post-prandial symptoms of excessive fullness, nausea, epigastric bloating and belching, occasionally accompanied by epigastric discomfort and heartburn;
 - (b) the medicinal product is sold or supplied in a container or package containing not more than 200 mg of domperidone; and
 - (c) the container or package is labelled to show a maximum dose of 10 mg of domperidone and a maximum daily dose of 40 mg of domperidone.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance domperidone maleate, where –

- (a) the medicinal product is sold in a container, or package, containing not more than 200 mg of domperidone maleate;
- (b) the container or package is labelled to show a maximum dose of 10 mg and a maximum daily dose of 40 mg; and
- (c) the medicinal product is indicated for use for the relief of postprandial symptoms of excessive fullness, nausea, epigastric bloating and belching, accompanied by epigastric discomfort and heartburn.
- 19 A medicinal product shall not be a prescription only medicine by reason that it contains the substance famotidine, where
 - (a) the container or package in which the medicinal product is sold or supplied is labelled to show a maximum dose of 10 mg and a maximum daily dose of 20 mg of famotidine for a maximum period of 14 days; and
 - (b) the medicinal product is indicated for
 - (i) the short-term symptomatic relief of heartburn, dyspepsia, indigestion, acid indigestion or hyperacidity, or
 - (ii) the prevention of the symptoms of heartburn, dyspepsia, indigestion, acid indigestion or hyperacidity where they are associated with the consumption of food or drink, including the prevention of sleep disturbance because of those symptoms.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance felbinac, where
 - (a) the maximum strength of the felbinac in the medicinal product does not exceed 3.17%, calculated in terms of weight in weight;
 - (b) the medicinal product is sold or supplied in a container, or package, containing not more than 50 g of the medicinal product;
 - (c) the container or package is labelled to show a maximum period of use of 7 days; and
 - (d) the medicinal product is indicated for external application for the relief of symptoms associated with soft tissue injury such as strains, sprains and contusions, in persons aged not less than 12 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance fluconazole, where
 - (a) the medicinal product is sold or supplied in a container, or package, containing not more than 150 mg of the medicinal product;
 - (b) the container or package is labelled to show a maximum dose of 150 mg of fluconazole; and
 - (c) the medicinal product is indicated for oral administration for the treatment of vaginal candidiasis or associated candidal balanitis, in persons aged not less than 16 years but less than 60 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance flunisolide, where –

- (a) the medicinal product is in the form of a non-pressurized nasal spray;
- (b) the maximum strength of the flunisolide in the medicinal product does not exceed 0.025%, calculated in terms of weight in volume;
- (c) the medicinal product is sold or supplied in a container, or package, containing not more than 240 metered doses of the medicinal product;
- (d) the container or package is labelled to show a maximum dose of 50 mcg per nostril and a maximum daily dose of 100 mcg per nostril of flunisolide in the case of persons aged not less than 16 years, and a maximum dose of 25 mcg per nostril and a maximum daily dose of 75 mcg per nostril in the case of children aged not less than 12 years but less than 16 years; and
- (e) the medicinal product is indicated for the prevention and treatment of seasonal allergenic rhinitus, including hay fever, in persons aged not less than 12 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance hydrocortisone, where
 - (a) the maximum strength of the hydrocortisone in the medicinal product does not exceed 0.5%, calculated in terms of weight in weight;
 - (b) the medicinal product is sold or supplied in a container, or package, containing not more than 15 g of the medicinal product; and
 - (c) the medicinal product is indicated for external use in combination with nystatin of a maximum strength of 3.0%, for intertrigo, in persons aged not less than 10 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance hydrocortisone, where
 - (a) the medicinal product is in the form of a cream, ointment or spray;
 - (b) the maximum strength of the hydrocortisone in the medicinal product does not exceed 1.0%, calculated in terms of weight in weight;
 - (c) the medicinal product is sold or supplied in a container, or package, containing
 - (i) where the medicinal product is in the form of a cream or ointment, not more than 15 g of the medicinal product, or
 - (ii) where the medicinal product is in the form of a spray, not more than 30 ml of the medicinal product;
 - (d) the medicinal product is indicated for external use, either alone or in conjunction with crotamiton in irritant dermatitis, contact allergic dermatitis, insect bite reactions or mild to moderate eczema, and either in combination with clotrimazole or miconazole nitrate for athlete's foot and candidal intertrigo or in combination with lignocaine for anal and perianal itch associated with haemorrhoids; and

- (e) the medicinal product is indicated for use in persons aged not less than 10 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance hydrocortisone acetate, where
 - (a) the medicinal product is in the form of a cream or ointment, or suppositories;
 - (b) the maximum strength of the hydrocortisone acetate in the medicinal product is equivalent to 1.0% of hydrocortisone, calculated in terms of weight in weight;
 - (c) the medicinal product is sold or supplied in a container, or package, containing
 - (i) where the medicinal product is in the form of a cream or ointment, not more than 15 g of the medicinal product, or
 - (ii) where the medicinal product is in the form of suppositories, not more than 12 suppositories;
 - (d) the medicinal product is indicated for external use -
 - (i) for irritant dermatitis, contact allergic dermatitis, insect bite reactions or mild to moderate eczema,
 - (ii) in combination with one or more of the following, namely benzyl benzoate, bismuth oxide, bismuth subgallate, peru balsam, pramoxine hydrochloride and zinc oxide, for haemorrhoids, or
 - (iii) in combination with miconazole nitrate, for tinea pedis or candidal intertrigo; and
 - (e) the medicinal product is indicated for use in persons aged not less than 10 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance hydrocortisone sodium succinate, where
 - (a) the medicinal product is in the form of pellets;
 - (b) the maximum strength of the hydrocortisone sodium succinate in the medicinal product is equivalent to 2.5 mg of hydrocortisone, calculated in terms of weight in weight;
 - (c) the medicinal product is sold or supplied in a container, or package, containing the equivalent of 50 mg of hydrocortisone; and
 - (d) the medicinal product is indicated for external use for aphthous ulceration of the mouth, in persons aged not less than 12 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance hydroxyzine hydrochloride, where
 - (a) the medicinal product is sold or supplied in a container, or package, containing not more than 750 mg of the medicinal product;
 - (b) the container or package is labelled to show a maximum dose of 25 mg, and to show a maximum daily dose of 75 mg in the case of persons aged not less than 12 years and a maximum daily dose of 50 mg in the case of children aged not less than 6 years but less than 12 years; and

- (c) the medicinal product is indicated for the management of pruritus associated with acute or chronic urticaria or atopic dermatitis or contact dermatitis, in persons aged not less than 6 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance hyoscine butylbromide, where
 - (a) the route of administration of the medicinal product is internal and is otherwise than by means of an inhaler;
 - (b) the medicinal product is sold or supplied in a container, or package, containing not more than 240 mg of the medicinal product; and
 - (c) the container or package is labelled to show a maximum dose of 20 mg and a maximum daily dose of 80 mg of the medicinal product.
- 29 A medicinal product shall not be a prescription only medicine by reason that it contains the substance Ibuprofen, where
 - (a) the medicinal product is indicated for the relief of rheumatic and muscular pain, pain of non-serious arthritic conditions, backache, neuralgia, migraine, headache, dental pain, dysmenorrhoea, feverishness, symptoms of colds and influenza; and either
 - (b) the route of the administration of the medicinal product is internal; and
 - (i) in the case of a prolonged release preparation the container or package is labelled to show a maximum dose of 600 mg and a maximum daily dose of 1200 mg, or
 - (ii) in any other case the container or package is labelled to show a maximum dose of 400 mg and a maximum daily dose of 1200 mg; or
 - (c) the route of administration of the medicinal product is external; and
 - (i) the maximum strength of the Ibuprofen in the medicinal product does not exceed 5%, or
 - (ii)
- (A) the maximum strength of the Ibuprofen in the medicinal product does not exceed 10%, and
- (B) the medicinal product is sold or supplied in a container or package containing not more than 50 g of medicinal product which is labelled to show a maximum dose of 125 mg and a maximum daily dose of 500 mg.
- 30 A medicinal product shall not be a prescription only medicine by reason that it contains the substance ketoconazole, where
 - (a) the medicinal product is in the form of a shampoo;
 - (b) the maximum strength of the ketoconazole in the medicinal product does not exceed 2%, calculated in terms of weight in weight;
 - (c) the medicinal product is sold or supplied in a container, or package, containing not more than 120 ml of the medicinal product

- and containing in the medicinal product not more than 2,400 mg of ketoconazole;
- (d) the container or package is labelled to show a maximum frequency of application of once every 3 days; and
- (e) the medicinal product is indicated for the prevention and treatment of dandruff and seborrhoeic dermatitis of the scalp.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance ketoprofen, where
 - (a) the maximum strength of the ketoprofen in the medicinal product does not exceed 2.5%, calculated in terms of weight in weight;
 - (b) the medicinal product is sold or supplied in a container, or package, containing not more than 30 g of the medicinal product; and
 - (c) the medicinal product is indicated only for treatment by external topical application, for rheumatic and muscular pain, in persons aged not less than 12 years, for a maximum period of 7 days.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance levocabastine hydrochloride, where
 - (a) the medicinal product is in the form of a nasal spray;
 - (b) the maximum strength of the medicinal product does not exceed the equivalent of 0.05% levocabastine;
 - (c) the medicinal product is sold or supplied in a container, or package, containing not more than 10 ml of the medicinal product; and
 - (d) the medicinal product is indicated for the symptomatic treatment of seasonal allergic rhinitis.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance levocabastine hydrochloride, where
 - (a) the medicinal product is in the form of aqueous eye drops;
 - (b) the maximum strength of the medicinal product does not exceed the equivalent of 0.05% levocabastine;
 - (c) the medicinal product is sold or supplied in a container, or package, containing not more than 4ml of the medicinal product; and
 - (d) the medicinal product is indicated for the symptomatic treatment of seasonal allergic conjunctivitis.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance loratedine, where
 - (a) the medicinal product is sold or supplied in a container, or package, containing not more than 100 mg of loratadine; and
 - (b) the container or package is labelled to show a maximum daily dose of 10 mg of loratadine.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance mebendazole, where –

- (a) the medicinal product is sold or supplied in a container, or package, containing not more than 800 mg of mebendazole;
- (b) the container or package is labelled to show a maximum dose of 100 mg of mebendazole; and
- (c) the medicinal product is indicated for oral use in the treatment of enterobiasis, in persons aged not less than 2 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance nedocromil sodium, where
 - (a) the maximum strength of the nedocromil sodium in the medicinal product does not exceed 2.0%, calculated in terms of weight in volume;
 - (b) the medicinal product is sold in a container, or package, containing not more than 3 ml of the medicinal product; and
 - (c) the medicinal product is indicated for the prevention, relief and treatment of seasonal and perennial allergic conjunctivitis.
- 37 A medicinal product shall not be a prescription only medicine by reason that it contains the substance nizatidine, where
 - (a) the container or package in which the medicinal product is sold or supplied is labelled to show a maximum dose of 75 mg of nizatidine and a maximum of 4 such doses in any period of 14 days; and
 - (b) the medicinal product is indicated only for the prevention of the symptoms of food-related heartburn, in persons aged not less than 16 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance nystatin, where
 - (a) the maximum strength of the nystatin in the medicinal product does not exceed 3.0%, calculated in terms of weight in weight;
 - (b) the medicinal product is sold in a container, or package, containing not more than 15 g of the medicinal product; and
 - (c) the medicinal product is indicated for external use in combination with hydrocortisone of a maximum strength of 0.5% for intertrigo, in persons aged not less than 10 years.
- 39 A medicinal product shall not be a prescription only medicine by reason that it contains the substance oxethazaine, where
 - (a) the medicinal product is sold or supplied in a container, or package, containing not more than 400 ml of oxethazaine; and
 - (b) the container or package is labelled to show a maximum dose of 10 ml and a maximum daily dose of 30 ml of oxethazaine.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance paracetamol, where
 - (a) the medicinal product is in the form of non-effervescent tablets or capsules;

- (b) the maximum strength of the medicinal product in each tablet or capsule does not exceed 500 mg;
- (c) the quantity (of tablets or capsules, or of any combination of tablets and capsules) that is sold or supplied in one container or package does not exceed 32:
- (d) the quantity (of tablets or capsules, or of any combination of tablets and capsules) that is sold or supplied to a person at any one time does not exceed 100; and
- (e) the medicinal product is indicated for use by administration wholly or mainly to persons aged not less than 12 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance paracetamol, where
 - (a) the medicinal product is in the form of non-effervescent tablets or capsules;
 - (b) the maximum strength of the medicinal product in each tablet or capsule does not exceed 250 mg;
 - (c) the quantity (of tablets or capsules, or of any combination of tablets and capsules) that is sold or supplied in one container or package does not exceed 32; and
 - (d) the quantity (of tablets and capsules, or of any combination of tablets and capsules) that is sold or supplied to a person at any one time does not exceed 100.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance piroxicam, where
 - (a) the maximum strength of the piroxicam in the medicinal product does not exceed 0.5%;
 - (b) the medicinal product is sold or supplied in a container, or package, containing not more than 30 mg of the medicinal product;
 - (c) the container or package is labelled to show a maximum period of use of 7 days; and
 - (d) the medicinal product is indicated for external application 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, in persons aged not less than 12 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance prochlorperazine maleate, where
 - (a) the medicinal product is in the form of tablets;
 - (b) the maximum amount of the prochlorperazine maleate in the medicinal product, in each tablet, does not exceed 3 mg;
 - (c) the medicinal product is sold or supplied in a container, or package, containing not more than 8 tablets; and
 - (d) the medicinal product is indicated only for nausea, and vomiting, in previously diagnosed migraine, in persons aged not less than 18 years.

- A medicinal product shall not be a prescription only medicine by reason that it contains the substance pyrantel embonate, where
 - (a) the medicinal product is sold or supplied in a container, or package, containing not more than 750 mg of the medicinal product;
 - (b) the container or package is labelled to show a maximum daily dose (to be taken as a single dose) of pyrantel embonate of 750 mg in the case of persons aged not less than 12 years, of 500 mg in the case of children aged not less than 6 years but less than 12 years, and of 250 mg in the case of children aged not less than 2 years but less than 6 years; and
 - (c) the medicinal product is indicated for the treatment of enterobiasis, in persons aged not less than 2 years.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance ranitidine hydrochloride, where
 - (a) the container or package in which the medicinal product is sold or supplied is labelled to show a maximum dose equivalent to 75 ml and a maximum daily dose equivalent to 300 ml of ranitidine for a maximum period of use of 14 days; and
 - (b) the medicinal product is indicated for the short-term symptomatic relief of heartburn, dyspepsia and hyperacidity, or the prevention of those symptoms when associated with the consumption of food and drink.
- A medicinal product shall not be a prescription only medicine by reason that it contains the substance sodium cromoglycate, where
 - (a) the medicinal product is in the form of aqueous eye drops;
 - (b) the maximum strength of the sodium cromoglycate in the medicinal product does not exceed 2%, calculated in terms of weight in volume;
 - (c) the medicinal product is sold or supplied in a container containing not more than 10 ml of the medicinal product; and
 - (d) the medicinal product is indicated for treatment of acute seasonal allergic conjunctivitis.
- 47 A medicinal product shall not be a prescription only medicine by reason that it contains the substance sodium cromoglycgate, where
 - (a) the medicinal product is in the form of an eye ointment;
 - (b) the maximum strength of the sodium cromoglycate in the medicinal product is 4%, calculated in terms of weight in weight;
 - (c) the medicinal product is sold or supplied in a container, or package, containing not more than 5 g of the medicinal product; and
 - (d) the medicinal product is indicated for the treatment of acute seasonal allergic conjunctivitis or perennial allergic conjunctivitis.
- A medicinal product shall not be a prescription only medicine by reason of the fact that it contains terbinafine, where –

- (a) the maximum strength of the terbinafine in the medicinal product does not exceed 1%;
- (b) the medicinal product is sold or supplied in a container, or package, containing not more than 30 g of the medicinal product; and
- (c) the medicinal product is indicated for external use as a gel for the treatment of tina corporis, tinea pedis and tinea cruris.
- 49 A medicinal product shall not be a prescription only medicine by reason of the fact that it contains terbinafine hydrochloride where
 - (a) the maximum strength of the terbinafine hydrochloride in the medicinal product does not exceed 1%;
 - (b) the medicinal product is indicated for external use for the treatment of tinea pedis and tinea cruris; and
 - (c) the medicinal product is sold or supplied in a container or package containing not more than 15 g of medicinal product.
- A medicinal product shall not be a prescription only medicine by reason of the fact that it contains terbinafine hydrochloride, where
 - (a) the maximum strength of the terbinafine hydrochloride in the medicinal product does not exceed 1%;
 - (b) the medicinal product is sold or supplied in a container containing not more than 30 ml of the medicinal product; and
 - (c) the medicinal product is indicated for external use as a spray solution for the treatment of tina corporis, tinea pedis and tinea cruris.
- A medicinal product shall not be a prescription only medicine by reason of the fact that it contains triamcinolone acetonide where
 - (a) the medicinal product is in the form of a non-pressurised nasal spray;
 - (b) the medicinal product is indicated for the treatment of symptoms of seasonal allergic rhinitis in persons aged 18 years and over for a maximum period of 3 months;
 - (c) the container or package is labelled to show a maximum dose of 110 mcg per nostril and a maximum daily dose of 110 mcg per nostril; and
 - (d) the medicinal product is sold or supplied in a container or package containing not more than 3.375 mg of triamcinolone acetonide.

SCHEDULE 2

EXEMPTION FOR CERTAIN PERSONS FROM ARTICLE 57(2) OF THE LAW

PART 129

(Articles 1(2)(a) and 8(1))

Conditions 1.(1) The sale or supply
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1 (1) The sale or supply
1 (1) The sale or supply
1.(1) The sale of suppry
shall be subject to the
presentation of an order,
signed by the principal of
the institution or the
appropriate head of
department in charge of a
specified course of
research.
(2) The order shall
specify-
(a) the name of the
institution for which the
prescription only medicine
is required;
(b) the purpose for which
the prescription only
medicine is required; and
(c) the total quantity
required.
(3) The sale or supply shall
be only for the purposes of
the education or research
with which the institution
is concerned.
2.(1) The sale or supply shall be subject to the
presentation of an order
signed by or on behalf of
any person listed in any of
paragraph 2(a), (b) and (c)
of column 1 of this Part of
this Schedule.
(2) The order shall specify
the status of the person
1 2 4 6 5 6 1 6 1 6 1 6 1 6 1 6 1 6 1 6 1 6 1

Column 1	Column 2	Column 3
Persons exempted	Prescription only	Conditions
-	medicines to which the	
	exemption applies	
or her;		signing it, and the amount
		of the prescription only
		medicine required.
(b) an authorized officer		(3) The sale or supply shall
within the meaning of the		be only in connection with
Food Safety (Jersey)		the exercise by the person
Law 1966 ³¹ ; and		of his or her statutory
		functions.
(c) a person duly		
authorized by the Minister		
under Article 96 or 97 of		
the Law.		
3. Persons selling or	3. All prescription only	3.(1) The sale or supply
supplying prescription only	medicines.	shall be subject to the
medicines to any person		presentation of an order
employed or engaged in		signed by or on behalf of
connection with a scheme		the person so employed or
for testing the quality and		engaged.
checking the amount of		(2) The order shall specify
drugs and appliances		the status of the person
supplied under the Health		signing it, and the amount
Insurance (Jersey)		of the prescription only
Law 1967 ³² , or under any		medicine required.
subordinate legislation		
made under that law.		(2) The color or supply shall
		(3) The sale or supply shall be only for the purposes of
		a scheme to which
		paragraph 3 of column 1 of
		this Part of this Schedule
		refers.
4. Certified midwives.	4. Prescription only	4. The sale or supply shall
4. Certified illiawives.	medicines containing any	be only in the course of the
	of the following	midwife's professional
	substances –	practice and, in the case of
	Chloral hydrate	Ergometrine maleate, only
	Dichloral-phenazone	when contained in a
	Ergometrine maleate	medicinal product that is
	Pentazocine hydrochloride	not for parenteral
	Phytomenadrone	administration.
	Triclofos sodium	
5. Persons lawfully	5. Prescription only	5. The sale or supply shall
conducting retail pharmacy	medicines (not being for	be subject to the
businesses.	parenteral administration)	presentation of an order
	that are of any of the	signed by a registered
	following descriptions –	ophthalmic optician.
	(a) eye drops, or eye	
	ointments, that are	

Column 1	Column 2	Column 3
Persons exempted	Prescription only	Conditions
-	medicines to which the	
	exemption applies	
	prescription only medicines	
	by reason only that they	
	contain –	
	(i) 30.0% Sulphacetamide	
	sodium; or	
	(ii) 0.5% Chloramphenicol;	
	(b) eye ointments that are	
	prescription only medicines	
	by reason only that they	
	contain –	
	(i) 30.0% Sulpha-cetamide	
	sodium; or	
	(ii) 0.5%Chloramphenicol;	
	or	
	(c) medicinal products that	
	are prescription only	
	medicines by reason only	
	that they contain any of the	
	following substances –	
	Atropine sulphate	
	Bethanecol chloride	
	Carbachol	
	Cyclopentolate	
	hydrochloride	
	Homatropine	
	hydrobromide	
	Hyoscine hydrobromide	
	Naphazoline hydrochloride Naphazoline nitrate	
	Neostigmine methyl-	
	sulphate	
	Physostigmine salicylate	
	Physostigmine sulphate	
	Pilocarpine hydrochloride	
	Pilocarpine nitrate	
	Tropicamide.	
6. Registered ophthalmic	6. Prescription only	6.(1) The sale or supply
opticians.	medicines listed in	shall be only in the course
1	paragraph 5 of column 2 of	of the optician's
	this Part of this Schedule.	professional practice.
		(2) The sale shall be only
		in an emergency.
7.(1) Holders of product	7. Prescription only	7. The sale or supply shall

Column 1	Column 2	Column 3
Persons exempted	Prescription only medicines to which the exemption applies	Conditions
licences.	medicines to which the licence relates.	be only –
(2) Holders of manufacturer's licences.		(a) to a pharmacist, 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 (b) of no greater quantity than is reasonably necessary for that purpose.
8. Pharmacists selling or supplying to persons to whom cyanide salts may be sold lawfully under the Poisons (Jersey) Law 1952 ³³ .	8. Amyl nitrite.	8. The sale or supply shall only be so far as is necessary to enable an antidote to be available to persons at risk of cyanide poisoning.

PART 2

(Articles 1(2) and 8(1))

Column 1	Column 2	Column 3
Persons exempted	Prescription only	Conditions
	medicines to which the	
	exemption applies	
1.(1) The Royal National	1. All prescription only	1. The supply shall be
Lifeboat Institution.	medicines.	only so far as is
(2) Certificated first aiders of		necessary for the
the Institution.		treatment of sick or
		injured persons.
2. The owner or the master of a	2. All prescription only	2. The supply shall be
ship that does not carry a doctor	medicines.	only so far as is
on board as part of the ship's		necessary for the
complement.		treatment of persons on
		the ship.
3. The operator or commander	3. Prescription only	3.(1) The supply shall be
of an aircraft.	medicines that –	only so far as is
	(a) are not for parenteral	necessary for the
	administration; and	immediate treatment of
	(b) have been sold or	sick or injured persons
	supplied to the operator	on the aircraft.
	or commander of the	(2) The supply shall be in
	aircraft in response to an	accordance with the
	order in writing signed	written instructions of a
	by a doctor.	doctor as to the
		circumstances in which
		prescription only
		medicines of the
		description in question
		are to be used on the
4. Dansana anthanina dha	4 Dunganintian auto	aircraft.
4. Persons authorized by	4. Prescription only	4. The supply shall be
licences granted under Article 4	medicines (being controlled drugs) whose	subject to the conditions, in the circumstances and
of the Misuse of Drugs	O ,	
(General Provisions) (Jersey)	supply is authorized by the licence.	to the extent specified in the licence.
Order 1989 ³⁴ to supply a	uic ncence.	uic licelice.
controlled drug.		

Column 1	Column 2	Column 3
5. Persons requiring prescription only medicines to enable them, in the course of any business carried on by them, to comply with any requirements under any enactment in respect of the medical treatment of their employees. 6. Persons operating an occupational health scheme.	5. Prescription only medicines specified in the enactment. 6. Prescription only medicines sold or supplied to such a person in response to an order in writing signed by a doctor or a registered nurse.	5.(1) The supply shall be only to enable the person to comply with any such requirements. (2) The supply shall be subject to such conditions and in such circumstances as may be specified in the enactment. 6.(1) The supply shall be only in the course of the scheme. (2) The person supplying the prescription only medicine shall be – (a) a doctor; or
		(b) 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 scheme.

PART 335

(Articles 1(2) and 8(2))

Column 1	Column 2	Column 3
Persons exempted	Prescription only	Conditions
	medicines to which the	
	exemption applies	
1. Chiropodists, registered	1. Prescription only	1. The administration shall
under the Health Care	medicines for parenteral	be only in the course of the
(Registration) (Jersey)	administration that contain,	chiropodist's professional
Law 1995 ³⁶ , who hold	as the sole active	practice.
certificates of competence	ingredient, not more than	
in the use of analgesics	one of the following	
issued by or with the	substances –	
approval of the		
Chiropodists Board of the		
United Kingdom.		
	Bupivacaine hydrochloride	
	Bupivacaine hydrochloride	
	with adrenaline, where the	
	maximum strength of the	
	adrenaline does not exceed	
	1 mg in 200 ml of	
	bupivacaine hydrochloride	
	Lignocaine hydrochloride	
	Lignocaine hydrochloride	
	with adrenaline, where the	
	maximum strength of the	
	adrenaline does not exceed	
	1 mg in 200 ml of	
	lignocaine hydrochloride	
	Mepivacaine hydrochloride	
	Prilocaine hydrochloride	
2. Certified midwives.	2. Prescription only	2. The administration shall
	medicines for parenteral	be only in the course of the
	administration that contain	midwife's professional
	any of the following	practice and, in the case of
	substances (but no other	Lignocaine, Lignocaine
	substance specified in	hydrochloride and
	column 1 of Part I of the	Promazine hydrochloride,
	First Schedule to this	shall be only while
	Order) –	attending on a woman in childbirth.
	Ergometrine maleate	
	Levallorphan tartrate	
	Lignocaine	
	Lignocaine hydrochloride	
	Naloxone hydrochloride	

Column 1	Column 2	Column 3
	Oxytocins,	
	Natural and Synthetic	
	Pentazocine lactate	
	Pethidine	
	Pethidine hydrochloride	
	Phytomenadione	
	Promazine hydrochloride.	
3. The owner or the master	3. All prescription only	3. The administration shall
of a ship that does not	medicines that are for	be only so far as is
carry a doctor on board as	parenteral administration.	necessary for the treatment
part of the ship's		of persons on the ship.
complement.		
4. The operator or	4. Prescription only	4.(1) The administration
commander of an aircraft.	medicines for parenteral	shall be only so far as is
	administration that have	necessary for the
	been sold or supplied to the	immediate treatment of
	operator or commander of	sick or injured persons on
	the aircraft in response to	the aircraft.
	an order in writing signed by a doctor.	
	by a doctor.	(2) The administration
		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.
5. Persons operating an	5. Prescription only	5.(1) The administration
occupational health	medicines for parenteral	shall be only in the course
scheme.	administralion that have	of the scheme.
	been sold or supplied to	(2) The person
	such a person in response	administering the
	to an order in writing	prescription only medicine
	signed by a doctor or a	shall be –
	registered nurse	(a) a doctor;
		(b) a person acting in
		accordance with the
		directions of a doctor; or
		(c) 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
	<u> </u>	the scheme.

Column 1	Column 2	Column 3
6. Persons who hold	6. The following	6. The administration shall
certificates of proficiency	prescription only	be only for the immediate,
in ambulance paramedical	medicines for parenteral	necessary treatment of sick
skills issued by or with the	administration –	or injured persons and, in
approval of the Secretary		the case of a prescription
of State of the United		only medicine containing
Kingdom, or persons who		Heparin sodium, shall be
are state registered		only for the purpose of
paramedics.		cannula flushing.
	(a) Diazemuls (product	
	licence number	
	10183/00001);	
	(b) Gelofusine (product	
	licence number	
	00183/5025R); and	
	(c) medicines containing	
	the substances Ergometrine	
	Maleate 500mcg per ml	
	with Oxytocin 5 iu per ml,	
	but no other active	
	ingredient:	
	(d) prescription only	
	medicines that contain one	
	or more of the following	
	substances, (but no other	
	active ingredient)-	
	Adrenaline acid tartrate	
	Benzylpenicillin	
	Frusemide	
	Glucose	
	Heparin sodium	
	Lignocaine hydrochloride	
	Metoclopramide	
	Morphine Sulphate	
	Nalbuphine hydrochloride	
	Naloxone hydrochloride	
	Polygeline	
	Sodium bicarbonate	
	Sodium chloride.	
	Streptokinase	

SCHEDULE 3

(Articles 9(3)(c) and (4)(a))

SUBSTANCES THAT MUST NOT BE CONTAINED IN A PRESCRIPTION ONLY MEDICINE EXEMPTED BY ARTICLE 9

Ammonium Bromide Amylobarbitone Amylobarbitone Sodium Barbitone Butobarbitone Sodium Butobarbitone Sodium Calcium Bromide Calcium Bromide Calcium Bromide Cyclobarbitone Cyclobarbitone Cyclobarbitone Cyclobarbitone Cyclobarbitone Cyclobarbitone Cyclobarbitone Embutramide Fencamfamin Hydrochloride Fluanisone Heptabarbitone Hexobarbitone Hexobarbitone Hexobarbitone Sodium Hydrobromic Acid Meclofenoxate Hydrochloride Methohexitone Sodium Methylphenobarbitone Pemoline Pentobarbitone Pentobarbitone Phenobarbitone Phenobarbitone Sodium Phenylmethylbarbituric Acid Piracetam Potassium Bromide Prolintane Hydrochloride Quinalbarbitone Quinalbarbitone Sodium Quinidine Phenylethylbarbiturate Secbutobarbitone Sodium Quinidine Phenylethylbarbiturate			
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Secbutobarbitone Sodium			
Sodium Bromide			
Strychnine Hydrochloride			
Tacrine Hydrochloride			
Thiopentone Sodium			
Note (for information): The restriction in Article 9(3)(c) is subject to Article 9 in respect of Phenobarbitone and Phenobarbitone Sod for use in the treatment of epilepsy.			

SCHEDULE 437

(Articles 6(2)(ca) and 6A(2))

CLINICAL MANAGEMENT PLAN

1 Information to be included in clinical management plan

A clinical management plan must include –

- (a) the name of the patient to whom the plan relates;
- (b) the illness or conditions in relation to which the supplementary prescriber may give a prescription or administer (or direct the administration of) a medicinal product;
- (c) the date on which the plan is to take effect and the date or dates when it is subject to review by the doctor or dentist who is a party to the plan;
- (d) the class or description of medicinal product that may be prescribed by a supplementary prescriber or administered by, or under the direction of, a supplementary prescriber;
- (e) any restrictions or limitations as to the strength or dose, or period of use, of any medicinal product which may be prescribed by or administered by, or under the direction of, the supplementary prescriber;
- (f) any relevant warnings about the known sensitivities of the patient to, or known difficulties of the patient with, particular medicinal products;
- (g) arrangements for the notification of suspected or known adverse reactions to
 - (i) the medicinal product referred to in paragraph (d), and
 - (ii) any other medicinal product taken at the same time or over the same period;
- (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.

ENDNOTES

Table of Legislation History

Legislation	Year and No	Commencement
Medicines (Prescription Only)	R&O.9140	1 January 1998
(Jersey) Order 1997		
Medicines (Prescription Only)	R&O.9326	1 January 1999
(Amendment) (Jersey)		
Order 1998		
Medicines (Prescription Only)	R&O.1/2000	1 February 2000
(Amendment No. 2) (Jersey)		
Order 2000		
Medicines (Prescription Only)	R&O.2/2001	1 February 2001
(Amendment No. 3) (Jersey)		
Order 2001		
Medicines (Prescription Only)	R&O.94/2002	1 October 2002
(Amendment No. 4) (Jersey)		
Order 2002		
Medicines (Prescription Only)	R&O.75/2003	13 August 2003
(Amendment No. 5) (Jersey)		
Order 2003		
Medicines (Prescription Only)	R&O.65/2004	12 July 2004
(Amendment No. 6) (Jersey)		
Order 2004		
Medicines (Prescription Only)	R&O.174/2005	11 November 2005
(Amendment No. 7) (Jersey)		
Order 2005		
States of Jersey (Amendments	R&O.45/2005	9 December 2005
and Construction Provisions		
No. 5) (Jersey) Regulations 2005		
Pharmacists and Pharmacy	L.6/2010	16 May 2010
Technicians (Registration)		
(Jersey) Law 2010		
Medicines (Prescription Only)	R&O.82/2013	1 July 2013
(Amendment No. 8) (Jersey)		
Order 2013		

Table of Renumbered Provisions

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6(c)	6(d)
THIRD SCHEDULE	SCHEDULE 3

Table of Endnote References

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This Order has been amended by the States of Jersey (Amendments and
                      Construction Provisions No. 5) (Jersey) Regulations 2005. The
                      amendments replace all references to a Committee of the States of Jersey
                      with a reference to a Minister of the States of Jersey, and remove and add
                      defined terms appropriately, consequentially upon the move from a
                      committee system of government to a ministerial system of government
                      chapter 20.625
3
                      chapter 08.680
                      chapter 26.500
                      chapter 15.240
                      chapter 20.625
                      chapter 20.750
                      chapter 20.300.50
<sup>9</sup>Article 1(1)
                      amended by R&O.2/2001, R&O.82/2013
                      chapter 15.360
                      chapter 20.625
<sup>12</sup> Article 2
                      amended by R&O.82/2013
<sup>13</sup> Article 5
                      substituted by R&O.82/2013
                      chapter 20.300
<sup>15</sup> Article 6
                      heading amended by R&O.82/2013
                      chapter 08.680.60
<sup>17</sup> Article 6(2)
                      amended by R&O.82/2013
<sup>18</sup> Article 6A
                      inserted by R&O.82/2013
                      chapter 08.680.60
20
                      chapter 20.625.80
21
                      chapter 08.680.60
22
                      chapter 20.625.80
<sup>23</sup> Article 9(3)
                      amended by R&O.9326, R&O.1/2000
                      chapter 08.680.60
<sup>25</sup> Article 10
                      substituted by R&O.82/2013
<sup>26</sup> Article 11
                      inserted by R&O.65/2004
<sup>27</sup> Schedule 1
                      Part 1 amended by R&O.9326, R&O.1/2000, R&O.2/2001,
                      R&O.94/2002, R&O.174/2005
<sup>28</sup> Schedule 1
                      Part 4 substituted by R&O.1/2000, amended by R&O.2/2001,
                      R&O.94/2002,
<sup>29</sup> Schedule 2
                      Part 1 amended by R&O.1/2000, L.6/2010
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                      chapter 20.225
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                      chapter 20.225
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                      chapter 26.500
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                      chapter 20.775
34
                      chapter 08.680.60
<sup>35</sup> Schedule 2
                      Part 3 amended by R&O.9326, R&O.1/2000, R&O.2/2001
                      chapter 20.300
<sup>37</sup> Schedule 4
                      inserted by R&O.82/2013
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