BERMUDA

PHARMACY AND POISONS ACT 1979

1979 : 26

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PART I
PRELIMINARY

Short title
1 This Act may be cited as the Pharmacy and Poisons Act 1979.
Interpretation

In this Act, unless the context otherwise requires—

“Association” means the professional association representing pharmacists in Bermuda;

“certificate of competence” means a certificate of competence granted by the Council under regulations made under section 15(1)(b);

“the Council” means the Pharmacy Council established by section 3;

“dentist” means a dental practitioner registered under the Dental Practitioners Act 1950 [title 30 item 4] or an exempted dental practitioner within the meaning of that Act;

“dispense,” with its grammatical variations, in relation to a medicine or a poison, means the preparation and supplying in such manner of a medicine or a poison on and in accordance with a prescription given by a duly qualified practitioner as to ensure the pharmaceutical and therapeutic suitability to the circumstances for which it is prescribed;

“drug” means a substance or combination of substances used, or for use in or on the body of a person or animal—

(a) to prevent, diagnose, treat or mitigate a disease, disorder or abnormal physical or mental state or symptom of them; or

(b) to restore, correct or modify organic functions, and includes a prescribed substance or combination of substances;

“drug product” means a manufactured product that contains a drug including tablets, pills, capsules, caplets, creams, powders, transdermal patches or liquids;

“functions” includes powers and duties;

“medicinal use” means—

(a) use by being administered to one or more human beings or animals; or

(b) use as an ingredient in the preparation of a substance or article which is to be administered to one or more human beings or animals;

for—

(i) treating or preventing disease;

(ii) diagnosing disease or ascertaining the existence, degree or extent of a physiological condition;

(iii) contraception;

(iv) inducing anaesthesia; or

(v) otherwise preventing or interfering with the normal operation of a physiological function, whether permanently or temporarily, and
whether by way of terminating, reducing or postponing, or increasing or accelerating, the operation of that function or in any other way;

“Minister” means the Minister responsible for Health;

“non-practising pharmacist” means a person who is registered as a pharmacist section 7A but who is not practising pharmacy in Bermuda;

“physician” means a medical practitioner registered under the Medical Practitioners Act 1950 [title 30 item 8] or an exempted medical practitioner within the meaning of that Act;

“poison” has the meaning assigned thereto in section 33;

“practitioner” includes any of the professions listed in the Second Schedule;

“prescribed” means prescribed by regulations;

“prescription” means a prescription issued by any of the practitioners listed in the Second Schedule;

“registered pharmacist” means a person registered pursuant to section 7(4) and (4B);

“registered pharmacy” has the meaning assigned thereto in section 17(3);

“Registrar” means the official for whose appointment section 7(1) provides;

“regulation” means regulation made under section 15, 22 or 48;

“relevant professional body”, in relation to registered pharmacists, means the Bermuda Pharmaceutical Association;

“Schedule 3 drug” has the meaning assigned thereto in section 25(6);

“Schedule 4 drug” has the meaning assigned thereto in section 28(1);

“veterinary practitioner” means a person who holds a certificate issued under the Veterinary Practitioners Act 2008.

[Section 2, “veterinary practitioner” amended by 2008 : 20 s.17 & Sch. 2 effective 9 July 2010; Section 2, “Schedule 3 drug” amended by 2011 : 31 s. 2 effective 10 August 2011; Section 2 “Association”, “drug”, “drug product”, “non-practising pharmacist” inserted, “practitioner” and “a prescription” deleted and substituted and “registered pharmacist” amended by 2013 : 48 s. 2 effective 24 December 2013]

PART II
THE PHARMACY COUNCIL

The Pharmacy Council
3 There shall be established a body called the Pharmacy Council, whose general function shall be to secure high standards of professional competence and conduct in the practice of pharmacy in Bermuda, and who shall have such other functions as may be assigned to the Council by any statutory provision.
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Membership of the Council

4 (1) The Council shall consist of—
   (a) a chairman appointed by the Minister;
   (aa) one member, who shall be a representative of the Association, appointed by the Minister after consulting with the Chief Medical Officer;
   (b) one member, who shall be a practitioner, appointed by the Minister after consulting with the Chief Medical Officer; and
   (c) four members elected by registered pharmacists from among themselves.

(2) The Council may co-opt a representative of the Bermuda Pharmacy Owners Association to any of their meetings but such a representative shall not have a vote.

[Section 4 subsection (1)(aa) inserted and subsection (1)(c) amended by 2013 : 48 s. 3 effective 24 December 2013]

Functions of the Council

4A The Council shall, in addition to any other function under this Act, make periodic reviews of the Act for the purpose of making recommendations to the Minister as to any necessary amendments to the Act generally, and with particular reference to the Third and Fourth Schedules.

Protection from personal liability

4B A member of the Council shall not be personally liable for damages for anything done or omitted to be done in the discharge or purported discharge of the Council’s functions under this Act unless the act or omission was in bad faith.

[Section 4B inserted by 2013 : 48 s. 4 effective 24 December 2013]

Proceedings of the Council, etc

5 The provisions in the First Schedule shall have effect with respect to the Council.

PART III

REGISTRATION OF PHARMACISTS

Offence to practise pharmacy if not registered

6 (1) It shall be unlawful for an individual to practise pharmacy unless at the time—
   (a) he is a registered pharmacist; and
   (b) he operates, or is employed at, premises which are a registered pharmacy.

(2) A person shall be deemed to be practising pharmacy for the purposes of this Act if, in the way of trade or business in Bermuda, he takes or uses a title, or holds himself out as engaging in a profession, being a title or profession to which this subsection applies.
The titles and professions to which subsection (2) applies are those of pharmacist, chemist, druggist, chemist and druggist, dispensing chemist and dispensing druggist, and any other suggesting a connexion with the business of compounding or dispensing medicines.

Registration as a pharmacist

(1) There shall be a Registrar, who shall be appointed by the Minister.

(2) The Registrar shall establish and maintain a register of pharmacists for the purposes of this Act.

(3) The register of pharmacists shall be kept at the offices of the Registrar, and be available for inspection by the public at all reasonable times without charge.

(4) Where a person who is qualified for registration as a pharmacist under this Act applies in the required form to the Registrar and pays the appropriate fee, the Registrar shall register him as a pharmacist under this Act by causing his name and the prescribed particulars relating to him to be entered in the register of pharmacists and giving him a certificate of registration in the prescribed form.

(4A) A person who has been registered under subsection (4) shall apply for re-registration every two years after the first registration.

(4B) The Registrar may approve an application for re-registration under subsection (4A) and issue a certificate of re-registration to the person applying.

(5) Any individual other than a disqualified person shall, for the purposes of subsection (4), be qualified for registration as a pharmacist under this Act if he—

(a) is fit and proper and possesses the appropriate qualifications and experience; and

(b) possesses a certificate of competence granted to him by the Council for passing a written exam in pharmacy set by the Council; and

(c) has had a minimum of six months’ practical experience of which not less than one month after graduation has been spent under the supervision in Bermuda of a registered pharmacist.

(5A) A person applying for re-registration under subsection (4A) shall—

(a) apply in the form required by the Council;

(b) pay the appropriate fee;

(c) continue to meet the qualifications, experience and conduct as required in subsection (5);

(d) meet the minimum amount of continuing professional development as required by the Council; and

(e) meet the number of practice hours as required by the Council.
(6) Where the Registrar refuses or fails to register a person who makes an application under subsection (4), or refuses or fails to re-register a person who makes an application under subsection (4A) (hereinafter in this section called “applicant”), the applicant may appeal to the Supreme Court.

(7) An applicant may appeal to the Supreme Court under subsection (6) within 28 days after the decision is made (in this section referred to as “the appeal period”).

(8) “appropriate fee” in subsections (4) and (5A) means the relevant fee prescribed in the Government Fees Regulations 1976.

(9) A list of registered pharmacists shall be published annually in the Gazette so soon as may be after the 1st day of January, and particulars of any alteration made in the register of pharmacists on or after that date in any year shall also be published in the Gazette.

[Section 7 subsections (1),(3),(4) and (5) amended, subsections (6) and (8) deleted and substituted, subsection (7) deleted and inserted and subsections (4B),(5A) and (7A) inserted by 2013 : 48 s. 5 effective 24 December 2013]

Re-registration as non-practising member

7A (1) A person who is registered under section 7(4) and is not practising pharmacy in Bermuda may re-register as a non-practising pharmacist in a form required by the Council and by paying the appropriate fee.

(2) The Registrar shall establish and maintain a register of non-practising pharmacists for the purposes of this Act.

(3) A person registered as a non-practising pharmacist shall not practise pharmacy in Bermuda.

(4) A non-practising pharmacist applying for re-registration to practise pharmacy shall—

(a) apply in the form required by the Council;
(b) pay the appropriate fee;
(c) continue to meet the qualifications, experience and conduct requirements in section 7(5); and
(d) meet the minimum amount and type of continuing professional development and practice hours as required by the Council.

(5) In subsection (1) “the appropriate fee” means the relevant fee prescribed in the Government Fees Regulations 1976.

[Section 7A inserted by 2013 : 48 s. 6 effective 24 December 2013]
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Code of Conduct
8  (1) It shall be the duty of the Council to prepare, and from time to time as they think fit amend, a code of conduct which the Council considers to be conduct that is proper for registered pharmacists in a professional respect (hereinafter in this Act called “the Code”).

(2) The Council shall send by post to each registered pharmacist at his address on the register of pharmacists a copy of the Code and of any amendment made to the Code.

(3) In exercise of their powers under section 10A the Council shall, subject to subsection (4), be guided by any relevant provision of the Code.

(4) Where an inquiry has been conducted by the Council under sections 10A or 10B, the Council may find a person guilty of negligence, incompetence or other improper conduct notwithstanding that the conduct in question is not prohibited by the Code, but they shall not find a person guilty of improper conduct if that conduct is authorized by the Code.

[Section 8 repealed and replaced by 2013 : 48 s. 7 effective 24 December 2013]

Pharmacy Profession Complaints Committee
9  (1) There shall be established, in accordance with the Fifth Schedule, a committee to be known as the “Pharmacy Profession Complaints Committee” (hereinafter in this Act called “the Committee”).

(2) The functions of the Committee are—

(a) to receive and investigate, or cause to be investigated, complaints against any registered pharmacist including any allegation that—

(i) the pharmacist’s registration was improperly obtained;

(ii) the pharmacist is guilty of professional misconduct;

(iii) the pharmacist is unfit to practise by reason of conviction of an indictable offence or adverse physical or mental health; or

(iv) the pharmacist is otherwise unfit to practise or to be registered; and

(b) to perform such other functions as may be prescribed.

(3) The Committee may investigate any complaint based on matters alleged to have occurred—

(a) inside or outside Bermuda; or

(b) at any time, whether or not at a time when the person complained against was registered as a pharmacist.

(4) A complaint referred to in subsection (2)(a)—

(a) shall be made by the complainant—
(i) if the complainant is a child or is physically or mentally unable to make
the complaint, by the parent or guardian, friend or a person acting on
behalf of the complainant; and

(ii) if the conduct complained of relates to a person who is dead, by the
person’s executor or personal representative;

(b) shall be in writing and addressed to the Committee;

(c) shall set out the matters alleged to constitute grounds for disciplinary
action to be taken against the pharmacist who is the subject of the
complaint;

(d) may be required by the Committee to be in a form approved by the
Committee.

(5) If the Committee considers that a complaint arose from a misunderstanding by
the complainant or between the complainant and the pharmacist complained of, the
Committee may, before proceeding further with the investigation of the complaint, require
the parties to appear before it in order to discuss the matter with a view to clarifying the
misunderstanding and resolving the matter informally.

(6) The Fifth Schedule has effect as to the appointment and proceedings of the
Committee and other matters relating to the Committee.

[Section 9 repealed and replaced by 2013 : 48 s. 7 effective 24 December 2013]

Investigation of complaint by Committee

10 (1) Where a complaint under section 9(4) is not resolved informally as provided in
section 9(5), the Committee shall investigate the complaint and determine whether, in its
opinion, the complaint—

(a) is frivolous or vexatious, is made in bad faith, is an abuse of process, or for
any other reason, ought not to be referred to the Council; or

(b) ought to be referred to the Council for decision.

(2) The Committee—

(a) shall give written notice to the pharmacist who is the subject of the
complaint that a complaint has been made, together with a summary of
the matters alleged in the complaint;

(b) shall request that the pharmacist who is the subject of the complaint show
cause in writing, within a specified time after the notice is given, explaining
why the matter should not be placed before the Council for determination; and

(c) may take evidence from witnesses on oath or affirmation, administered by
the Chairman.

(3) If the Committee determines that a complaint is frivolous or vexatious, is made
in bad faith, is an abuse of process or otherwise ought not to be considered by the
Committee, it shall dismiss the complaint and give written notice to the complainant of the dismissal and the reasons for the dismissal.

(4) If the Committee determines that a complaint ought to be referred to the Council for decision, the Committee shall, as soon as practicable, refer the matter to the Council.

Inquiry into complaint by Council

10A (1) If, pursuant to an investigation under section 10, the Committee places the matter before the Council for determination, the Council shall inquire into the matter.

(2) For the purposes of an inquiry of this section, the Council—

(a) may take evidence from witnesses on oath or affirmation, and for that purpose the Chairman of the Council may administer an oath or affirmation;

(b) shall afford the registered pharmacist and the Committee, or a member of the Committee, every facility—

(i) to appear before the Council;

(ii) to be represented by a barrister and attorney;

(iii) to call or cross-examine witnesses; and

(iv) generally to make a full defence or explanation in the matter of the complaint.

(3) Following its inquiry, the Council shall make a decision as to whether the complaint is proved or not proved, in whole or in part, together with reasons for its decision.

(4) If the Council decides that a complaint is not proved, in whole or in part, it shall dismiss the complaint to the extent that it is not proved.

(5) If the Council decides that a complaint is proved, in whole or in part, it shall record a finding to that effect and it may make any order of a disciplinary nature that it sees fit in respect of a pharmacist against whom the complaint is made, including an order—

(a) admonishing the pharmacist;

(b) suspending the pharmacist from practice as a pharmacist for such period as it sees fit or for an indefinite period;

(c) striking the name of the pharmacist off the register;

(d) imposing conditions or limitations with regard to the pharmacist’s practice as a pharmacist.

(6) The Council shall give written notice, to the pharmacist against whom a complaint is made, of its decision under subsection (3) and any order made by the Council under subsection (5), together with reasons.
(7) The pharmacist against whom the complaint is made may appeal against a decision or order of the Council in the manner provided in section 14.

(8) Any proceedings in connection with the holding of an inquiry by the Council under this section shall, for the purpose of the provisions of the Criminal Code Act 1907 relating to perjury, be deemed to be judicial proceedings.

(9) A member of the Council who was involved in the matter complained of may not participate in an inquiry by the Council under this section.

(10) A person who is suspended from practice under this section shall, for the duration of the suspension, be deemed not to be registered.

[Section 10A inserted by 2013: 48 s. 8 effective 24 December 2013]

**Inquiry by Council of its own initiative**

10B (1) In the absence of a complaint, the Council may, of its own initiative, hold an inquiry into any matter referred to in section 9(2) that could have formed the subject of an investigation by the Committee.

(2) The provisions of section 10A that apply in respect of an inquiry by the Council under that section shall apply to an inquiry under this section with any necessary modification.

[Section 10B inserted by 2013: 48 s. 8 effective 24 December 2013]

**Surrender of registration**

11 The Minister may order the Registrar to erase from the register of pharmacists the name of a registered pharmacist against whom no matter of complaint is pending under sections 10A and 10B, if the registered pharmacist applies to the Minister for the purpose and surrenders to him his certificate of registration.

[Section 11 amended by 2013: 48 s. 9 effective 24 December 2013]

**Restoration of name to register**

12 (1) A person whose name has been removed from the register under section 11, or whose name has been struck from the register, or who has been suspended from practice under section 10A or 10B, may make an application to the Council, in a form determined by the Council, for his name to be restored to the register or for his suspension to be terminated.

(2) An application under subsection (1) for the restoration of a name to the register of pharmacists shall not be made to, or be considered by, the Council—

(a) within twelve months after the date of removal, striking off or suspension;

(b) within twelve months after a previous application under that subsection; or

(c) where the Council in the direction ordering the erasure appointed a period within which another application should not be made under that subsection, within that period.
(3) On receipt of an application, the Council shall decide whether to restore the applicant's name to the register or to terminate his suspension, after considering the following matters—

(a) the character and professional ability of the applicant;
(b) the nature of the matter in respect of which the applicant’s name was struck from the register or for which the applicant was suspended;
(c) the conduct of the applicant after his name was struck from the register or after he was suspended;
(d) any other circumstances appearing to the Council to be relevant.

(4) The Council shall give written notice to the applicant of its decision, together with reasons.

(5) An applicant may appeal against the decision of the Council in the manner provided in section 14.

[Section 12 repealed and replaced by 2013 : 48 s. 10 effective 24 December 2013]

Proof of registration

A certificate signed by the Permanent Secretary to the Minister certifying that a person named in the certificate is or, as the case may be, is not, a registered pharmacist and, in the case of a person to whom the certificate refers as being a registered pharmacist, specifying the date of registration, shall be admissible in any proceedings as prima facie evidence of the facts stated in the certificate.

Appeals

(1) A person aggrieved by any decision of the Council referred to in subsection (2) may, within 28 days after the date on which the decision is given to the person by the Council, appeal to the Supreme Court against the decision.

(2) The person referred to in subsection (1) may appeal against the following decisions—

(a) a decision not to issue or renew a registration certificate;
(b) a decision not to issue a re-registration certificate;
(c) a decision to remove the name of a person from the register;
(d) a decision to remove, or alter, any entry in the register in respect of a person;
(e) a decision not to restore a person’s name to the register;
(f) a decision not to terminate a person’s suspension.

(3) On an appeal under this section the Supreme Court may make such order in the matter as it thinks proper, including an order as to the costs of the appeal.

(4) An order of the Supreme Court under subsection (2) is final.
(5) The practice and procedure to be followed in relation to an appeal under this section are as prescribed by rules of court.

(6) The Council may appear as respondent on such appeal and, whether they appear at the hearing of the appeal or not, they shall be deemed to be a part to the appeal for the purpose of enabling directions to be given as to the costs or expenses of the appeal.

[Section 14 repealed and replaced by 2013 : 48 s. 11 effective 24 December 2013]

Regulations for this part

15 (1) The Minister may make regulations—

(a) regulating the making of applications for registration as a pharmacist under this Act and providing for the evidence to be produced in support of such applications;

(b) prescribing professional standards that are to be met by registered pharmacists;

(c) prescribing the procedure to be followed on an inquiry held pursuant to sections 10A and 10B.

(2) Regulations made under subsection (1) shall be subject to the negative resolution procedure.

[Section 15 amended by 2013 : 48 s. 9 effective 24 December 2013]

PART IV

REGISTRATION OF PHARMACIES

Register of pharmacies

16 (1) The Registrar shall establish and maintain a register of pharmacies for the purposes of this Act.

(2) The register of pharmacies shall be kept at the offices of the Registrar, and be available for inspection by the public at all reasonable times without charge.

[Section 16 subsection (2) amended by 2013 : 48 s. 12 effective 24 December 2013]

Registration of premises as registered pharmacies

17 (1) Where an application for the registration of premises as a registered pharmacy is made by any person (hereafter in this Part called an “applicant”) to the Registrar on the prescribed form accompanied by the appropriate fee, the Registrar shall, subject to sections 18, 20 and 21(1), enter the prescribed particulars relating to those premises in the register of pharmacies.

(2) In subsection (1) “the appropriate fee” means the relevant fee prescribed in the Government Fees Regulations 1976.
In this Act “to register premises as a registered pharmacy” means to enter the prescribed particulars relating to them in the register of pharmacies pursuant to subsection (1), and any premises in relation to which the prescribed particulars are so entered are in this Act referred to as a “registered pharmacy”.

A list of registered pharmacies shall be published annually in the Gazette so soon as may be after the 1st day of January, and particulars of any alteration made in the register of pharmacies on or after that date in any year shall also be published in the Gazette.

It shall be an offence for any premises to bear any sign or other representation that it is a pharmacy, drug-store, dispensary or other words representing any such premises as being registered as a pharmacy under this Act unless such premises are in fact so registered, or for any person to represent himself as being a pharmacist, apothecary, druggist, dispenser or any other description, whether of the foregoing classes or not, calculated to represent that he is registered as a pharmacist under this Act, unless he is so registered.

Unfit premises: new applications

If it appears to the Minister that premises in respect of which an application under section 17 has been made fail in a material respect to comply with the requirements of regulations made under section 22(1)(a) which are for the time being in force, he may determine to issue to the applicant a certificate of unfitness under this section certifying that the premises are unsuitable for registration as a registered pharmacy.

Before the Minister issues a certificate of unfitness under this section, he shall serve on the applicant a notice stating what he proposes and his reasons therefor.

If within fourteen days after receipt of a notice under subsection (2) the applicant makes representations in writing to the Minister, or gives notice in writing to the Minister of his desire to be heard with respect to the Minister’s proposal to issue such a certificate, the Minister shall not issue the certificate before he has considered the applicant’s representations in writing or, where the applicant gave notice of his desire to be heard, his oral representations if made within a reasonable time.

Where the Minister, after considering any such representations as aforesaid, determines not to issue a certificate of unfitness under this section in respect of the premises in question, he shall notify the applicant and the Registrar of his decision, and the Registrar shall forthwith register the premises as a registered pharmacy.

Where the Minister, after considering any such representations as aforesaid, determines that a certificate of unfitness ought to be issued in respect of the premises in question, he shall issue the certificate by serving it on the applicant, and he shall also serve a copy of the certificate on the Registrar.

A certificate of unfitness issued under this section shall state the reasons for its issue.
(7) Except in accordance with the directions of the Supreme Court given under section 20(2), the Registrar shall not register as a registered pharmacy premises in respect of which a certificate of unfitness has been issued under this section.

Unfit premises: registered pharmacies

19 (1) Where the Minister is of opinion that a registered pharmacy fails in a material respect to comply with the requirements of regulations made under section 22(1)(a) which are for the time being in force, the Minister shall serve on the operator of the pharmacy a notice stating his intention to issue a certificate of unfitness under this section in respect of the pharmacy, and the Minister’s reasons therefor; and section 18(3) to (6) shall have effect mutatis mutandis in relation to notices and certificates under this section as they have effect in relation to notices and certificates of unfitness under that section.

(2) Where a certificate of unfitness is issued under this section, the registered pharmacy to which the certificate relates shall cease to be a registered pharmacy with effect from the date of the taking effect of the certificate under section 21.

Appeals

20 (1) Any person aggrieved by the issue of a certificate of unfitness under section 18 or 19 may, at any time within twenty-eight days after the service of the certificate upon him, appeal under this section to the Supreme Court, and upon any such appeal that Court shall have power to confirm or revoke the issue of the certificate.

(2) Where the Supreme Court revokes a certificate of unfitness issued under section 18, the Court shall give such directions as the case requires with regard to the registration of the premises as a registered pharmacy under section 17.

(3) The powers of the Chief Justice to make rules under section 62 of the Supreme Court Act 1905 [title 8 item 1] shall extend to the making of rules regulating the practice and procedure to be followed on, and the fees to be paid in connexion with, any such appeal.

(4) The Registrar may appear as the respondent on any such appeal and, whether he appears at the hearing of the appeal or not, he shall be deemed to be a party to the appeal for the purpose of enabling directions to be given as to the costs or expenses of the appeal.

When certificates of unfitness take effect

21 (1) Without prejudice to section 18(7), where an appeal is not brought against the issue of a certificate of unfitness under that section, or where such an appeal is brought but is withdrawn or struck out for want of prosecution, the certificate shall take effect on the expiration of the time for appealing or, as the case may be, on the withdrawal or striking out of the appeal; but otherwise such a certificate shall take effect if and when the appeal is dismissed and not otherwise.

(2) Where an appeal is not brought against the issue of a certificate of unfitness under section 19, or where such an appeal is brought but is withdrawn or struck out for want of prosecution, the certificate shall take effect on the expiration of thirty days after the expiration of the time for appealing or, as the case may be, upon the expiration of thirty days after the withdrawal or striking out of the appeal; but otherwise such a certificate shall
take effect upon the expiration of thirty days after the dismissal of the appeal and not otherwise.

**Regulations for this Part**

22 (1) The Minister may make regulations under this section with respect to registered pharmacies—

(a) prescribing standards for their maintenance and operation, including provision for space, equipment and facilities;

(b) imposing requirements as to the circumstances in which a registered pharmacist must, or (as the case may require) need not, be present in a registered pharmacy;

(c) prescribing the books and records to be kept, and providing for the examination by or on behalf of the Minister of such books and records;

(d) prescribing the returns to be made, and information to be forwarded, to the Minister.

(2) Regulations made under this section shall be subject to the negative resolution procedure.

**PART V**

**CONTROL OF PRESCRIPTIONS AND IMPORTATION**

**Prescriptions to be in a certain form**

23 (1) Subject to the provisions of this section, a prescription of any substance shall not be made by a practitioner unless it is on a valid prescription form which includes the information as provided in regulation 3A of the Pharmacy and Poisons (Control of Prescriptions) Regulations 1979.

(2) Nothing in subsection (1) shall make it unlawful for a registered pharmacist to execute a prescription that is transmitted to him by telephone by a practitioner where the practitioner's voice is known to him and he honestly believes the voice of the person transmitting the prescription to be that of the practitioner.

(3) Subsection (1) shall not apply to a practitioner who transmits a prescription to a registered pharmacist by telephone if the prescription is for a ten-day supply of the medicine prescribed; so, however, that in no case such a prescription be refilled by the registered pharmacist.

(4) The original of every prescription dispensed by him shall bear a number and shall be preserved by the registered pharmacist on a file kept for that purpose in the pharmacy and he shall, where requested to do so by another registered pharmacist, furnish a copy thereof to that other registered pharmacist unless the prescribing practitioner has forbidden the furnishing of such a copy.
(5) A copy of a prescription furnished to another registered pharmacist shall contain the following information:

(a) the name and address of the prescribing practitioner and of the person for whom the substance has been prescribed;
(b) the name of the substance prescribed, its strength and quantity, and directions for its use;
(c) the dates of the first and last dispensing of the substance prescribed and the number of refills (if any) remaining; and
(d) the number of the prescription and the name and address of the pharmacy.

(6) Where a request is made for a prescription to be refilled at a pharmacy other than at which the substance prescribed was first dispensed, the registered pharmacist to whom the request is made shall communicate with the pharmacy at which the substance was first dispensed for the purpose of obtaining a copy of the prescription and the pharmacy at which the substance prescribed was first dispensed shall make a record of the date, the name and address of the pharmacy where the prescription is refilled. In the event that a third pharmacy is in possession of the original prescription, that pharmacy must be informed as well of the fact of the refilling of the prescription and of the date, name and address of the pharmacy where the prescription is refilled. A registered pharmacist who refills a prescription shall make a record of the date and quantity of the substance dispensed and he shall initial the record.

(7) A registered pharmacist may, at the request of a person under medical treatment and where the circumstances constitute an emergency, supply a Schedule 3 drug in relation that person without a prescription being presented to him:

Provided that in no circumstances whatever shall he supply a drug which is also specified in Schedule 2 of the Misuse of Drugs Regulations 1973.

(8) Before a registered pharmacist may supply a Schedule 3 drug under subsection (7) he must satisfy himself by means of questions put to the person requesting the drug—

(a) that there is a genuine and urgent need by the person for the Schedule 3 drug;
(b) that it is not practicable in the circumstances of the particular case for a prescription to be obtained from a practitioner immediately;
(c) that treatment with the particular Schedule 3 drug has been previously prescribed by a practitioner for the person requesting it; and
(d) that the dose which he will supply is appropriate to the need of the person.

(9) The supply of a Schedule 3 drug in the circumstances specified in subsection (8) shall not in any case exceed five days’ supply except—

(a) where the drug is in the form of an ointment or cream, or is a preparation in an aerosol container for the relief of asthma, and in these cases the supply shall consist of the smallest package or container available:
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(b) an oral contraceptive in which case the full cycle may be dispensed; or

c) an antibiotic in liquid form for oral administration, in which case the
smallest quantity that will provide a full course of treatment may be
supplied.

(10) The container or package of a Schedule 3 drug supplied pursuant to subsection
(7) shall bear a label showing—

(a) an identification number;
(b) the date of supply;
(c) the name of the person to whom supplied
(d) the name and address of the supplying pharmacy;
(e) the name, quantity, directions for use, and where appropriate, the
pharmaceutical form and strength of the drug;
(f) the words EMERGENCY SUPPLY marked thereon; and
(g) the initials of the registered pharmacist.

(11) The registered pharmacist shall also keep a book entitled “Emergency Supply
Book” in which shall be entered the particulars at subsection (10)(a) to (f) (inclusive).

[Section 23 subsection (1) repealed and replaced and subsections (3)-(7) (10) and (11) amended by 2013 : 48 s. 14 effective 24 December 2013]

Validity of a prescription
23A A prescription shall be valid for one year from the date as shown on a valid
prescription form.

[Section 23A inserted by 2013 : 48 s. 15 effective 24 December 2013]

Supply by registered pharmacist of equivalent medicines
24 (1) Where a registered pharmacist receives for execution a prescription which does
not prohibit an alternative equivalent drug or drug product from being supplied under the
prescription—

(a) it shall be required for the registered pharmacist to supply under the
prescription any drug or drug product available to the pharmacist at the
location of sale—

(i) which is in his opinion the chemical and therapeutic equivalent of the
drug or drug product specified in the prescription; and

(ii) if taking all relevant factors into account, the price that he charges and
accepts for the drug or drug product he supplies is less than that which
he would have charged and accepted for the drug or drug product
specified;
(b) it shall be lawful for the registered pharmacist to supply under the prescription any drug or drug product—

(i) which is in his opinion the chemical and therapeutic equivalent of the drug or drug product specified in the prescription; and

(ii) if taking all relevant factors into account, the prices that he charges and accepts for the drug or drug product he supplies is the same as that which would have charged and accepted for the drug or drug product specified.

(2) A drug or drug product supplied by a registered pharmacist under subsection (1) must be a drug or drug product accepted by the Council as the chemical and therapeutic equivalent of the drug or drug product specified in the prescription in question.

[Section 24 repealed and replaced by 2013 : 48 s. 16 effective 1 February 2014]

Restrictions on the importation of medicines

25 (1) A person shall only import into Bermuda for medicinal use medicines that are obtained from foreign manufacturers or foreign wholesalers if those medicines are eligible for sale in the United States of America, Canada or a country in the European Union in accordance with the regulatory standards of the relevant country.

(2) A person who acquires medicine from abroad for distribution or sale in Bermuda shall register with the Minister in accordance with regulations made under this Act by the Minister.

(3) Any person who fails to comply with this section or any regulations made under this Act commits an offence.

(4) A person who fails to comply with this section or any regulations made under this Act may have any medicines being imported by him forfeited to the Crown.

(5) The Minister may make regulations to prescribe the requirements for—

(a) the registration of a person under subsection (2); and

(b) the importation of medicines.

(6) In this section—

"manufacturer" means a person involved in the production, preparation, propagation, conversion, processing, packaging or labelling of medicine;

"medicine" means any substance specified in the Third Schedule (in this Act referred to as a "Schedule 3 drug");

"wholesaler" means a person who obtains medicine for distribution or delivery to persons other than consumers.

(7) The negative resolution procedure shall apply to regulations made under this section.

[Section 25 repealed and replaced by 2011 : 31 s. 3 effective 10 August 2011]
Declaration relating to imported medicines
26  [Repealed by 2011 : 31 s. 4]
[Section 26 repealed by 2011 : 31 s. 4 effective 10 August 2011]

PART VI
CONTROL OF DRUGS

Certain substances to be sold on prescription only
27  (1) Subject to any provision made by any regulation, no person shall for medicinal use sell any Schedule 3 drug otherwise than under a prescription.

(2) In this section and section 28 “sell” or “sale” means sell or sale by retail.

Certain substances to be available at pharmacies only
28  (1) Subject to subsection (3) and to any provision made by any regulation, no person shall for medicinal use keep for sale, or sell, any substance specified in Part I or Part II of the Fourth Schedule (in this Act referred to as a “Schedule 4 drug”) elsewhere than at a registered pharmacy.

(2) Subject to subsection (3) and to any provision made by any regulation, no person shall for medicinal use keep for sale, or sell, any substance specified in Part II of the Fourth Schedule unless he is a registered pharmacist.

(3) Subsection (1) or (2) shall not apply to a practitioner as respects anything done by him in the course of his practice as such.
[Section 28 substituted by 1989:56 effective 15 January 1990]

Restrictions on dispensing
29  Subject to any provision made by any regulation no person other than a registered pharmacist or a practitioner acting in the course of his practice as such shall manufacture or compound or dispense any Schedule 3 or Schedule 4 drug.

Prohibition on giving away Schedule 3 or Schedule 4 drugs
30  (1) Subject to subsection (2), no person shall make a gift of any Schedule 3 or Schedule 4 drug to any person who is not a practitioner or a registered pharmacist.

(2) Subsection (1) shall not apply—

(a) to a practitioner who makes a gift of a Schedule 3 or Schedule 4 drug; or

(b) to a registered pharmacist who makes a gift of a Schedule 4 drug,
to another person for use by that person for the medical or dental treatment of a human being or animal.
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Unfit drugs
31  [Repealed by 2013 : 48 s. 17]

[Section 31 repealed by 2013 : 48 s. 17 effective 24 December 2013]

Health and safety requirements
31A  (1) For the purposes of sections 31B, 31C, 31D and 31E, a drug or drug product fails to comply with a general health and safety requirement if it is not reasonably safe having regard to all the circumstances including—

(a) the manufacturer of a drug or drug product, or any regulatory authority that granted a drug or drug product marketing authorisation, issuing a recall or any form of notice of warning for the drug or drug product;

(b) marketing authorisation of a drug or drug product granted by the regulatory authority in the United States, Canada or the European Union, or another jurisdiction that the United States, Canada or the European Union has a mutual recognition agreement with, is denied, suspended or discontinued due to reasons of quality, safety or efficacy;

(c) the storage, distribution, supply, security, or handling of the product compromised its safety, quality or efficacy due to standards set by the manufacturer or regulatory authority that granted marketing authorization for the drug or drug product;

(d) the drug or drug product is not properly labelled to allow for—

(i) its safe consumption;

(ii) the determination of—

(A) the amount of active ingredients;

(B) its proper use;

(C) the content;

(e) any other risk to public or individual health as specified by the Minister after consultation with the Chief Medical Officer.

(2) A person is guilty of an offence under this section if he—

(a) supplies any drug or drug product which fails to comply with the health and safety requirement or any prescribed standard;

(b) offers or agrees to supply any such drug or drug product; or

(c) exposes or possesses such drug or drug product for supply.

and is liable on summary conviction to a fine of $10,000 or imprisonment for 6 months, or both.

[Section 31A inserted by 2013 : 48 s. 18 effective 24 December 2013]
Orders and notices to prohibit supply of a drug or drug product

31B  (1) The Minister may—

(a) make orders ("prohibition orders") prohibiting persons from supplying, or offering to supply, exposing for supply or possessing for supply any drug or drug product which the Chief Medical Officer considers is not safe and which are described in the orders;

(b) serve on any person a notice ("prohibition notice") prohibiting the person from supplying, or offering to supply, agreeing to supply, exposing for supply or possessing for supply any drug or drug product which the Chief Medical Officer considers is not safe and described in the notice;

(c) serve on any person a notice ("notice to warn") requiring the person to publish, in a form and manner and on occasions specified in the notice and at his own expense a warning about any drug or drug product so specified which the Chief Medical Officer considers is not safe and which the person supplies or has supplied.

(2) A person who contravenes a prohibition order, a prohibition notice or a notice to warn is liable on summary conviction to a fine of $10,000 or to imprisonment for 6 months, or both.

[Section 31B inserted by 2013 : 48 s. 18 effective 24 December 2013]

Suspension notices

31C  (1) Where the Minister has reasonable grounds for suspecting that any health and safety requirement provided in section 31A has been contravened in relation to any drug or drug product, he may serve a notice ("a suspension notice") prohibiting the person on whom it is served, for such period ending not more than six months after the date of the notice as specified therein, from supplying the drug or drug product, offering to supply them, agreeing to supply them or exposing them for supply without the consent of the Minister.

(2) A suspension notice shall—

(a) describe the drug or drug product in a manner to sufficiently identify it;

(b) set out the grounds on which the Minister suspects that a safety provision has been contravened in relation to the drug or drug product; and

(c) state that the person on whom the notice is served may apply under section 31D for an order setting aside the notice.

(3) The consent of the Minister under subsection (1) may impose such conditions on the doing of anything for which the consent is required as the Minister considers appropriate.

(4) Any person who contravenes a suspension notice is guilty of an offence and is liable on summary conviction to a fine of $10,000 or to imprisonment for 6 months, or both.

[Section 31C inserted by 2013 : 48 s. 18 effective 24 December 2013]
**Application to set aside a suspension notice**

31D  (1) Any person having an interest in any drug or drug product in respect of which a suspension notice is in force may apply to a magistrate for an order setting aside the notice.

(2) On an application under subsection (1), the magistrate shall not make an order setting aside the suspension notice unless he is satisfied that there has been no contravention of any safety provision in relation to any drug or drug product.

[Section 31D inserted by 2013 : 48 s. 18 effective 24 December 2013]

**Power to obtain information**

31E  (1) If the Minister considers that, for the purpose of deciding whether to make, vary or revoke a prohibition order or to serve, vary or revoke a prohibition notice or to serve or revoke a notice to warn, he requires information which another person is likely to be able to furnish, the Minister may serve on the other person a notice requiring the person—

(a) to furnish to the Minister within a period specified in the notice, such information as is so specified;

(b) to produce such documents as are specified in the notice at a time and place so specified and to permit a person appointed by the Minister for the purpose of taking copies of the documents at that time and place.

(2) A person is guilty of an offence if he—

(a) fails, without reasonable cause, to comply with a notice served on him under subsection (1); or

(b) in purporting to comply with a requirement which by virtue of subsection (1)(a) is contained in a notice served on him under that subsection, furnishes information which he knows is false in a material particular or recklessly furnishes information which is false in a material particular.

(3) A person guilty of an offence under—

(a) subsection 2(a) of that subsection, is liable on summary conviction to a fine of $10,000 or to imprisonment for 6 months; and

(b) subsection 2(b) of that subsection, is liable on summary conviction to a fine of $15,000 or to imprisonment for 12 months.

(4) No information obtained by virtue of this section shall be disclosed except—

(a) for the purpose of any criminal proceedings or any investigation with a view to such proceedings;

(b) for the purpose of enabling the Minister to decide whether to make, vary or revoke safety regulations or a prohibition order or whether to serve, vary or revoke a prohibition notice or to serve or revoke a notice to warn; or
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(c) in a prohibition notice, a notice to warn or a warning published as required by a notice to warn or in a warning about goods which is published by the Minister;

but the prohibition on disclosure imposed by this subsection does not apply to publicised information.

(5) A person who discloses information in contravention of subsection (4) is liable on summary conviction to a fine of $10,000 or to imprisonment for 6 months, or both.

[Section 31E inserted by 2013 : 48 s. 18 effective 24 December 2013]

PART VII

CONTROL OF POISONS

Prohibition of sale of poison without licence
32 Subject to the provisions of this Part, it shall be unlawful for a person to offer for sale, or sell, any poison unless he holds a licence for the purpose under section 34.

Poisons
33 Any substance specified in the Fifth Schedule shall be a poison for the purposes of this Act.

Licences to sell poisons
34 (1) Any person who makes application to the Minister in the prescribed form and pays the appropriate fee provided for under the Government Fees Act 1965 [title 15 item 18] may be granted a licence by the Minister under this section, and a person holding such a licence is in this Act referred to as a "licensed seller of poisons".

(2) The Minister may refuse to grant a licence under this section to any person who for any reason relating to that person or his premises and appearing to the Minister to be sufficient is not fit to hold such a licence.

(3) A licence under this section shall not entitle the holder to sell any poison for a use that is a medicinal use, and it shall accordingly be an offence against this Act for a licensed seller of poisons to sell a poison if he knows or has reason to believe that the poison will be applied to a use that is a medicinal use.

(4) The licence of a licensed seller of poisons shall lapse if he does not on or before the 31st day of December pay to the Minister the appropriate annual fee provided for under the Government Fees Act 1965 [title 15 item 18].

(5) A list of the licensed sellers of poisons shall be published annually in the Gazette so soon as may be after the 1st day of January, and particulars of any alterations made in the number of licensed sellers of poisons on or after that date in any year shall also be published in the Gazette.
Revocation of licences
35 Subject to section 36, the Minister may revoke a licence granted under section 34 for any reason such as is mentioned in section 34(2), and shall give notice in writing to the holder of the licence of his decision to revoke the licence and the reasons for the decision.

Appeals
36 (1) Any person aggrieved by the revocation of a licence under section 35 may within twenty-eight days after receiving notice of the decision appeal to the Supreme Court, and upon any such appeal that Court shall have power to confirm or revoke the Minister’s decision.

(2) The powers of the Chief Justice to make rules under section 62 of the Supreme Court Act 1905 shall extend to the making of rules regulating the practice and procedure to be followed on, and the fees to be paid in connexion with, any such appeal.

(3) The Minister may appear as respondent on any such appeal and, whether he appears at the hearing of the appeal or not, he shall be deemed to be a party to the appeal for the purpose of enabling directions to be given as to the costs or expenses of the appeal.

(4) Where an appeal is not brought against the decision of the Minister to revoke a licence under section 35, or where an appeal is brought but is withdrawn or struck out for want of prosecution, the decision shall take effect on the expiration of the time for appealing or, as the case may be, on the withdrawal or the striking out of the appeal; but otherwise such a decision shall take effect if and when the appeal is dismissed and not otherwise.

Poisons Book
37 (1) Every person who sells poison shall maintain a book (in this Act called the “Poisons Book”) in such form as the Minister may approve for the purpose of keeping the records called for by subsection (2).

(2) Every such person shall enter and keep in the Poisons Book, in relation to every sale by him of a poison, a record of—

(a) the date of the sale;

(b) the kind and quantity of the poison sold;

(c) the name and address of the purchaser; and

(d) the purpose stated by the purchaser for the purchase,

and he shall obtain the signature of the purchaser to, and himself sign, the entry in the Book.

Sale of poisons to unknown persons prohibited
38 A person shall not sell poison to any person that is not known to him, except in the presence of a third person who—

(a) is known to the seller; and
(b) declares to the seller that the purchaser is known to him; and
(c) in confirmation of his declaration signs the entry in the Poisons Book.

Sale of poisons to persons under 18 prohibited
39 (1) Subject to subsection (2), it shall be an offence against this Act for any person
to sell poison to a person under 18 years of age.

(2) It shall be a defence for a person charged with an offence against subsection
(1) that he believed on reasonable grounds (the proof whereof shall be on him) that the
purchaser was 18 years or over.

Labelling of poisons
40 Subject to any provision made by any regulation, no person shall sell any poison
to any other person unless the word “poison” and the name and business address of the
seller and the date of the sale are displayed in clear and legible writing on the surface of the
receptacle in which the poison is contained.

Sale of poisons to intoxicated persons etc. prohibited
41 It shall be an offence against this Act for any person to sell poison to another person
whom he knows, or has cause to believe, to be intoxicated by drink or drugs or to be of
unsound mind.

Method of keeping poisons
42 In the keeping of poisons it shall be the duty of every licensed seller of poisons to
ensure—

(a) that every bottle, vessel, box or package containing poison has attached to
it a label bearing the name of the article and also some distinctive mark to
show that poison is contained therein;

(b) that poison is kept in accordance with one or other of the following systems,
that is to say,—

(i) in a bottle or vessel tied over, capped, locked or otherwise secured in a
manner different from that in which vessels containing articles that are
not, or do not contain, poison are secured in the same premises;

(ii) in a bottle or vessel rendered distinguishable by touch from bottles or
vessels in which articles that are not, or do not contain, poison are kept
in the same premises;

(iii) in a bottle, vessel, box or package kept in a room or cupboard set apart
for dangerous articles.

Statement of proportion of poison in preparations
43 (1) Subject to subsections (2) and (3), it shall be the duty of every person selling
a preparation containing poison to ensure that there is set out on a label attached to the
preparation the proportion, whether expressed as a percentage or otherwise, which such poison bears to the total content of the preparation.

(2) In the case of a preparation listed in the official British Pharmacopoeia or the British Pharmaceutical Codex or any supplement thereto, it shall be a sufficient compliance with subsection (1) if that preparation—

(a) when sold either with or without dilution or admixture, is described by its name or synonym or abbreviated name used in the Pharmacopoeia, Codex or supplement with the addition of the letters B.P. or B.P.C., as the case may be; and

(b) when sold with dilution or admixture, is described by the proportion which the preparation bears to the mixture of which it forms a part.

Liquid preparations containing poison

It shall be the duty of every person selling any liquid preparation containing poison to ensure—

(a) that the preparation is not sold otherwise than in bottles, tins, drums or casks sufficient to withstand without leakage the ordinary risks of transit;

(b) that every such bottle, tin, drum or cask has the legend “Poison - not to be taken internally” indelibly printed, marked or branded in easily legible letters in a conspicuous position apart from the label, and that there is thereon a label bearing the same legend; and

(c) when such a liquid is sold in bottles, that such bottles are of a distinctive character so as to be easily distinguishable by touch from other bottles.

PART VIII
MISCELLANEOUS

Wholesale transactions

Subject to any provision made by any regulation, no person shall by wholesale sell or otherwise dispose of any schedule 3 or Schedule 4 drug or poison to any person that is not entitled to sell that drug or poison by retail.

(2) A sale or disposal of a drug or poison is a sale or disposal by wholesale for the purposes of this Act if it is a sale or disposal to a person who buys or receives the drug or poison for the purpose of selling or disposing of the drug or poison to some other person; and in this Act “sale by retail” or “sell by retail” means sale or sell otherwise than by wholesale.

Dispensing records

Where any person supplies a Schedule 3 or Schedule 4 drug or poison (hereafter in this section referred to as a “substance”) under a prescription—
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(a) he shall mark in clear and legible writing on a paper accompanying the substance—

(i) his initials;

(ii) his name, address and telephone number (if any) or, where the substance is supplied from a registered pharmacy, the name, address and telephone number (if any) of the registered pharmacy;

(iii) the name of the customer to whom the substance is supplied;

(iv) the directions for using the substance;

(v) the number assigned to the prescription;

(vi) the quantity of the substance supplied;

(vii) the brand or trade name, the generic name, the name of the manufacturer and the strength of the substance supplied;

(viii) whether the prescription is to be refilled, and if so, the number of times;

(ix) the date when the prescription is filled; and

(x) the name of the practitioner who issued the prescription;

(b) he shall, or, where the substance is supplied from a registered pharmacy, the operator of the pharmacy shall, for the period of two years (or, where the prescription was repeated, two years after the last time it was repeated) retain the original of the prescription.

[Dishonest sales

47 It shall be an offence against this Act for any person keeping for sale, or offering for sale, or selling, any Schedule 3 or Schedule 4 drug or poison falsely to represent to any person—

(a) that it is a substance that it is not; or

(b) that it contains a substance that it does not contain; or

(c) that it is unadulterated when it has been adulterated.

Regulations for Parts VI and VII

48 (1) The Minister may make regulations under this section—

(a) prescribing the amount or proportion of any substance that is to be contained in a Schedule 3 or Schedule 4 drug or a poison;

(b) prescribing the types of, and labelling for, containers to be used for containing a Schedule 3 or Schedule 4 drug or a poison;

(c) regulating the manner in which, and the conditions subject to which, Schedule 3 or Schedule 4 drugs or poisons are to be prescribed by
practitioners, including the conditions under which Schedule 3 or Schedule 4 drugs or poisons may be supplied on a second or subsequent occasion without a further prescription having to be prepared;

(d) regulating the manner in which records are to be kept of the purchase and sale of Schedule 3 or Schedule 4 drugs or poisons;

(e) designating poisons that may be sold by persons not otherwise authorized by this Act for the purpose, and authorizing and regulating the sale of such poisons by such persons or by classes of such persons;

(f) designating Schedule 3 or Schedule 4 drugs and poisons that may be sold by persons not otherwise authorized by this Act for the purpose, and authorizing and regulating the sale without prescription by such persons or by classes of such persons of such drugs and poisons to owners of birds or animals for the treatment of the birds or animals;

(g) [deleted]

(h) generally for carrying out the purposes of sections 31A, 31B, 31C, 31D, 31E, 51, 51A and 51B.

(2) Regulations made under this section shall be subject to the negative resolution procedure.

[Section 48 subsection (1) amended by 2013 : 48 s. 20 effective 24 December 2013; subsection (1)(g) deleted by 2014 : 36 s. 2 effective 22 December 2014]

Minister may by order amend the Third or Fourth Schedule

48A (1) The Minister may, on the recommendation of the Chief Medical Officer or the Council, by order amend the Third or Fourth Schedule.

(2) The negative resolution procedure shall apply to an order made under subsection (1).

[Section 48A inserted by 2014 : 36 s. 2 effective 22 December 2014]

Minister may provisionally add or remove drugs in the Third or Fourth Schedule

48B The Minister may, on the recommendation of the Chief Medical Officer or the Council, by Notice in the Gazette, provisionally list, or remove, the drugs in the Third or Fourth Schedule and such drugs shall be considered listed in, or removed from, the Third or Fourth Schedule for a period not exceeding 30 days or until the Minister issues an order either adding to, or deleting from, the Third or Fourth Schedule such drugs, whichever occurs earlier.

[Section 48B inserted by 2014 : 36 s. 2 effective 22 December 2014]

Minister may obtain reports on drugs and poisons

49 (1) The Minister may by notice in writing served upon any practitioner or any registered pharmacist require him to report to the Minister in writing the quantity of any
Schedule 3 or Schedule 4 drug or any poison that he has purchased or sold or, in the case of a practitioner, prescribed, as the case may be, during the period stated in the notice.

(2) Where—

(a) the Minister has reason to believe (whether or not because of a report made to him pursuant to a notice served under subsection (1)) that a practitioner or a registered pharmacist has purchased or sold, or a practitioner has prescribed, excessive or otherwise unreasonable amounts of a Schedule 3 or Schedule 4 drug or a poison during a particular period; or

(b) a practitioner or registered pharmacist fails to make a report that he has been properly required under subsection (1) to make; or

(c) a report such as aforesaid appears to the Minister to be incomplete,

then, but without prejudice to any other power that is available to the Minister or any other person, the Minister may report the matter to the Council in the case of a registered pharmacist, or to the relevant professional body in the case of a practitioner, for such action as the Council or that body may think fit to take.

(3) In subsection (2) “relevant professional body”, in relation to a practitioner, means the body appearing to the Minister to be the body having professional disciplinary control over the practitioner.

Minister may obtain information on prices

50 (1) The Minister may by notice in writing under this section served upon any practitioner or the operator of any registered pharmacy require him to supply to the Minister in writing such information as may be specified pursuant to subsection (2).

(2) A notice under this section may demand information relating to—

(a) the price at which any substance was purchased by any person; and

(b) the price at which any substance was sold by any person to any member of the public,

in the conduct, or for the purposes, of the practice of the practitioner or the business of the registered pharmacy as the case may be during the period specified in the notice, and may demand any other information relating to, or connected with, the prices of substances so purchased or sold which the Minister may consider is required for establishing whether the prices charged to the public for such substances during the period were fair and reasonable.

Inspections

51 (1) It shall be the duty of the Minister, by means of inspection and otherwise, to take all reasonable steps to enforce, and secure compliance by registered pharmacists and others with the provisions of this Act or any regulation, and the Minister shall for that purpose appoint such number of inspectors as in his opinion is required.
(2) Any inspector may, for the purposes of enforcement of this Act or any regulations, make test purchases or otherwise ascertain whether any provisions of this Act or any regulations or of an order under this Act are being complied with.

(3) An inspector appointed under this section who has reasonable cause to believe that an offence under this Act or regulations has been committed shall, for the purpose of enforcing and securing compliance with the said provisions have power—

(a) at all reasonable times and on production, if required, of his credentials, enter any registered pharmacy or place of business (other than premises or parts of premises used as a dwelling house) and while there he may—

(i) inspect any drug or drug product found;

(ii) examine any procedure;

(iii) seize and detain drugs or drug products for testing;

(iv) seize and detain goods or documents which he believes may be required as evidence in proceedings under this Act;

(v) for the purpose of exercising his powers to seize drugs or drug products under this section and to the extent that it is reasonably necessary in order to ensure compliance with any provision of this Act, require any person having authority to do so to break open any container, and if the person does not comply, the inspector may do so himself.

(4) An inspector who seizes drugs, drug products or documents in exercise of his powers under subsection (3) shall, in a written statement specifying the nature and amount of items seized, inform the person from whom they are seized.

(5) For the purpose of proceedings taken or transactions made under this Act, the written statement of an inspector given under subsection (4) has effect as a receipt for the drug, drug products or documents seized.

(6) A magistrate who is satisfied by sworn information in writing that there are reasonable grounds to believe that—

(a) goods, books or documents which an inspector has power to inspect are on any premises and that their inspection is likely to disclose evidence of the commission of an offence under this Act or the regulations; or

(b) an offence under this Act or the regulations has been, is being, or is about to be committed on any premises;

and that—

(c) admission to the premises has been or is likely to be refused and that notice of intention to apply for a warrant under this subsection has been given to the occupier; or

(d) an application for admission or the giving of the notice mentioned in paragraph (c) would defeat the object of the entry or that the premises are
unoccupied or that the occupier is temporarily absent and it might defeat
the object of the entry to await his return,

may by warrant under his hand, which shall continue in force for a period of one month,
authorise any inspector to enter the premises, if need be by force.

(7) An inspector who enters premises by virtue of this section may take with him
such other persons and equipment as appears necessary to him, and on leaving premises
which he enters by virtue of a warrant under subsection (6), where either the premises are
unoccupied or the occupier is temporarily absent, he shall affix a notice in a conspicuous
place stating that the premises were entered for the purpose of this section, and as far as
practicable shall leave the premises as effectively secured against trespassers as he found
them.

(8) A person who—

(a) wilfully obstructs an inspector acting in the exercise of any power conferred
on him under subsections (3) to (7);

(b) wilfully fails to comply with any requirement properly made to him by an
inspector under subsections (3) to (7);

(c) without reasonable cause fails to give an inspector acting under
subsection (3) to (7), such assistance or information as he may reasonably
require of the person for the performance of the inspector’s functions;

(d) in giving information as mentioned in paragraph (c) makes a statement
which he knows to be false;

(e) not being an inspector purports to act as an inspector under this Act;

(f) discloses to another person, where the disclosure is not made in the
performance of his duty—

(i) information with respect to a manufacturing process or trade secret
obtained by him in premises which he has entered by virtue of
subsections (3) to (7); or

(ii) information otherwise obtained by him under this Act,

is guilty of an offence and is liable on summary conviction to a fine of
$10,000 or imprisonment for 6 months, or both.

(9) An inspector appointed under this section shall have power with the consent
of the Minister to institute summary proceedings in respect of an offence against this Act
or any regulation, and to conduct any such proceedings notwithstanding that he is not a
barrister and attorney.

(10) If a person wilfully delays or obstructs an inspector in the exercise of any of his
powers under this section, or refuses to allow any sample to be taken in accordance with
the provisions of this section, or fails without reasonable excuse to give any information
which he is duly required under this section to give, he is guilty of an offence against this Act.

Section 51 repealed and replaced by 2013 : 48 s. 22 effective 24 December 2013

Notice of test

51A (1) Where drugs or drug products seized or purchased by an inspector in pursuance of this Act are submitted to a test, the inspector shall—

(a) if the drugs or drug products were seized, inform the person from whom they were seized of the result of the test;

(b) if the drugs or drug products were purchased and the test leads to proceedings for an offence under this Act, inform the person from whom the goods were purchased of the result of the test;

and where as a result of the test proceedings for an offence are instituted against a person, the inspector shall allow the person to have the goods tested independently if it is reasonably practicable to do so.

(2) The Minister may by order provide for the testing of drugs or drug products seized or purchased by an inspector in pursuance of this Act and in particular may in those orders provide that the test be carried out at the Ministry's expense in a manner, by a person, and at a laboratory or testing facility specified in the order.

Section 51A inserted by 2013 : 48 s. 23 effective 24 December 2013

Compensation

51B (1) Where in the exercise of his powers under section 51 an inspector seizes and detains any drugs or drug products, and the owner suffers loss by reason of the goods being seized or by reason that, during the detention, the goods are lost or damaged or deteriorate, unless the owner is convicted of an offence under this Act committed in relation to the goods, the owner is entitled to compensation for the loss so suffered.

(2) Any disputed question as to the right to or the amount of any compensation payable under this section shall on the written application of the owner or of the Attorney-General be determined as follows—

(a) if the amount of the compensation claimed does not exceed $10,000, by a magistrate; or

(b) if the amount of the compensation claimed exceeds $10,000, by a judge of the Supreme Court,

in like manner as if the magistrate or the judge were a single arbitrator appointed pursuant to the provisions of the Arbitration Act 1986, and the provisions of that Act shall apply accordingly.

Section 51B inserted by 2013 : 48 s. 23 effective 24 December 2013
PHARMACY AND POISONS ACT 1979

Service of documents
52 Any notice or other document required or authorized by any provision of this Act to be served on any person, or to be given or sent to any person, may be served, given or sent—

(a) by delivering it to him; or

(b) by sending it by post to him at his usual or last-known residence or place of business in Bermuda; or

(c) in the case of a body corporate, by delivering it to the secretary or clerk of the body corporate at its registered or principal office or sending it by post to the secretary or clerk of that body corporate at that office.

Transitional
53 (1) Every person who immediately before 1 January 1980 was registered as a pharmacist under the Pharmacists Registration Act 1928 (now repealed) shall be deemed on and after that date to be a registered pharmacist within the meaning of this Act, but subject to the provisions of this Act.

(2) For such period (and no longer) beginning on 1 January 1980 as the Minister may appoint for the purpose by notice made under this subsection and published in the Gazette every set of premises which immediately before that date was being operated as a pharmacy, being premises to which this subsection applies, shall be deemed to be a registered pharmacy within the meaning of this Act.

(3) Subsection (2) applies to premises in respect of which the operator of those premises notifies the Minister in writing by 1 February 1980 of his wish to have the benefit of that subsection apply to those premises.

(4) Every person who immediately before 1 January 1980 was the holder of a licence granted to him under section 2 of the Poisons Act 1930 (now repealed) shall be deemed on and after that date to be a licensed seller of poisons within the meaning of this Act, but subject to the provisions of this Act.

Student pharmacists
54 (1) Nothing in section 6 shall have effect in relation to a student pharmacist acting in accordance with a permit granted to him under this section.

(2) The Minister may grant a permit under this section to a student pharmacist to compound or dispense any substance specified in the Third, Fourth or Fifth Schedule, subject to the conditions specified in the permit.

(3) A permit under this section must contain a condition that the permit-holder when acting under the permit shall do so under the direct personal control and supervision of a registered pharmacist who is named in the permit and who has endorsed the permit in acknowledgement of his responsibility thereunder; and (but without prejudice to any liability of the permit-holder apart from this Act) any act done by the permit-holder under, or in reliance upon, the authority of the permit shall for the purposes of this Act be deemed to be the act of that registered pharmacist.
(4) The Minister may without notice at any time in writing revoke a permit granted under this section.

(5) In this section “student pharmacist” means a person who has satisfied the Minister that he is undergoing a course of training that will qualify him in due course to receive a certificate of competence from the Council.

Offences

55 (1) Any person who contravenes or fails to comply with any duty or prohibition imposed upon him by or under any provision to which this section applies commits an offence against this Act.

(2) The provisions to which this section applies are sections 6, 23, 25, 27 to 30, 32, 37, 38, 40, 42 to 46, 49 and 50.

(3) Any person committing an offence against this Act may be proceeded against either summarily or on indictment—

(a) Punishment on summary conviction: imprisonment for 12 months or a fine of $5,000, or both such imprisonment and fine;

(b) Punishment on conviction on indictment: imprisonment for 3 years or a fine of $15,000, or both such imprisonment and fine;

(4) The power to make regulations under section 15, 22 or 48 includes the power to constitute offences for contravention of, or failure to comply with, any such regulation and to fix punishments, including imprisonment (but not exceeding the scale of punishments for which subsection (3) of this section provides), for any such offence.

(5) Where an offence committed against this Act or any regulation by a body corporate is proved to have been committed with the consent or connivance of, or to be attributable to any neglect on the part of, any director, manager, secretary or other similar officer of the body corporate, or any person who was purporting to act in any such capacity, he, as well as the body corporate, commits an offence against this Act and is liable to be proceeded against and punished accordingly.

Repeal

56 [omitted]

Commencement

57 [omitted]
THE PHARMACY COUNCIL

1 A member of the Council shall hold office for the period of one year or for such longer or shorter period as the Minister may determine.

2 A member of the Council shall be eligible for re-appointment or re-election to membership of the Council.

3 A person appointed or elected to fill the place of a member of the Council who vacates office before the expiry of his term of office shall hold office for so long only as the member whose place he fills would have held the office.

4 Where a member of the Council vacates his office three months or less before the expiry of his term of office, the vacancy need not be filled.

5 A member of the Council may resign his office at any time by giving notice in writing to the Minister of his resignation.

6 The Minister may declare the office of a member of the Council vacant if—
   (a) the Minister is satisfied that the member is unable through mental or physical incapacity to perform the functions of his office; or
   (b) the member has failed without adequate cause to attend three successive meetings of the Council.

7 The Council may act notwithstanding any vacancy in their membership, and no act of the Council shall be invalid by reason only of a defect in the appointment of a member.

8 Subject to the foregoing provisions of this Schedule the Council may regulate their own procedure.

9 (1) The Council may, in its discretion, appoint from among its own members or from among other persons, such number of committees as it thinks fit for purposes which, in the opinion of the Council, would be more expeditiously carried out or managed by such committees.

   (2) The Chairman of any committee appointed under subsection (1) shall be a member of the Council.
10  [deleted]

[First Schedule amended by 2013: 48 s. 24 effective 24 December 2013; paragraph 10 deleted by 2014: 36 s. 2 effective 22 December 2014]
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SECOND SCHEDULE

LIST OF PRACTITIONERS

Physician- for the purposes of medical treatment of human beings

Dentist- for the purposes of dental treatment of human beings

Veterinary Practitioner- for the purposes of animal treatment

Optometrist- subject to the restrictions and requirements under section 10 and Schedule 2 of the Optometrists and Opticians Act 2008

Advanced Practice Nurse- subject to the restrictions and requirements under 8B(1) and (2) of the Nursing Act 1997

[Second Schedule repealed and replaced by 2013 : 48 s. 25 effective 24 December 2013]
THIRD SCHEDULE

(DRUGS OBTAINABLE ONLY ON PRESCRIPTION EXCEPT WHERE SPECIFIED IN THE FOURTH SCHEDULE (PART I AND PART II))

Note: The following annotations used in this Schedule have the following meanings:

md        (maximum dose)  i.e. the maximum quantity of the substance contained in the amount of a medicinal product which is recommended to be taken or administered at any one time.

mdd       (maximum daily dose) i.e. the maximum quantity of the substance that is contained in the amount of a medicinal product which is recommended to be taken or administered in any period of 24 hours.

mg        milligram

ms         (maximum strength)  i.e. either or, if so specified, both of the following:

(a) the maximum quantity of the substance by weight or volume that is contained in the dosage unit of a medicinal product; or

(b) the maximum percentage of the substance contained in a medicinal product calculated in terms of w/w, w/v, v/w, or v/v, as appropriate.

external use means for application to the skin, teeth, mucosa of the mouth, throat, nose, eye, ear, vagina or anal canal when a local action only is necessary and extensive systemic absorption is unlikely to occur. Note: the following are not regarded as for external use: throat sprays, throat pastilles, throat lozenges, throat tablets, nasal drops, nasal sprays, nasal inhalations or teething preparations.

parenteral use means administration by breach of the skin or mucous membrane.

1
Abacavir
ABC Liniment
Acamprosate
Acarbose
Acebutolol
Acepiffline
Acepromazine
Acetanilide
Acetarsol
Acetazolamide
Acetohexamide
Acetorphine
Acetrizoic Acid
Acetyl Sulphafurazole
Acetyl Sulphamethoxypyridazine
Acetylcarmbromal
Acetylccholine
Acetylcysteine
Acetyldigitoxin
Acetyldihydrocodeine
Acetylpromazine
Acetylstrophanthidin
Acitretin
Aconiazide
Aconite Belladonna and Chloroform Liniment BPC 1968
Aconite Root
Aconitine
Actinomycin C
Actinomycin D
Acyclovir
Adapalene
Adefovir
Adicillin
Adiphenine
Admune Influenza Vaccine
Adrenaline
Adrenocortical Extract
Adriamycin
Aerosoxacin
Aesculin
Agomelatine
Albamycin preparations
Albamycin T preparations
Albumin Human
Albumin Microspheres Human (3M)
Albuterol
Alclofenac
Alcuronium Chloride
Aldosterone
Alendronate
Alfacalcidol
Alfentanil
Alfuzosin
Algestone
Algestone Acetonide
Algestone Acetophenide
Aliskiren
Alitretinoin
Alkavervir
Allobarbitone
Allopurinol
Allyloestrenol
Allylprodine
Almotriptan
Alphacetylmethadol
Alphadolone Acetate
Alphameprodine
Alphamethadol
Alphaprodine
Alphaxalone
Alprazolam
Alprenolol
Alprostadil
Alseroxylon
Amantadine
Ambenonium Chloride
Ambrisantan
Ambuside
Ambutonium Bromide
Aminocaproic Acid
Amikacin
Amiloride
Aminodarone
Aminogluthimide
Aminophylline
Aminopterin
Aminorex (and Methyl Derivative)
Aminosalicylic Acid
Amiodarone
Amiphenazole
Amitriptyline
Amlodipine
Ammonium Bromide
Ammonium Chloride - in inhalers
Amoxycillin
Amoxycillin Trihydrate
Amphetamine
Amphomycin
Amphotericin
Ampicillin
Ampicillin Trihydrate
Amyl Nitrite Vitrellae BPC
Amylobarbitone
Amylocaine - in preparations for local ophthalmic use
Anaesthetics - all inhalational
Anagrelide
Anastrozole
Ancrod
Androsterone
Aneurine
Angiotensin Amide
Anileridine
Antazoline
Anterior Pituitary Extract
Anti-lymphocyte Immunoglobulin
Antimony
Apiol
Apomorphine
Apramycin
Aprepitant
Aprobarbitone
Aprotinin
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Arecoline
Arecoline-acetarsol
Aripiprazole
Arprinocid
Arsanilic Acid
Arsenic
Arsphenamine
Asparaginase
Asparaginase, L-Atenolol
Atamestane
Atazanavir
Atenolol
Atomoxetine
Atorvastatin
Atovaquone
Atracurium Besylate
Atropine Eye Drops B.P. - in preparations for local ophthalmic use
Atropine Eye Ointment B.P. - in preparations for local ophthalmic use
Atropine - in inhalers
Atropine Methobromide - in preparations for local ophthalmic use
Atropine Methobromide - in inhalers
Atropine Oxide - in preparations for local ophthalmic use
Atropine Oxide - in inhalers
Azacyclonol
Azaperone
Azapropazone
Azarbine
Azathioprine
Azelaic Acid
Azidocillin
Azithromycin
Bacampicillin
Bacitracin Methylene Disalicylate
Baclofen
Balsalazide
Bambermycin
Bamipine
Barbitone
Barbituric Acid - and derivatives
Barium Carbonate
Barium Chloride
Barium Sulphate
Barium Sulphide
Beclamide
Beclomethasone
Belladonna Herb
Belladonna Root
Bemegride
Benactyzine
Benapryzine
Benazepril
Bendazac
Bendrofluazide
Benethamine Penicillin
Benoxaprofen
Benperidol
Benserazide
Benzaflbrate
Benzathine Penicillin
Benzbromarone
Benzestrol
Benzethidine
Benzhexol
Benzilonium Bromide
Benzocaine - for local ophthalmic use
Benzocaine
Benzocaine
Benzocaine
Benzoyl Peroxide - in concentrations greater than 10%
Benzoylsulphanilamide, N-
Benzphetamine
Benzquinamide
Benzthiazide
Benztrone Injections
Benztropine Mesylate
Benzylmorphine
Benzylpenicillin
Betameprodine
Betamethasone
Betamethadol
Betamethasone
Betaprodine
Betaxolol
Bethanechol Chloride
Bethanidine
Bexarotene
Bezafibrate
Bezitramide
Bicalutamide
Bimatoprost
Biorphen Oral Solution
Biperiden
Bismuth Glycollylarsanilat
Bisoprolol
Bleomycin Sulphate
Bolandiol
Bolasterone
Bolazine
Boldenone Undecylenate
Bolanol
Bolmantalate
Bosentan
Bretylium Tosylate
Brimonidine
Brinzolamide
Bromazepam
Bromhexine
Bromocriptine
Bromvaletone
Brotoplan
Budesonide
Bufexamac
Bufotenine
Bumetanide
Buphenine
Bupivacaine - in preparations for local ophthalmic use
Buprenorphine
Bupropion
Buspirone
Busulphan
Butacaine - in preparations for local ophthalmic use
Butalbital
Butanilicaine - in preparations for local ophthalmic use
Butaperazine
Butobarbitone
Butorphanol
Butriptyline
Butylchloral Hydate
Cabergoline
Cadexomer
Calcipotriol
Calcitriol
Calcitonin
Calcitriol
Calcium 5-allyl-5-N-Butylbarbiturate
Calcium Acetate
Calcium Aminosalicylate
Calcium Amphomycin
Calcium Benzamidosalicylate
Calcium Bromide
Calcium Bromolactobionate
Calcium Carbimide
Calcium Folinate
Calcium Leucovorin preparations
Calcium Sulphaloxate
Calusterone
Camphorated Opium tincture BP
Camazepam
Candesartan
Candicidin
Cannabidiol
Cannabidiol and derivatives
Cannabis
Cannabis resin
Cantharadin
Capreomycin sulphate
Captodiamine
Captopril
Caramiphen
Carbachol
Carbamazepine
Carbenicillin
Carbenoxolone
Carbidopa
Carbidopa Monohydrate
Carbimazole
Carbocisteine
Carbon Tetrachloride
Carboxymethylcysteine
Carfecillin
Carfentanil
Carisoprodol
Carmustine
Carperidine
Carphenazine
Carvedilol
CCNU Capsules
Cathine
Cefaclor
Cefadroxil
Cefdinir
Cefixime
Cefotaxime
Cefoxitin
Cefpodoxime Proxetil
Cefsulodin
Cefuroxime
Celecoxib
Centella Asiatica - extract and active principals thereof (if for internal use)
Cephalexin
Cephaloglycin
Cephaloram
Cephaloridine
Cephalosporin C
Cephalosporin E
Cephalosporin N
Cephalothin Sodium
Cephamandole
Cephalizin Sodium
Cephradine
Cerium Oxalate
Chemocycline preparations
Chenodeoxycholic Acid
Chloral Antipyrine
Chloral Betaine
Chloral Formamide
Chloral Glycerolate
Chloral Hydrate
Chloralose
Chloralurethane
Chlorambucil
Chloramphenicol
Clorazepic acid
Chlordiazepoxide
Chlorhexadol
Chlorisondamine Chloride
Chlormadinone Acetate
Chlormerodrin
Chlormethiazole
Chlormezanone
Chlorodyne BPC
Chloroform - for inhalational use
Chloroform and Morphine Tincture BPC
Chloroquine
Chlorothiazide
Chlorotrianisene
Chlorphenoxamine
Chlorphentermine
Chlorpromazine
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Chlorthalidone
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<tr>
<td>Chlorzoxazone</td>
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Clopidogrel
Cloprostenol Sodium
Clorazepate
Cloroxolone
Clorprenaline
Clostebol Acetate
Clotiazepam
Clotrimazole
Cloxacillin
Cloxazolam
Clozapine
Cocaine
Cocculus Indicus
Cocillana Compound Syrup BPC 1949
Codeine - for non-parental use with ms greater than 8mg calculated as base
Co-dergocrine Mesylate
Colchicine
Colestipol
Colistin
Collagen preparations - if for implantation under the skin
Collagenase - when sold or recommended as a debriding agent
Colocynth and Jalap Compound Tablets BPC 1963
Conine
Conium Leaf
Contraceptives - oral
Corticotrophin
Cortisone
Cortodoxone
Cotarnine Chloride
Co-Trifamole
Co-trimoxazole
Coumarin derivatives
Cropropamide
Crotamiton
Crotethamide
Croton Oil
Croton Seed
Cuemid
Curare
Cyclandelate - in nausea and vomiting in pregnancy
Cyclizine
Cycobarbitone
Cyclobenzaprine
Cyclofenil
Cyclomethycaine
Cyclopentamine
Cyclophosphamide
Cyclophosphamide
Cyclopsorin preparations
Cyclothiazide
Cycrimine
Cyproheptadine
Cyproterone Acetate
Cycrimine
Cytarabine
Dabigatran
Dacarbazine
Dactinomycin
Danazol
Dantrolene
Dapsone
Darifenacin
Dasatinib
Daunorubicin
Deanol
Debrisoquine
Deferasirox
Deferiprone
Dehydrocholic Acid
Dehydroemetine
Dehydroepiandrosterone
Delmadinone Acetate
Delorazepam
Demecarium Bromide
Demeclocycline
Deoxycortone
Deoxyribonuclease
Deptropine
Dequalinium Chloride
Deserpidine
Desferrioxamine
Desfluorotriamcinolone
Desipramine
Deslanoside
Desloratadine
Desmopressin
Desogestrel
Desomorphine
Desonide
Desoxymethasone
Dexamethasone
Dexamphetamine (Dextroamphetamine, Dexamfetamine)
Dexetimide
Dextranomer preparations - for medicinal use
Dextromethorphan
Dextromoramide
Dextropropoxyphene
Dextrothyroxine
Diamorphine
Diampromide
Diazepam
Diazoxide
Dibenzepin
Dichloralphenazone
Dichlorophenarsine
Dichlorphenamide
Diclofenac Sodium
Dicloxacillin
Dicobalt Edetate
Dicyclomine
Didanosine
Dienoestrol
Diethanolamine Fusidate
Diethyl Carbamazine Citrate
Diethyl Propion
Diethylamide Ethyl Benzilate
Diethylamine Acetarsol
Diethylstilboestrol - and derivatives if for medicinal use
Diethyliambutene
Difenoxin - (1-(3-cyano-3,3-diphenylpropyl)-4-phenyl piperidine-4-carboxylic acid)
Diflorasone
Diffucortolone Valerate
Diffunisal
Digitalis Leaf
Digitalis prepared
Digitoxin
Digoxin
Dihydergot preparations
Dihydrallazine Sulphate
Dihydrocodeine
Dihydrocodeinone O-Carboxymethylxime
Dihydroergocornine
Dihydroergocristine
Dihydroergocryptine
Dihydroergotamine
Dihydroergotoxine
Dihydromorphine
Dihydrostreptomycin
Di-iodohydroxquinoline
Diloxanide Furoate
Diltiazem
Dimenoxadole
Dimepheptanol
Dimepregnen
Dimercaprol
Dimethisoquin - in preparations for local ophthalmic use
Dimethisterone
Dimethothiazine
Dimethyl Sulphoxide
Dimethylthiambutene
Dimethyltubocurarine
Dinitrodiarylphenylsulphonylethyleneamine
Dinitrophenol, 2,4- and derivatives if for medicinal use
Dinoprost
Dinoprostone
Dioxaphethyl Butyrate
Diphenhydramine - for parenteral use
Diphenidol
Diphenoxylate
Diphetarsone
Diphylline
Dipipanone
Dipivefrin
Diprenorphine
Diprophylline
Dipropyltryptamine
Dipyridamole
Dipyrene
Disodium Etidronate
Disopyramide
Distigmine Bromide
Disulfiram
Disulphamide
Dobutamine
Domperidone
Donepezil Hydrochloride
Dopamine
Dorzolamide
Dothiepin
Doxapram
Doxazosin
Doxepin
Doxorubicin
Doxycycline
Doxycycline Calcium Chelate
Dronabinol (Marinol)
Dronedarone
Droperidol
Drostanolone
Drotebanol
Duloxetine
Dutasteride
Dydrogesterone
Dyflos
Ecgone - any derivative of ecgonine which is convertible to ecgonine or to cocaine
Econazole
Ecothiopate Iodide
Ectyl urea
Edogestrone
Edrophonium Chloride
Efavirenz
Efornithine
Eletriptan
Eltrombopag
Embutramide
Emepriprum Bromide
Emeside preparations
Emetine
Emtricitabine
Emylcamate
Enalapril
Enestebol
Enflurane - for inhalational use
Entacapone
Entecavir
Ephedrine
Ephedrine - in inhalers
Epicillin
Epinastine
Epioestriol
Epithiazide
Epitiostanol
Eplerenone
Epoprostenol
Ergometrine Maleate
Ergot - prepared
Ergotamine
Ergotoxine
Erlotinib
Erythrityl Tetrinitrate
Erythromycin
Escitalopram
Esomeprazole
Estazolam
Estradiol
Estramustine Phosphate
Etafedrine
Etamiphylline
Ethacrynite Acid
Ethambutol
Ethamivan
Ethamsylate
Ethanolamine Oleate
Ethchlorvynol
Ethabenecid
Ether - for inhalational use
Ethiaizide
Ethinamate
Ethinyloestradiol
Ethionamide
Ethisterone
Ethoglucid
Ethoheptazine Citrate
Ethopropazine
Ethosuximide
Ethotoin
Ethulose
Ethyl Acetanilide
Ethyl Alcohol - for internal use 45%
Ethyl Biscoumacetate
Ethyl loflazepate
Ethyl N-heptyloxyacetate - if for internal use
Ethylmethylthiambutene
Ethylmorphine - if for non-parenteral use and (a) in undivided preparations with ms 2.5% (calculated as base); or(b) in single-dose preparations with ms per dosage unit 100mg (calculated as base)

Ethylestrenol
Ethylstibamine
Ethynodiol Diacetate
Etidronate Disodium
Etodolac
Etomidate
Etonizazene
Etoposide
Etophine
Etoxeridine
Etravirine
Etretinate
Etymemazine
Everolimus
Exemestane
Exenatide
Ezetimibe
Factor XIII Concentrate
Factorate
Famciclovir
Famotidine
Famprofazone
Fazadinium Bromide
Febuxostat
Felodipine
Fencamfamin
Fenclofenac
Fenethylline
Fenfluramine (not in combination with Phentermine)
Fenofibrate
Fenoprofen
Fenoterol
Fenpipramide
Fenpiprane
Fentanyl
Fenproporex
Fentiazac
Fentin Compounds
Feprazone
Ferrous Arsenate
Ferrous salts - for parenteral use
Fesoterodine
Fexofenadine
Fibrinolysin
Finasteride
Flavoxate
Flecainide
Floctafenine
Florantyrone
Floxapen preparations
Fluanisone
Fluclorolone Acetonide
Flucloxacillin
Fluconazole
Flucytosine
Fludarabine
Fludiazepam
Fludrocortisone Acetate
Flufenamic Acid
Flugestone
Flumethasone
Flumethiazide
Flunitrazepam
Flunixin
Fluocinolone Acetonide
Fluocinonide
Fluocortolone
Fluromazine
Flurometholone

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Fluorouracil
Fluoxetine
Fluoxymesterone
Flupenthixol
Flupentixol
Fluperolene Acetate
Fluphenazine
Fluprednidene Acetate
Fluprednisolone
Fluprostanol
Flurandrenolone
Flurazepam
Flurbiprofen
Fluspirilene
Flutamide
Fluticasone
Fluvastatin
Fluvioxamine
Folic Acid
Follicle stimulating hormone
Formebolone
Formocortol
Formosulphathiazole
Formoterol
Fosfestrol Tetrasodium
Fosinopril
Framycetin Sulphate
Frovatriptan
Frusemide
Fumagillin
Furaltadone
Furazolidone
Furethidine
Furoxone preparations
Fusafungine
Fusidic Acid
Gabapentin
Galantamine
Gallamine Triethiodide
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Gefitinib
Gelsemine
Gelsemium
Gemfibrozil
Gentamicin
Gestrenol
Glafenine
Glibenclamide
Glibornuride
Gliclazide
Glimepiride
Glipamide
Glipizide
Gliquidone
Glutethimide
Glyburide
Glyceryl Trinitrate preparations
Glycopyrronium Bromide
Glymidine
Glytona
Gonadotraphon LH
Goserelin
Gramicidin
Granisetron
Gravigard
Griseofulvin
Growth hormone
Guanethidine
Guanoclor
Guanoxan
Hachimycin
Halazepam
Halcinonide
Halobetasol
Haloperidol
Haloprogan
Halopyramine
Halothane
Haloxazolam
Halquinol
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Heparin
Heptobarbitone
Heptaminol
Hetacillin
Hexachlorophane
Hexamethonium
Hexamine
Hexobarbitone
Hexoestrol
Histidine, L-
Homatropine -
Homatropine Hydrobromide
Homatropine Methylbromide
Hyaluronidase
Hydralazine
Hydargaphen
Hydrobromic Acid
Hydrochlorothiazide
Hydrocodone
Hydrocortamate
Hydrocortisone
Hydroflumethiazide
Hydrogen cyanide
Hydromorphinol
Hydromorphone
Hydroxy-3-nitrophenylarsonic Acid, 4-
Hydroxychloroquine
Hydroxycholecalciferol, 1,a-
Hydroxymethylgramicidin
Hydroxyphentidene
Hydroxyprogesterone
Hydroxyurea
Hydroxyzine
Hygromycin B
Hyoscine -
Hyoscine Butylbromide -
Hyoscine Hydrobromide -
Hyoscine Methobromide -
Hyoscyamine -
in preparations for local ophthalmic use
in inhalers
in inhalers
in inhalers
in inhalers
Hypnomidate Concentrate
Ibandronate
Ibuprofen
Idarubicin
Idoxuridine
Ifosfamide
Imatinib
Imipramine
Imiquimod
Immunoglobulins
Indapamide Hemihydrate
Indinavir
Indomethacin
Injectables - all
Injections - all preparations for human use
Inosine Pranobex
Intra-uterine contraceptive devices
Intravenous Fluids - all
Iodoxamic Acid
Iopanoic Acid
Ipratropium Bromide
Iprindole
Iproniazid
Irbesartan
Isoaminile
Isocarboxazid
Isoconazole preparations
Isoetharine
Isoflurane - if for inhalational use
Isomethadone
Isomethaptenene
Isoniazid
Isoprenaline
Isopropamide Iodide
Isopropylaminophenazone
Isosorbide Dinitrate preparations
Isosorbide Mononitrate preparations
Isotretinoin
Isoxsuprine
Isradipine
Itraconazole
Ivabradine
Jaborandi
Kanamycin Sulphate
Ketamine
Ketazolam
Ketobemidone
Ketoconazole
Ketoprofen
Ketorolac Trometamol
Ketotifen
Khellin
Labetolol
Lacosamide
Lamivudine
Lamotrigine
Lanatoside
Lansoprazole
Lanthanum
Lapatinib
Latamoxef
Latanoprost - in preparations for local ophthalmic use
Lead and Opium Lotion BPC 1959
Lead Arsenate
Lefetamine
Leflunomide
Letrozole
Leuprolide
Levatolorphan
Levetiracetam
Levocetirizine
Levodopa
Levofoxacin
Levomethorphan
Levomoramide
Levonorgestrel
Levophenacylmorphan
Levorphanol
Levothyroxine
Lidoflazine
Lignocaine - in preparations for local ophthalmic use
Lincomycin
Linezolid
Liothyronine
Lisdexamfetamine (Lisdexamphetamine)
Lisinopril
Lithium Carbonate
Lithium Sulphate
Lobeline
Lodoxamide
Lofentanil
Lofepramine
Lomustine
Loperamide
Lopinavir
Loprazolam
Lorazepam
Lorcaserin
Lormetazepam
Losartan
Loteprednol Etabonate
Loxapine
Luteinising hormone
Lynoestrenol
Lypressin
Mafenide
Magnesium Bromide
Magnesium Fluoride
Magnesium Glutamate
Mandragora Autumnalis
Mannomustine
Maprotiline
Maraviroc
Mazindol
Mebanazine
Mebeverine
Mebezonium Iodide
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Mebhydrolin
Mebolazine
Mecamylamine
Mechlorethamine
Mecillinam
Meclofenamic Acid
Meclofenoxate
Mecloqualone
Meclozine - if sold or recommended for the prevention of nausea of pregnancy
Medazepam
Medicinal Opium - if in preparations from which the opium cannot be readily recovered in amounts which constitute a risk to health and, also if in liquid preparations with ms 0.2% (calculated as anhydrous morphine base); in solid preparations with ms 0.2% (calculated as anhydrous morphine base)
Medigoxin
Medrogestone
Mefenamic Acid
Mefenorex
Mefloquine
Mefruside
Megestrol
Meglumine Diatrizoate
Melarsonyl Potassium
Melarsoprol
Melengestrol
Meloxicam
Melphalan
Memantine
Menadiol - if for parenteral route
Menotrophin
Mepazine
Mepenzolate
Mephenesin
Mephenoxolone
Mephentermine
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Mepitiostane
Mepivacaine - in preparations for local ophthalmic use
Meprobamate
Mepazinol
Mepyramine
Mequitazine
Mercaptorpurine
Mercuderamide
Mesabolone
Mersalyl Acid
Mesalamine
Mescaline
Mesna
Mesoridazine
Mestanolone
Mesterolone
Mestranol
Metabutethamine - in preparations for local ophthalmic use
Metaldehyde - if for medicinal use
Metaraminol
Metaxalone
Metazocine
Metformin
Methacycline
Methadone
Methadyl Acetate
Methallenoestril
Methandienone
Methandriol
Methaqualone
Metharbitone
Methazolamide
Methdilazine
Methenamine
Menholone
Methicillin
Methimazole
Methionine - all isomers
Methisazone
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Methixene
Methohexitone
Methoin
Methoserpidine
Methotrexate
Methotrimeprazine
Methoxamine
Methoxsalen
Methoxyflurane - for inhalational purposes
Methoxyphenamine
Methsuximide
Methylaminorex
Methyclothiazide
Methyl benzoinoate
Methyl-3-Piperidylbenzilate, N-
Methylacetanilide, N-
Methylamphetamine
Methyldesorphine
Methyldihydromorphine
Methyldihydromorphinone
Methyldopa
Methylephedrine
Methylergometrine
Methylergonovine
Methylparafynol
Methylpentynol
Methylphenidate
Methylphenobarbitone
Methylprednisolone
Methylsulphonol
Methyltestosterone
Methylthioaracil
Methyclothiazide
Methyprylone
Methysergide
Metiguanide Tablets
Metindizate
Metirosine
Metoclopramide
Metolazone
Metomidate
Metopon
Metopimazine
Metoprolol
Metribolone
Metronidazole
Metryrapone
Mexiletine
Mezlocillin
Mianserin
Mibolerone
Miconazole
Midazolam
Midodrine
Mifepristone
Minocycline
Minoxidil
Mirtazapine
Misoprostol
Mithramycin
Mitobronitol
Mitomycin C
Mitopodozide
Mitotane
Moclobemide
Modafinil
Moexipril
Molindone
Mometasone
Monensin
Monosulfram - for internal use
Montelukast
Morazone
Morpheridine
Morphine - in liquid preparations with ms 0.2% (calculated as anhydrous morphine base); in solid preparations with ms 0.2% (calculated as
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Morphine Methobromide
Moxifloxacin
Mupirocin
Mustine
Mycophenolate Mofetil
Myrophine

N-Ethylamphetamine
N-Ethyl-3-Piperidylbenzilate
Nabilone
Nabiximols (Sativex)
Nabumetone
Nadolol
Naftilin
Naftidrofuryl Oxalate
Nalbuphine
Nalidixic Acid
Nalorphine
Naloxone
Naltrexone
Nandrolone
Naphazoline
Naproxen
Narasin
Naratriptan
Natamycin
Nateglidine
Nealbarbitone
Nedocromil
Nefopam
Nelfinavir
Neoarsphenamine
Neocinchophen
Neomycin
Neostigmine
Nepafenac -

anhydrous morphine base); in pentavalent nitrogen derivatives
morphine N-Oxide and other pentavalent nitrogen morphine derivatives

in preparations for local ophthalmic use
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Nepenthe Oral Solution
Netilmicin
Nevirapine
Nialamide
Nicardipine
Nicocodine
Nicodicodine
Nicodicodine (6-nicotinoyldihydrocodeine)
Nicomorphine
Nicotinaldehyde Thio-semicarbazone
Nicotine - for human use (except in natural substances)
Nicoumalone
Nifedipine
Nifenazone
Niflumic Acid
Nifuratel
Nikethamide
Nilotinib
Nimetazepam
Nimorazole
Niridazole
Nitrazepam
Nitrofurantoin
Nitrofurazone
Nitroprusside Sodium
Nitroxoline
Nizatidine
Nomifensine
Noracymethadol
Noradrenaline
Norboletone
Norclostebol
Norcodeine
Nordazepam aka Nordiazepam
Norethandrolone
Norethindrone
Norethisterone
Norethynodrel
Norfloxacin
Norgestrel (d-Norgestrel)
Norlevorphanol
Normethadone
Normorphine
Norpipanone
Nortriptyline
Novobiocin
Nux Vomica Seed
Nux Vomica Tincture BP
Nystatin
Octacosactrin
Oestradiol
Oestriol
Oestrogenic substances, conjugated
Oestrone
Ofloxacin
Olanzepine
Oleandomycin
Olmesartan
Olopatadine - in preparations for local ophthalmic use
Omeprazole
Ondansetron
Opipramol
Opium, raw
Opium, Tincture BP
Oral Contraceptives - all
Orciprenaline
Orlistat
Orphenadrine
Orthocaine - in preparations for local ophthalmic use
Oseltamivir
Ouabain
Ovandrotone
Ovarian Gland, dried
Oxabolone
Oxacillin
Oxamniquine
Oxanamide
Oxandrolone
Oxantel Pamoate
Oxatomide
Oxazepam
Oxazolam
Oxcarbazepine
Oxedrine
Oxethazaine
Oxolinic Acid
Oxophenarsine
Oxprenolol
Oxtriphylline
Oxybuprocaine -
except in preparations for local ophthalmic use

Oxybutynin
Oxycodone
Oxymesterone
Oxymetholone
Oxymorphone
Oxypertine
Oxyphenbutazone
Oxyphenacylmine
Oxyphenonium Bromide
Oxytetracycline
Oxytetracycline Dihydrate
Oxytocins -
natural and synthetic

Paliperidone
Pancrelipase
Pancuronium Bromide
Pantoprazole
Papaverine
Papaverine -
in inhalers
Papaveroline
Pareidione Capsules
Paraldehyde
Paramethadione
Paramethasone Acetate
Parathyroid Gland
Paregoric B.P.
Pargyline
Paromomycin
Paroxetine
Pavaveroline 2-sulphonic Acid
Pazopanib
Pecilocin
Pemoline
Pempidine
Penemecillin
Penbutolol
Penethamate
Penicillamine
Penicillins - all
Pentacosactride
Pentaerythritol Tetranitrate
Pentazocine
Penthenenate bromide
Pentobarbitone
Pentolinium Tartrate
Pentosan Polysulfate Sodium
Pentoxifylline
Pentrium Tablets
Pergolide
Perhexiline
Pericyazine
Perindopril
Perphenazine
Pethidine
Phacetoperane
Phenacemide
Phenadoxone
Phenaglycodol
Phenampromide
Phenarsone Sulphoxylate
Phenazocine
Phenazine

except in preparations for local ophthalmic use
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Phenazine and Caffeine Citrate
Phenazine Salicylate
Phenbicillin Potassium
Phenbutrazate
Phencyclidine
Phendimetrazine
Phenazine Sulphate
Phenethicillin Potassium
Phenethyamine -

Pheneturide
Phenglutarimide
Phenindione
Pheniprazine
Phenmetrazine
Phenobarbitone
Phenol - for parenteral use
Phenomorphan
Phenoperidine
Phenoxybenzamine
Phenoxyethylpenicillin
Phenprocoumon
Phensuximide
Phentermine
Phentermine Resin Complex
Phentolamine
Phentoxate
Phenylaminosalicylate
Phenylbutazone
Phenylephrine - if for ophthalmic or nasal administration:
above 1%w/v
Phenyldimedone - and its derivatives
Phenylmethylbarbituric Acid
Phenylpropanolamine
Phenytoin
Phenytoin Sodium
Pholcodine - if for non-parenteral use and in undivided preparations with ms 2.5% (calculated as base) if for non-parenteral use and in single-dose preparations with ms per dosage unit 100 mg (calculated as base)

Phthalysulphacetamide
Phthalysulfathiazole
Physostigmine
Phytonadione
Phytonadione
Picrotoxin
Pilocarpine
Pimecrolimus
Phytomenadione
Phytonadione
Pimozide
Pinazepam
Pindolol
Pioglitazone
Pipamazin
Pipenzolate Bromide
Piperacetazine
Piperazine Oestrone Sulphate
Piperidolate
Piperilate
Pipobroman
Pipothiazine
Pipradol
Piracetam
Pirbuterol
Pirenzepine
Piretanide
Piritramide
Piroxicam
Pituitary extract
Pituitary Gland (whole dried) - if in inhalers
Pituitary powdered (posterior lobe) - if in inhalers
Pivampicillin
Pivmecillinam
Pizotifen
Podophyllotoxin
Podophyllum
Podophyllum Indian
Podophyllum Resin
Poldine Methylsulphate
Polidexide
Poliovaccines - all
Polymyxin B Sulphate - if for parenteral use
Polynoxylin
Polyoestradiol Phosphate
Polysaccharide Iron Complex
Polythiazide
Posaconazole
Potassium Aminosalicylate
Potassium Arsenite
Potassium Bromide
Potassium Chloride - if for non-parenteral medicinal use
Potassium Clorazepate
Potassium Gluconate
Potassium Hydroxyquinolone
Potassium Perchlorate
Practolol
Pralidoxime
Pramipexole
Prasterone
Pravastatin
Prazepam
Prazosin
Prednisolone
Prednisone
Pregabalin
Prenylamine Lactate
Prethcamide
Prilocaine - except in preparations for local ophthalmic use
Primaquine Phosphate
Primidone
Probenecid
Probucol
Procainamide
Procainamide Durules
Procaine - except in preparations for local ophthalmic use
Procaine Penicillin
Procarbazine
Prochlorperazine
Procyclidine
Prodilidine
Progesterone
Proguanil
Proheptazine
Prolactin
Proligestone
Prolintane
Promazine
Promethazine - if for parenteral use
Propafenone
Propanidid
Propantheline Bromide
Proparacaine - in preparations for oral, parenteral and ophthalmic use
Properidine
Propetandrol
Propicillin
Propiomazine
Propiram
Propranolol
Propylhexedrine - if in inhalers
Propylidone
Propylthiouracil
Propyphenzone
Proquamezine
Prostaglandins - all
Protamine
Proteline
Prothionamide
Prothipendyl
Protirelin
Protoveratrines A and B
Protriptyline
Proxymetacaine - except in preparations for local ophthalmic use
Proxyphylline
Pseudoephedrine
Psilocybin
Pyrantel
Pyrazinamide
Pyridostigmine Bromide
Pyrimethamine
Pyroglutamyl, L-histidyl-L-proline Amide, L
Pyrovalerone
Quetiapine
Quinagolide
Quinalbarbitone
Quinapril
Quinbolone
Quinestradiol
Quinestrol
Quinethazone
Quingestanol
Quinidine
Quinine
Quinine and Urea
Quinuronium Sulphate
Rabeprazole
Racemethorphan
Racemoramide
Racemorphan
Racephedrine
Ragwort
Raloxifene
Raltegravir
Ramipril
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Ranitidine
Rasagiline
Rauwolfia (Serpentina and Vomitoria)
Razoxane
Repaglinide
Reproterol
Rescinnamine
Reserpine
Retinol - for oral use in preparations containing more than 10,000 units per dosage unit if for parenteral use
Ribavirin
Rifamide
Rifampicin
Rifamycin
Riluzole
Rimexolone
Rimiterol
Risedronate
Risperidone
Ristocetin
Ritodrine
Ritonavir
Rivaroxaban
Rivastigmine
Rizatriptan
Rolitetracycline Nitrate
Ropinirole
Rosiglitazone
Rosuvastatin
Rotigotine
Roxibolone
Rufinamide
Sabadilla
Salazosulphadimidine
Salbutamol
Salcatonin
Salmefamol
Salmeterol
Salsalate
Sandostatin
Saquinavir
Saxagliptin
Secbutobarbitone
Selegiline
Sertraline
Sevelamer Carbonate
Silandrone
Sildenafil
Silver Nitrate - if for medicinal use
Silver Sulphadiazine
Simvastatin
Sirolimus
Sissomicin
Sitagliptin
Sodium Aminosalicylate
Sodium Antimonylgluconate
Sodium Apolate
Sodium Arsanilate
Sodium Arsenate
Sodium Arsenite
Sodium Aurothiomalate
Sodium Bromate
Sodium Bromide
Sodium Cacodylate
Sodium Cromoglycate
Sodium Ethacrynate
Sodium Fluoride
Sodium Fusidate
Sodium Iodide preparations - for internal use
Sodium Methylarsinate
Sodium Monofluorophosphate - no restriction if in dentifrices and ms 1.14%
Sodium Nitroprusside
Sodium Stibogluconate
Sodium Tauroglycocholate
Sodium Tetradecyl Sulphate
Sodium Valproate
Solapsone
Solifenacin
Somatotropin (Somatrophin)
Somatrem
Sorafenib
Sotalol
Spectinomycin
Spiramycin
Spironolactone
Stannous Fluoride
Stanolone
Stanozolol
Stavudine
Stenbolone
Stibocaptate
Stibophen
Stilboestrol
Streptodornase
Streptokinase
Streptomycin
Streptozocin
Strontium Bromide
Strophanthin-K
Strychnine
Styramate
Succinamide
Succinylsulphathiazole
Sucralfate
Sufentanil
Sulbutiamine
Sulfacytine
Sulfadieramide
Sulfadoxine
Sulfametopyrazine
Sulfamonomethoxine
Sulfapyrazole
Sulfasuxidine Tablets
Sulfoxone
Sulindac
Sulphabromomethazine
Sulphacetamide
Sulphachlorpyridazine
Sulphadiazine
Sulphadimethoxine
Sulphadimidine
Sulphaethidole
Sulphafurazole
Sulphafurazole Diethanolamine
Sulphaguanidine
Sulphafloxacin
Sulphamerazine
Sulphamethizole
Sulphamethoxazole
Sulphamethoxydiazine
Sulphamethoxypyridazine
Sulphamethylphenazole
Sulphamezathine preparations
Sulphamoprine
Sulphamoxole
Sulphanilamide
Sulphanitran
Sulphaphenazole
Sulphapyridine
Sulphaquinoxaline
Sulphasalazine
Sulphasomidine
Sulphathiazole
Sulphathiourea
Sulphatolamide
Sulphaurea
Sulphinpyrazone
Sulphomyxin
Sulphonamide
Sulpiride
Sulthiame
Sumatriptan
Sunitinib

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Suprofen
Sutilains - when sold or recommended as a debriding agent
Suxamethonium Bromide
Suxamethonium Chloride
Suxethonium Bromide
Tacrine
Tacrolimus
Tadalafil
Talampicillin
Tamoxifen
Tamsulosin
Tapentadol
Tazarotene
Teclothiazide Potassium
Telithromycin
Telmisartan
Temazepam
Temozolomide
Tenofovir
Terazosin
Terbutaline
Testosterone
Testosterone 17B Choral Hemiacetal
Tetrabenazine
Tetracaine - if for parenteral or ophthalmic use
Tetracosaactrin
Tetracycline
Tetracycline Phosphate Complex
Tetrasodium Fostestrol
Tetrazepam
Thalidomide
Thallium Acetate
Thebacon - and its salts
Thebaine
Theobromine
Theophylline
Thiambutosine
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Thiethylperazine
Thiocarlide
Thioguanine
Thiomesterone
Thiopropazate
Thioproperazine
Thioridazine
Thiosinamine
Thiosinamine and Ethyl Iodide
Thiotepa
Thiothixene
Thiouracil
Thrombin preparations
Thymoxamine
Thyroid
Thyrotrophin
Thyrotrophin releasing hormone
Thyroxine Sodium
Tiagabine
Tianulin
Tiaprofenic Acid
Ticarcillin
Tigloidine
Tilidate
Timolol
Tinidazole
Tioguanine/Thioguanine
Tiotropium
Tipranavir
Tizanidine
Tobramycin
Tocainide
Tofenacin
Tolazamide
Tolazoline
Tolbutamide
Tolcapone
Tolmetin Sodium Dihydrate
Tolperisone
Tolterodine
Topiramate
Torasemide
Toremifene
Totaquine
Tramadol
Trandolapril
Tranexamic Acid
Tranylcypromine Sulphate
Travoprost
Trazodone
Tretamine
Tretinoin
Triacyctyleoleandomycin
Triamcinolone
Triamcinolone Acetonide
Triamterene
Triaziquone
Triazolam
Tribenoside
Tribromoethyl Alcohol
Trichloroethylene - for inhalational purposes
Triclofos Sodium
Tricyclamol Chloride
Tridione preparations
Trienbolone Acetate
Trifluoperazine
Trifluperidol
Triflupromazine
Trifluridine
Trihexphenidyl
Triiodothyronine Injection
Triiodothyroprionic Acid
Trilostane
Trimeperidine
Trimeprazine
Trimetaphan
Trimetazidine
Trimethadione
<table>
<thead>
<tr>
<th>Name</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trimethoprim</td>
<td></td>
</tr>
<tr>
<td>Trimipramine</td>
<td></td>
</tr>
<tr>
<td>Trimustine</td>
<td></td>
</tr>
<tr>
<td>Trioxsalen</td>
<td></td>
</tr>
<tr>
<td>Tripsin</td>
<td>if for internal use</td>
</tr>
<tr>
<td>Trometamol</td>
<td></td>
</tr>
<tr>
<td>Tropicamide</td>
<td></td>
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<tr>
<td>Trospium</td>
<td></td>
</tr>
<tr>
<td>Troxidone</td>
<td></td>
</tr>
<tr>
<td>Tryptamine</td>
<td>Tryptamine or ring-hydroxy tryptamine derivatives formed by substitution at</td>
</tr>
<tr>
<td></td>
<td>the nitrogen atom of the sidechain with one or more alkyl substituents; their</td>
</tr>
<tr>
<td></td>
<td>salts; their esters and ethers; their salts (None of these derivatives</td>
</tr>
<tr>
<td></td>
<td>specified above is thought to be commercially available)</td>
</tr>
<tr>
<td>Tryptophan, L-</td>
<td></td>
</tr>
<tr>
<td>Trypure</td>
<td></td>
</tr>
<tr>
<td>Tubocurarine Chloride</td>
<td></td>
</tr>
<tr>
<td>Tybamate</td>
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<tr>
<td>Tylosin</td>
<td></td>
</tr>
<tr>
<td>Tylosin Phosphate</td>
<td></td>
</tr>
<tr>
<td>Tylosin Tartrate</td>
<td></td>
</tr>
<tr>
<td>Tyrothricin</td>
<td></td>
</tr>
<tr>
<td>Uramustine</td>
<td></td>
</tr>
<tr>
<td>Umeclidinium</td>
<td></td>
</tr>
<tr>
<td>Urea</td>
<td>if for medicinal use</td>
</tr>
<tr>
<td>Urea Stibamine</td>
<td></td>
</tr>
<tr>
<td>Uredofos</td>
<td></td>
</tr>
<tr>
<td>Urethane</td>
<td></td>
</tr>
<tr>
<td>Uridine-5-triphosphoric Acid</td>
<td></td>
</tr>
<tr>
<td>Urokinase</td>
<td></td>
</tr>
<tr>
<td>Ursodeoxycholic Acid</td>
<td></td>
</tr>
<tr>
<td>Vaccines</td>
<td>all</td>
</tr>
<tr>
<td>Valacyclovir</td>
<td></td>
</tr>
<tr>
<td>Valerian preparations</td>
<td></td>
</tr>
<tr>
<td>Valproic Acid</td>
<td></td>
</tr>
<tr>
<td>Valsartan</td>
<td></td>
</tr>
</tbody>
</table>
PHARMACY AND POISONS ACT 1979

Vancomycin
Vardenafil
Varenicline
Venlafaxine
Verapamil
Veratrine
Veratrum (green and white)
Vidarbine
Vigabatrin
Viloxazine
Vinbarbitone
Vinblastine
Vincristine
Vindesine
Viomycin
Virginiamycin
Vitamin B12 - with intrinsic Factor Concentrate
Vitamin D - above 50,000 I.U. per dosage unit
Voriconazole

Warfarin
Xantinol Nicotinate
Xipanide
Xylazine
Yohimbine

Zafirlukast
Zaleplon
Zeranol
Zidovudine
Zimelidine
Zipeprol
Zinc Sulphate - if for oral use with md greater than 200mg
Ziprasidone
Zolmitriptan
Zolpidem
Zonisamide
Zopiclone
1. 1-Ethyl-3-piperidyl-alpha-cyclopentyl mandelate
   1. Methyl-4-phenylpiperidine-4-carboxylic acid
   2. Methyl-3-morpholino-1,1-diphenylpropane-Levomethorphan carboxylic acid
   4. Chloromethandienone
   4. Cyano-1-methyl-4-Isomethadone phenylpiperidine
   4. Cyano-2-dimethylamino-4,4-Hydromorphone diphenylbutane
   4. Phenylpiperidine-4-carboxylic acid ethyl ester
   α-Methylphenethylhydroxylamine

2. Any ester or ether or substance for the time being specified in paragraph 1.

3. Any salt of a substance for the time being specified in paragraph 1 or 2.

FOURTH SCHEDULE

(Sections 28(1); 28(2))

PART I

DRUGS OBTAINABLE ONLY AT REGISTERED PHARMACIES

Note: The following annotations used in this Schedule have the following meanings:

<table>
<thead>
<tr>
<th>Annotation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>md</td>
<td>(maximum dose) i.e. the maximum quantity of the substance contained in the amount of a medicinal product which is recommended to be taken or administered at any one time.</td>
</tr>
<tr>
<td>mdd</td>
<td>(maximum daily dose) i.e. the maximum quantity of the substance that is contained in the amount of a medicinal product which is recommended to be taken or administered in any period of 24 hours.</td>
</tr>
<tr>
<td>mg</td>
<td>milligram i.e. either or, if so specified, both of the following:</td>
</tr>
<tr>
<td>ms</td>
<td>(maximum strength) (a) the maximum quantity of the substance by weight or volume that is contained in the dosage unit of a medicinal product; or (b) the maximum percentage of the substance contained in a medicinal product calculated in terms of w/w, w/v, v/w, or v/v, as appropriate.</td>
</tr>
<tr>
<td>external use</td>
<td>means for application to the skin, teeth, mucosa of the mouth, throat, nose, eye, ear, vagina or anal canal when a local action only is necessary and extensive systemic absorption is unlikely to occur. Note: the following are not regarded as for external use: throat sprays, throat pastilles, throat lozenges, throat tablets, nasal drops, nasal sprays, nasal inhalations or teething preparations.</td>
</tr>
<tr>
<td>parenteral use</td>
<td>means administration by breach of the skin or mucous membrane.</td>
</tr>
</tbody>
</table>

---

1. Acetomenaphthone - in preparations for external use and ms 0.2%
2. Acetylcholine - in preparations for external use (ms 1.3% of the crude drug)
3. Aconite root - in preparations for external use and ms 0.02%
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- Aconitine - Eye drops, neutral BPC in preparations for external use
- Acriflavine
- Adrenaline
- Albendazole
- Alkaline Eye Drops BPC
- Allantoin
- Aloprin
- Aluminium acetate - for medicinal use
- Aluminium chloride - alcoholic solutions
- Aluminium oxide - for human use
- Alverine
- Ambucetamide
- Amethocaine - in all other preparations for non-parenteral use
- Aminacrine - in preparations for non-parenteral use
- Amylocaine
- Antazoline - if for nasal or ophthalmic administration
- Aspirin - if enteric coated or formulated in any other way so as to delay absorption
- Azatadine
- Bamethan
- Belladonna herb - in preparations for external use, in preparations for internal use and mdd 1mg of the alkaloids
- Belladonna root - in preparations for external use, in preparations for internal use and mdd 1mg of the alkaloids
- Benorylate
- Benzamine lactate
- Benzocaine - if in preparations for non-parenteral use with ms more than 1%
- Benzoyl peroxide - in concentrations of 10% or less
- Benzydamine preparations
- Benzyl Benzoate preparations
- Betaine
- Borax BP
- Boric Acid BP
- Bromelains
- Bromodiphenhydramine
- Brompheniramine
- Buclizine
- Buclosamide
- Buphenine - in preparations for internal use with md 6mg and mdd 18mg
<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
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</thead>
<tbody>
<tr>
<td>Bupivacaine</td>
<td>in preparations for non-parenteral use</td>
</tr>
<tr>
<td>Butacaine</td>
<td>in preparations for non-parenteral use</td>
</tr>
<tr>
<td>Butanilicaine</td>
<td>in preparations for non-parenteral use</td>
</tr>
<tr>
<td>Butethamide</td>
<td></td>
</tr>
<tr>
<td>Butoxyethyl nicotinate</td>
<td></td>
</tr>
<tr>
<td>Butylaminobenzoate</td>
<td>for topical use only</td>
</tr>
<tr>
<td>Calcium</td>
<td>if for oral administration</td>
</tr>
<tr>
<td>Glucogalactogluconate</td>
<td></td>
</tr>
<tr>
<td>Calcium polystyrene sulphonate</td>
<td></td>
</tr>
<tr>
<td>Calcium resonium</td>
<td></td>
</tr>
<tr>
<td>Calcium with Vitamin D tablets</td>
<td></td>
</tr>
<tr>
<td>BPC</td>
<td></td>
</tr>
<tr>
<td>Cantharidin</td>
<td>in preparations for external use and ms 0.01%</td>
</tr>
<tr>
<td>Caramiphen</td>
<td>in liquid preparations and ms 0.1% (calculated as base), in tablet preparation and ms 7.5mg (calculated as base)</td>
</tr>
<tr>
<td>Carbaryl preparations</td>
<td></td>
</tr>
<tr>
<td>Carbenoxolone</td>
<td>in gels and ms 2%, in pellets with md 5mg and mdd 25mg</td>
</tr>
<tr>
<td>Carbetapentane citrate</td>
<td></td>
</tr>
<tr>
<td>Carinoxamine</td>
<td></td>
</tr>
<tr>
<td>Castor oil</td>
<td>if for ophthalmic use</td>
</tr>
<tr>
<td>Cetylpyridium chloride</td>
<td>if for internal use</td>
</tr>
<tr>
<td>Charcoal</td>
<td>if for internal use</td>
</tr>
<tr>
<td>Chloral hydrate</td>
<td>in preparations for external use</td>
</tr>
<tr>
<td>Chlorcyclizine</td>
<td></td>
</tr>
<tr>
<td>Chlordantoin</td>
<td></td>
</tr>
<tr>
<td>Chlorhexidine</td>
<td>if for administration into the nasal or oral cavities,</td>
</tr>
<tr>
<td>-</td>
<td>if for use specifically as a bath additive,</td>
</tr>
<tr>
<td>Chlorpheniramine</td>
<td>if impregnated onto gauze dressing for direct application to a wound</td>
</tr>
<tr>
<td>Chlorphenoxyethanol</td>
<td></td>
</tr>
<tr>
<td>Chlorprenaline</td>
<td></td>
</tr>
<tr>
<td>Chlorpyriline Citrate</td>
<td></td>
</tr>
<tr>
<td>Chlorxylenol</td>
<td>for application to the skin</td>
</tr>
<tr>
<td>Cholebrin tablets</td>
<td></td>
</tr>
<tr>
<td>Choline magnesium trisalicylate</td>
<td></td>
</tr>
<tr>
<td>Choline salicylate</td>
<td></td>
</tr>
<tr>
<td>Chymotrypsin</td>
<td></td>
</tr>
<tr>
<td>Cinchocaine</td>
<td>in preparations for non-parenteral use and ms 3% (calculated as base)</td>
</tr>
<tr>
<td>Cineole</td>
<td></td>
</tr>
<tr>
<td>Substance</td>
<td>Notes</td>
</tr>
<tr>
<td>----------------------------</td>
<td>----------------------------------------------------------------------</td>
</tr>
<tr>
<td>Cinnarizine</td>
<td>in preparations for external use, in preparations for internal use for treatment of mouth ulcers with ms 35mg and mdd 250mg</td>
</tr>
<tr>
<td>Clioquinol -</td>
<td></td>
</tr>
<tr>
<td>Coal tar -</td>
<td>in preparations Conium leaf in preparations for external use and ms 7% of the crude drug</td>
</tr>
<tr>
<td>Creosote -</td>
<td>if for medicinal use</td>
</tr>
<tr>
<td>Crotamiton</td>
<td></td>
</tr>
<tr>
<td>Cyanocobalamin -</td>
<td>if in a formulation in which it is the sole active ingredient and is for internal use</td>
</tr>
<tr>
<td>Cyclizine -</td>
<td>if in preparations 1% and less</td>
</tr>
<tr>
<td>Cyclomethacaine -</td>
<td>if for nasal administration</td>
</tr>
<tr>
<td>Cyclopentamine -</td>
<td>if for oral administration and maximum dose 15mg</td>
</tr>
<tr>
<td>Cyclopropane</td>
<td></td>
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<tr>
<td>Cyteal</td>
<td></td>
</tr>
<tr>
<td>Danthron</td>
<td></td>
</tr>
<tr>
<td>Deanol -</td>
<td>in preparations for internal use and mdd 26 mg</td>
</tr>
<tr>
<td>Dequalinium -</td>
<td>chloride in external paint preparations and ms 1% in throat lozenges or throat pastilles and ms 0.25mg</td>
</tr>
<tr>
<td>Dexamfetamine</td>
<td></td>
</tr>
<tr>
<td>Dextromethorphan -</td>
<td>in preparations for internal use with md 15 mg (calculated as base) and mdd 75 mg (calculated as base)</td>
</tr>
<tr>
<td>Di-iodohydroxyquinoline -</td>
<td>for topical preparations for the skin Diabetic Diagnostic Reagents all Diabetic diagnostic tests</td>
</tr>
<tr>
<td>Diatrizoate sodium -</td>
<td>for non-parenteral use</td>
</tr>
<tr>
<td>Dibromopropamidine -</td>
<td>for ophthalmic use</td>
</tr>
<tr>
<td>Dichlorphen</td>
<td></td>
</tr>
<tr>
<td>Dicophane</td>
<td></td>
</tr>
<tr>
<td>DiHydroxystearcoglycerol</td>
<td></td>
</tr>
<tr>
<td>Dimenhydrinate</td>
<td></td>
</tr>
<tr>
<td>Dimethindene</td>
<td></td>
</tr>
<tr>
<td>Dimethisoquin -</td>
<td>in preparations for non-parenteral use</td>
</tr>
<tr>
<td>Dimethyldimethanobenzal tartrate</td>
<td></td>
</tr>
<tr>
<td>Dioctylsodium sulphosuccinate</td>
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</tr>
<tr>
<td>Diphenhydramine</td>
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</tr>
<tr>
<td>Diphenylpyruline</td>
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<tr>
<td>Dithranol preparation</td>
<td></td>
</tr>
<tr>
<td>Docusate sodium</td>
<td></td>
</tr>
<tr>
<td>Domiphen Bromide -</td>
<td>if for oral use</td>
</tr>
<tr>
<td>Doxylamine</td>
<td></td>
</tr>
<tr>
<td>Medicine</td>
<td>Specification</td>
</tr>
<tr>
<td>---------------------------</td>
<td>-------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Embamine</td>
<td>in preparations for internal or external use and ms 1%</td>
</tr>
<tr>
<td>Emetine</td>
<td></td>
</tr>
<tr>
<td>Ephedrine</td>
<td>in nasal sprays or nasal drops and ms 2%, in preparations for external use, in preparations for internal use (except nasal sprays or nasal drops) with md 30 mg and mdd 60 mg</td>
</tr>
<tr>
<td>Ethyl alcohol</td>
<td>for medicinal use</td>
</tr>
<tr>
<td>Famotidine</td>
<td>in preparations for internal use with ms 10 mg</td>
</tr>
<tr>
<td>Ferrous salts</td>
<td>for internal use where the ferrous salt is the sole active ingredient</td>
</tr>
<tr>
<td>Fluorescein</td>
<td></td>
</tr>
<tr>
<td>Fluothane</td>
<td></td>
</tr>
<tr>
<td>Folic Acid</td>
<td>if in preparations for internal use and mdd 500 micrograms</td>
</tr>
<tr>
<td>Frangula preparations</td>
<td></td>
</tr>
<tr>
<td>Gamma Benzene hexachloride</td>
<td></td>
</tr>
<tr>
<td>Gelsemine</td>
<td>in preparations for internal or external use and ms 0.1%</td>
</tr>
<tr>
<td>Gelsemium</td>
<td>in preparations for internal use with md 25mg of the crude drug and mdd 75mg of the crude drug</td>
</tr>
<tr>
<td>Glutaraldehyde</td>
<td></td>
</tr>
<tr>
<td>Glycopyrronium bromide</td>
<td>in preparations for internal use with md 1 mg and mdd 2 mg</td>
</tr>
<tr>
<td>Grindelia liquid extract</td>
<td></td>
</tr>
<tr>
<td>Guaiacol</td>
<td></td>
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<tr>
<td>Guar gum</td>
<td></td>
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<tr>
<td>Gynomin</td>
<td></td>
</tr>
<tr>
<td>Halibut-Liver Oil Capsules</td>
<td></td>
</tr>
<tr>
<td>Heparin</td>
<td>in preparations for external use</td>
</tr>
<tr>
<td>Heparinoid</td>
<td></td>
</tr>
<tr>
<td>Hexachlorophane</td>
<td>if in preparations for external use in:</td>
</tr>
<tr>
<td></td>
<td>(a) soaps with ms more than 0.1% but not more than 2%</td>
</tr>
<tr>
<td></td>
<td>(b) products other than soaps or aerosols with ms more than 0.1% but not more than 0.75%</td>
</tr>
<tr>
<td>Hexamidine isethionate</td>
<td></td>
</tr>
<tr>
<td>Histapyrodine</td>
<td></td>
</tr>
<tr>
<td>Histidine, L</td>
<td>if for use as an ingredient in dietary or nutritional products as an aminoacid</td>
</tr>
<tr>
<td>Homatropine hydrobromide</td>
<td>if in preparations for internal use with md 0.2mg and mdd 0.6mg</td>
</tr>
<tr>
<td>Hydrargaphen</td>
<td>in preparations for local application to the skin</td>
</tr>
<tr>
<td>Hydroxymethylgramicidin</td>
<td>if in throat lozenges or throat pastilles</td>
</tr>
<tr>
<td>Compound</td>
<td>Usage and Dosage</td>
</tr>
<tr>
<td>--------------------------------</td>
<td>---------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Hyoscine</td>
<td>in preparations for external use, in preparations for internal use with ms 0.15%</td>
</tr>
<tr>
<td>Hyoscine butylbromide</td>
<td>in preparations for external use, in preparations for internal use (other than inhalers) with md 3mg and mdd 9mg; or</td>
</tr>
<tr>
<td>Hyoscine hydrobromide</td>
<td>in preparations for external use, in preparations for internal use with md 300 micrograms and mdd 900 micrograms</td>
</tr>
<tr>
<td>Hyoscine methobromide</td>
<td>in preparations for external use, in preparations for internal use with md 2.5 mg and mdd 7.5 mg</td>
</tr>
<tr>
<td>Hypermellose</td>
<td>for ophthalmic use</td>
</tr>
<tr>
<td>Inositol nicotinate</td>
<td></td>
</tr>
<tr>
<td>Iocetamic acid</td>
<td>if for oral administration</td>
</tr>
<tr>
<td>Iodinated glycerin</td>
<td></td>
</tr>
<tr>
<td>Ipecacuanha</td>
<td></td>
</tr>
<tr>
<td>Isopropamide iodide</td>
<td>in preparations for internal use with md 2.5 mg (as isopropamide ion) and mdd 5 mg (as isopropamide ion)</td>
</tr>
<tr>
<td>Isothipendyl</td>
<td></td>
</tr>
<tr>
<td>Ispagula husk</td>
<td></td>
</tr>
<tr>
<td>Jaborandi</td>
<td>in preparations for external use and ms more than 0.025% of the alkaloids in the medicinal product;</td>
</tr>
<tr>
<td>Kaolin Poultice BPC</td>
<td></td>
</tr>
<tr>
<td>Lachesine Eye Drops BPC</td>
<td></td>
</tr>
<tr>
<td>Lactulose</td>
<td></td>
</tr>
<tr>
<td>Lead Subacetate Solution,</td>
<td></td>
</tr>
<tr>
<td>Dilute BPC</td>
<td></td>
</tr>
<tr>
<td>Lead Subacetate Solution,</td>
<td></td>
</tr>
<tr>
<td>Strong BPC</td>
<td></td>
</tr>
<tr>
<td>Lignocaine</td>
<td>in preparations for external use and ms 0.7% in preparations for non-parenteral use</td>
</tr>
<tr>
<td>Lindane</td>
<td></td>
</tr>
<tr>
<td>Lithium carbonate</td>
<td>in preparations for internal use with md 5mg (calculated as base) and mdd 15 mg (calculated as base)</td>
</tr>
<tr>
<td>Lithium sulphate</td>
<td>in preparations for internal use with md 5mg (calculated as base) and mdd 15 mg (calculated as base)</td>
</tr>
<tr>
<td>Lobeline</td>
<td>in preparations for external use, in preparations for internal use with md 3mg and mdd 9mg (calculated as base)</td>
</tr>
<tr>
<td>Loratadine</td>
<td>in tablets with ms 10 mg in syrup with ms 5 mg/5 ml</td>
</tr>
</tbody>
</table>
### PHARMACY AND POISONS ACT 1979

<table>
<thead>
<tr>
<th>Substance</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mafenide</td>
<td>in eye drops and ms 5%</td>
</tr>
<tr>
<td>Magnesium citrate</td>
<td></td>
</tr>
<tr>
<td>Malathion preparations</td>
<td></td>
</tr>
<tr>
<td>Mebeverine</td>
<td>if in preparations for internal use with md 100mg and mdd 300mg</td>
</tr>
<tr>
<td>Meclozine</td>
<td></td>
</tr>
<tr>
<td>Medicinal opium</td>
<td>in liquid preparations with ms 0.02% (calculated as anhydrous morphine base)</td>
</tr>
<tr>
<td>-</td>
<td>in solid preparations with ms 0.04% (calculated as anhydrous morphine base)</td>
</tr>
<tr>
<td>Menadiol</td>
<td>for internal use excluding parenteral route</td>
</tr>
<tr>
<td>Mepenzolate bromide</td>
<td>in preparations for internal use with md 25mg and mdd 75mg</td>
</tr>
<tr>
<td>Mepivacaine</td>
<td>in preparations for non-parenteral use</td>
</tr>
<tr>
<td>Mepyramine</td>
<td>if for non-parenteral use Mercuric oxide if for human use</td>
</tr>
<tr>
<td>MetabutethamineB</td>
<td>in preparations for non-parenteral use</td>
</tr>
<tr>
<td>Methapyrilene</td>
<td></td>
</tr>
<tr>
<td>Methoxamine</td>
<td>in nasal sprays or nasal drops not containing liquid paraffin as a vehicle and ms 0.25%</td>
</tr>
<tr>
<td>Methylephedrine</td>
<td>in preparations for internal use with md 30mg and mdd 60mg</td>
</tr>
<tr>
<td>Methylhydroxybenzoate</td>
<td></td>
</tr>
<tr>
<td>Miristalkonium chloride</td>
<td></td>
</tr>
<tr>
<td>Monosulfiram</td>
<td>for external use</td>
</tr>
<tr>
<td>Naphazoline</td>
<td>in nasal sprays or nasal drops not containing liquid paraffin as a vehicle and ms 0.05%</td>
</tr>
<tr>
<td>-</td>
<td>if in eye drops and ms 0.25%</td>
</tr>
<tr>
<td>Natuderm Cream</td>
<td></td>
</tr>
<tr>
<td>Niclosamide</td>
<td></td>
</tr>
<tr>
<td>Nicotinic Acid</td>
<td>for internal use</td>
</tr>
<tr>
<td>Nicotinyl alcohol</td>
<td>for internal use</td>
</tr>
<tr>
<td>Nizatidine</td>
<td>in preparations for internal use with ms 75 mg</td>
</tr>
<tr>
<td>Orthocaine</td>
<td>if in preparations for non-parenteral use</td>
</tr>
<tr>
<td>Oxolamine</td>
<td></td>
</tr>
<tr>
<td>Oxybuprocaine</td>
<td>if in preparations for non-parenteral use</td>
</tr>
<tr>
<td>Oxymetazoline</td>
<td></td>
</tr>
<tr>
<td>Oxyphenonium bromide</td>
<td>in preparations for internal use with md 5mg and mdd 15mg</td>
</tr>
</tbody>
</table>
PHARMACY AND POISONS ACT 1979

- Padimate
- Pancreatin
- Papaverine - in preparations for internal use with md 50mg (calculated as base) and mdd 150mg (calculated as base)
- Penthienate methobromide - in preparations for internal use with md 5mg and mdd 15mg
- Penthrane - if in preparations for non-parenteral use
- Phenacaine
- Phenazine - in preparations for external use
- Phenindamine
- Pheniramine
- Phenol - for all medicinal use
- Phenylephrine - if for internal use (excluding parenteral route) with md 20mg and mdd 40mg
  if for ophthalmic or nasal administration; with a maximum strength of 1%w/v
- Phenylpropanolamine - in nasal sprays or nasal drops and ms 2% in preparations for internal use (except controlled release capsules, nasal sprays or nasal drops) with md 50mg and mdd 150mg
- Phenyltoloxamine
- Pholcodine - if for non-parenteral use and in undivided preparations with ms 1.5% (calculated as base) and md 20 mg (calculated as base)
  if for non-parenteral use and in single-dose preparations with ms per dosage unit 1.5% (calculated as base) and md 20 mg (calculated as base)
- Phosphorylcolamine
- Phytomenadione - if for non-parenteral use in preparations for internal use with md 5mg and mdd 15mg
- Pipenzolate bromide - in preparations for internal use with md 5mg and mdd 15mg Piperazine
- Piperidolate - if in preparations for internal use with md 50mg and mdd 150mg
- Podophyllum resin - in preparations for external use and ms 20%
- Poldine methylsulphate - in preparations for internal use with md 2mg and mdd 6mg Polvinyl alcohol if for ophthalmic use
- Polystyrene sulphonate resins - for use as an enema
- Ponoxylan
- Potassium arsenite - if in preparations for internal or external use and ms 0.0127%
- Potassium citrate preparations
Potassium guaicol sulphonate
Povidone iodine preparations - all
Pramoxine
Prilocaine - if in preparations for non-parenteral use
Procaine - if in preparations for non-parenteral use
Promethazine
Propantheline bromide - if in preparations for internal use with md 15 mg and mdd 45mg
Propamidine
Proxamine
Proxymetacaine - if in preparations for non-parenteral use
Pseudoephedrine - if in preparations for internal use with md 60mg and mdd 180mg
Pumilio pine oil
Pyrrobutamine phosphate
Quinine - in preparations for internal use with md 100mg and mdd 300mg (calculated as base)
Racephedrine - in nasal sprays or nasal drops and ms 2% in preparations for external use
Resonium A - in preparations for internal use (except nasal sprays or nasal drops) with md 30mg and mdd 60mg
Resorcinol preparations - if for medicinal use
Retinol - in preparations containing 10,000 units or less
Rose Bengal - if for ophthalmic use
Salicylamide
Salicylic Acid - if for medicinal use
Scarlet Red Ointment
Selenium sulphide
Senna
Sodium alkylsulphoacetate - if for rectal administration
Sodium apolate - if in preparations for external use
Sodium arsenite - if in preparations for internal or external use and ms 0.013%
Sodium Cellulose phosphate - if for internal use
Sodium cromoglycate - if in preparations for use by being administered through the nose
Sodium fluoride - in preparations for use in the prevention of dental caries, other than dentifrices, in the form of: tablets or drops and
Sodium ipodate capsules
Sodium iron edetate
Sodium Perborate - in preparations for oral use
Sodium picosulphate
Sodium pidolate
Squalane
Squill preparations - for human use
Sterculia preparations
Streptodornase - if in preparations for external use
Streptokinase - if in preparations for external use
Succinamide - in products for decontaminating water
Terpin hydrate - if for medicinal use
Tetracaine
Tetrahydrofurfuryl salicylate
Tetrahydrozoline
Thiabendazole
Thiomersal - when used as a skin antiseptic
Tolazoline - if in preparations for external use
Totaquine - if in preparations for internal use with md 100mg and mdd 300mg
Tramazoline
Tripelennamine
Triprolidine
Tripotassium dicitratobismuthate
Trypsin - if for external use
Tryptophan, L- if used as an ingredient in dietary or nutritional products as an essential amino-acid; or in preparations for external use
Turpentine oil - if for internal use
Tyloxapol Tyrothricin - if in throat lozenges or throat pastilles
Urea - if for application to the skin
Urea hydrogen peroxide - if for aural use
Vanillylnonamide
Viprynium
Vitamin D 1000-50,000 I.U. per dosage unit
Xylometazoline

Zinc Sulphate - if for oral use
Zinc Sulphate and Adrenaline Eye Drops
Zinc Sulphate Eye Drops BPC

2 Any ester or ether or substance for the time being specified in paragraph 1.
3 Any salt of a substance for the time being specified in paragraph 1 or 2.

PART II

DRUGS OBTAINABLE ONLY FROM REGISTERED PHARMACISTS AT REGISTERED PHARMACIES

1
Acyclovir - in preparations for topical use and ms 5%
Adrenaline
Ammonium Chloride and Morphine Mixture BP
Astemizole

Bacitracin - in topical preparations for auricular or local ophthalmic use
Cannabidiol with tetrahydrocannabinol less than 1%
Chloroform - except for inhalational use
Clemestine
Clotrimazole - if in preparations for external or vaginal use
Cyproheptadine

Diclofenac - in topical preparations
Econazole - in cream, powder or solution for external use if in preparations for external vaginal use
Enflurane - except for inhalational use
Ether - except for inhalational use Ethyl alcohol for external use
Ethyl alcohol - for external use
<table>
<thead>
<tr>
<th>Compound</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ethylmorphine</td>
<td>in undivided preparations with ms 0.2% (calculated as base) and with md 7.5mg (calculated as base); or in single dose preparations with ms per dosage unit 0.2% (calculated as base) and 7.5mg (calculated as base)</td>
</tr>
<tr>
<td>Folic acid</td>
<td>if in preparations for internal use and md 500 micrograms and mdd 1000 micrograms</td>
</tr>
<tr>
<td>Gramicidin</td>
<td>in preparations for external use and ms 0.02% in topical preparations for auricular or local ophthalmic use</td>
</tr>
<tr>
<td>Haloprogin</td>
<td>in preparations for external use Halothane except for inhalational use</td>
</tr>
<tr>
<td>Homatropine</td>
<td>in preparations for external use, in preparations for internal use with md 0.15mg and mdd 0.45mg</td>
</tr>
<tr>
<td>Homatropine methylbromide</td>
<td>in preparations for internal use with md 2mg and mdd 6mg</td>
</tr>
<tr>
<td>Hyaluronidase</td>
<td>if in preparations for external use Hydrocortisone in preparations for topical use ms 0.5% in preparations for external use and ms 1%</td>
</tr>
<tr>
<td>Hydrocortisone</td>
<td>in preparations for topical use ms 0.5% in preparations for external use and ms 1%</td>
</tr>
<tr>
<td>Hydrogen cyanide</td>
<td>in preparations for internal or external use and ms 0.1%</td>
</tr>
<tr>
<td>Ibuprofen</td>
<td>if for use in rheumatic and muscular pain, backache, neuralgia, migraine, headache, dental pain, dysmenorrhoea, feverishness, symptoms of colds and influenza and with md 400mg and mdd 1200mg</td>
</tr>
<tr>
<td>Injections</td>
<td>except insulin products</td>
</tr>
<tr>
<td>Insulin</td>
<td>all</td>
</tr>
<tr>
<td>Iodine, aqueous solution</td>
<td>for internal use</td>
</tr>
<tr>
<td>Isoconazole</td>
<td>if in preparations for external or vaginal use</td>
</tr>
<tr>
<td>Isoconazole preparations</td>
<td>if for application to the skin (excluding mucous membranes)</td>
</tr>
<tr>
<td>Isoflurane</td>
<td>except for inhalational use</td>
</tr>
<tr>
<td>Itraconazole</td>
<td>if in preparations for external or vaginal use</td>
</tr>
<tr>
<td>Ketoconazole</td>
<td>if in preparations for external or vaginal use</td>
</tr>
<tr>
<td>Ketoprofen</td>
<td>in preparations for internal use with ms 75 mg</td>
</tr>
<tr>
<td>Levamizole</td>
<td></td>
</tr>
<tr>
<td>Levonorgestrel</td>
<td>if for use as an emergency contraception</td>
</tr>
<tr>
<td>Loperamide</td>
<td>if for the treatment of acute diarrhoea</td>
</tr>
</tbody>
</table>
Mebendazole - for external application to the skin, or for oral use or vaginal use
Miconazole - in topical preparations with ms 2%
Minoxidil - in liquid preparations with ms 0.02% (calculated as anhydrous morphine base) and md 3 mg (calculated as anhydrous morphine base) in solid preparations with ms 0.04% and 300 micrograms per dosage unit (calculated as anhydrous morphine base) with md 3mg (calculated as anhydrous morphine base)
Morphine - in liquid preparations with ms 0.02% (calculated as anhydrous morphine base) and md 3 mg (calculated as anhydrous morphine base) in solid preparations with ms 0.04% and 300 micrograms per dosage unit (calculated as anhydrous morphine base) with md 3mg (calculated as anhydrous morphine base)
Neomycin - in preparations for external use with ms 3.5 mg per gram
Nicotine - in oral preparations and md 2mg
Nicotine - if in liquid form for inhalation via any electronic delivery system with ms 21mg
Nicotine - in patches with ms 21 mg/24 hours
Nizatidine - in capsules with ms 75 mg
Phenazopyridine
Phenylpropanolamine - in controlled release capsules with md 75 mg and mdd 150mg
Polymyxin B sulphate with ms 75 mg
Ranitidine - in preparations for use on the skin
Silver Nitrate - Eye drops and ms 2% Eye ointment and ms 4%
Sodium cromoglycate - Insulin
Syringes -
Terfenadine for other use
Trichloroethylene -

2 Any ester or ether or substance for the time being specified in paragraph 1.

3 Any salt of a substance for the time being specified in paragraph 1 or 2.

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FIFTH SCHEDULE

PHARMACY PROFESSION COMPLAINTS COMMITTEE

1 The Committee shall consist of three members appointed by the Minister, as follows—
   (a) one from a list of at least three registered pharmacists in good standing who is nominated by the Association;
   (b) one from a list of at least three registered pharmacists in good standing nominated by the Council;
   (c) one professionally qualified person who is not a registered pharmacist.

2 A person who is a member of the Council may not be appointed as a member of the Committee.

3 Appointment as a member under paragraph 1 shall be for a term not exceeding three years and a member is eligible for reappointment.

4 The Minister may appoint a second person to act as alternate to a member appointed under paragraph 1.

5 An alternate member shall be appointed in accordance with the requirements for the appointment of the member, and his term of appointment shall, if not sooner terminated, end at the expiration of the term of the member.

6 There shall be a Chairman of the Committee who shall be appointed annually by the Minister from among the members of the Committee to hold office until 31 December of the year for which he was appointed, and who shall be eligible for reappointment as Chairman.

7 If at any time the Chairman ceases to be a member of the Committee, or for any other reason ceases to be the Chairman, the Minister shall, as soon as may be, appoint from among the members of the Committee another person to be Chairman in his stead.

8 Three members of the Committee shall form a quorum at any meeting.

9 (1) Where any complaint is before the Committee, a member of the Committee shall advise the Chairman if he is personally acquainted with the facts of the case and may, with leave of the Chairman, withdraw on that ground or for any other reason which the Chairperson deems sufficient; and the Chairman may himself withdraw on any such ground.

   (2) Where a member has so withdrawn, the Chairman may request the Minister to appoint a member of equal standing as the withdrawn member to be a member of the Committee for the purpose of those proceedings, and the Minister may make such appointment, whereupon the person so appointed shall be deemed to be a member of the Committee for such purpose.

10 Fees shall be paid to members of the Committee in accordance with the Government Authorities Fees Act 1971.
PHARMACY AND POISONS ACT 1979

11 The validity of any act or proceedings of the Committee shall not be affected by any vacancy among the members of the Committee or by any defect in the appointment of a member of the Committee or of the Chairperson.

12 The Committee shall, not later than 31 January after the end of each calendar year, submit a report on its activities for the preceding year to the Council.

13 Subject to this Act, the Committee shall regulate its own proceedings.

[Fifth Schedule repealed and replaced by 2013 : 48 s. 28 effective 24 December 2013]
PHARMACY AND POISONS ACT 1979

SIXTH SCHEDULE

(Section 26B)

PROHIBITION ORDERS, PROHIBITION NOTICES AND NOTICES TO WARN

PART I

PROHIBITION ORDERS

1 If the Minister proposes to make a prohibition order ("an order"), then, subject to paragraph 5, he shall before he makes the order—

(a) publish, in such manner as he thinks fit a notice stating—

(i) that he proposes to make the order and, in such terms as he thinks fit, the proposed effect of the order; and

(ii) that any person may make representations in writing to the Minister about the proposed order before a date specified in the notice (which must be after the expiration of the period of 28 days beginning with the date of the first publication of the notice); and

(b) consider any such representations made within that period.

2 The effect of an order must not be more restrictive, but may be less restrictive, than the proposed effect of it as stated in the notice.

3 Without prejudice to the power to make a further order and subject to paragraph 4, an order shall cease to have effect at the expiration of a period specified in the order which must not be longer than 12 months beginning with the date on which the order comes into force.

4 An order may revoke a previous order or may vary it otherwise than providing for it to be in force after expiration of 12 months beginning with the date of the coming into force of the previous order.

5 Paragraphs 1 and 2 shall not apply to an order if the order contains a statement that in the opinion of the Minister the risk of danger connected with the drug or drug product to which the order relates is such that the order must be made without delay.

PART II

PROHIBITION NOTICES

Preliminary

6 In this Part—

“notice” means a prohibition notice;
“notification” means a notification in writing;
“the trader” in relation to a proposed notice or an actual notice means the person 
on whom the proposed notice is proposed to be served or on whom the actual 
notice has been served.

7 A notice must specify the date on which it comes into force.

**General Procedure**

8 If the Minister proposes to serve a notice in respect of any drug or drug product, 
then, subject to paragraph 14, he shall before he serves the notice serve on the trader a 
notification—

(a) stating that the Minister proposes to serve on him a notice in respect of the 
drug or drug product; and

(b) specifying the drug or drug product in a manner sufficient to identify them 
and stating that, for the reasons set out in the notification, the Minister 
considers that the drug or drug products are not safe; and

(c) stating that the trader may make representations, in writing or both in 
writing and orally, for the purpose of satisfying the Minister that the drug 
or drug product is safe but that if the trader intends to make such 
representations he must, before the expiration of the period of 14 days 
beginning with the day when the notification is served on him, inform the 
Minister of his intention indicating whether the representations are to be 
in writing only or both in writing and oral.

9 Subject to paragraph 14, the Minister shall not serve a notice on the trader in 
respect of any drug or drug product before the expiration of the period of 14 days beginning 
with the day on which the Minister served on him a notification in pursuance of paragraph 
8 relating to the drug or drug products; and if within that period the trader informs the 
Minister as mentioned in paragraph 8(c), then—

(a) the Minister shall not serve a notice on the trader in consequence of the 
notification before the expiration of the period of 28 days beginning with 
the day aforesaid; and

(b) if during that period the trader makes to the Minister such written 
representations as are mentioned in paragraph 8(c) the Minister shall not 
serve a notice on the trader in consequence of the notification before the 
Minister has considered the report of a person appointed in pursuance of 
paragraph 10 in consequence of the representations.

10 Where, in consequence of the service on the trader of a notification in pursuance 
of paragraph 8, the trader informs the Minister as mentioned in paragraph 8(c) within the 
period so mentioned and makes to the Minister within that period or the fourteen days
beginning with the end of that period such written representations as are so mentioned, the
Minister shall—

(a) appoint any person to consider the written representations; and

(b) if the trader informed the Minister in pursuance of paragraph 8(c) that the
representations would be both written and oral, inform the trader of the
place and time (which must not be before the expiration of the fourteen
days and of seven days beginning with the day when the information is
given to the trader) at which the oral representations may be made to the
person appointed;

and the trader or his representative may at that place and time make to the person
appointed oral representations for the purpose of satisfying the Minister that the drug or
drug product in question is safe and may call and examine witnesses in connection with
the representations.

11 The person appointed in pursuance of paragraph 10 to consider written
representations with respect to any drug or drug product shall, after considering the
representations, any oral representations made in pursuance of that paragraph with
respect to the drug or drug product and any statements made by witnesses in connection
with the oral representations, make a report (including recommendations) to the Minister
about the representations and the proposed notice.

12 If at any time after the Minister has served a notification on the trader in pursuance
of paragraph 8 the Minister decides not to serve a notice on him in consequence of the
notification, the Minister shall inform him of the decision; and after the Minister informs
him of the decision the notification and anything done in consequence of it in pursuance of
the preceding paragraphs shall be disregarded for the purposes of those paragraphs.

13 Where a notification is served on the trader in respect of any drug or drug product
in pursuance of paragraph 8, a notice served on him in consequence of the notification may
relate to some only of those the drug or drug product.

Special Procedure

14 Paragraphs 8 to 13 do not apply to a notice which contains a statement that the
Minister considers that the risk of danger connected with the drug or drug product to which
the notice relates is such that the notice must come into force without delay; and references
to a notice in paragraphs 15 to 18 are to a notice containing such a statement.

15 A notice in respect of any drug or drug product must—

(a) state that, for the reasons set out in the notice, the Minister considers that
the drug or drug product is not safe; and

(b) state that the trader may, at such time as the trader thinks fit, make
representations in writing to the Minister for the purpose of satisfying him
that the drug or drug product is safe.
If representations in writing about a notice are made by the trader to the Minister, the Minister shall consider the representations and either revoke the notice and inform the trader that he has revoked it or—

(a) appoint a person to consider the representations; and

(b) serve on the trader a notification stating that he may make to the person appointed oral representations for the purpose mentioned in paragraph 15 and specifying the place and time (which, except with the agreement of the trader, must not be before the date of service of the notification) at which the oral representations may be made,

and the trader or his representative may at that place and time make to the person appointed oral representations and may call and examine witnesses in connection with the representations.

The person appointed in pursuance of paragraph 16 to consider written representations with respect to any drug or drug product shall, after considering the representations, any oral representations made in pursuance of that paragraph with respect to the goods and any statements made by witnesses in connection with the oral representations, make a report including recommendations to the Minister about the representations and the notice in question.

Where the Minister has appointed a person in pursuance of paragraph 16 to consider any representations relating to a notice then, without prejudice to the operation of paragraphs 19 and 20, paragraphs 16 and 17 shall not apply to any subsequent representations in writing about the notice.

Other Representations

If at any time the trader on whom a notice has been served makes representations in writing to the Minister for the purpose of satisfying him that the drug or drug product to which the notice relates is safe and, by virtue of paragraph 18, paragraph 16 does not apply to the representations, the Minister shall consider the representations and serve on the trader, before the expiration of one month beginning with the day when the Minister receives the representations, a notification stating—

(a) that the Minister will revoke the notice or vary it or declines to do so; or

(b) that the Minister has appointed a person to consider the representations and that the trader may make to the person appointed, at a place specified in the notification and a time so specified (which, except with the agreement of the trader, must not be before the expiration of the period of twenty-one days beginning with the date of service of the notification), oral representations for the purpose,

and the trader or his representative may at that place and time make to the person appointed oral representations and may call and examine witnesses in connection with the representations.
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20 The person appointed in pursuance of paragraph 19 to consider written representations with respect to any drug or drug product shall, after considering the representations, any oral representations made in pursuance of that paragraph with respect to the drug or drug product and any statements made by witnesses in connection with the oral representations, make a report including recommendations to the Minister about the representations and the notice in question.

Miscellaneous

21 The Minister may revoke or vary a notice by serving on the trader a notification stating that the notice is revoked or, as the case may be, is varied as specified in the notification; but the Minister shall not have power to vary a notice so as to make the effect of the notice more restrictive for the trader.

22 The Minister shall consider any report made to him in pursuance of paragraph 17 or 20 and, after considering the report, inform the trader of his decision with respect to the notice in question.

PART III
NOTICES TO WARN

23 If the Minister proposes to serve on a person a notice to warn in respect of any drug or drug product, the Minister shall, before he serves the notice, serve on the person a notification in writing—

(a) containing a draft of the notice and stating that the Minister proposes to serve on the person such a notice in the form of the draft;

(b) stating that, for the reasons set out in the notification, the Minister considers that the drug or drug product specified in the draft is not safe; and

(c) stating that the person may make representations, in writing or both in writing and orally, for the purpose of satisfying the Minister that the drug or drug product is safe but that if the person intends to make such representations he must, before the expiration of the period of fourteen days beginning with the day when the notification is served on him, inform the Minister of his intentions indicating whether the representations are to be in writing only or both in writing and oral.

24 Paragraphs 9 to 13 and 21 shall with the necessary modifications have effect in relation to a notice to warn as they have effect in relation to a prohibition notice but as if—

(a) the reference to paragraph 14 in paragraph 9 were omitted;

(b) for the references to paragraph 8 in paragraphs 9, 10, 12 and 13 there were substituted references to paragraph 23;
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(c) in paragraph 13 for the words from “relate” onwards there were substituted the words “be less onerous than the draft of the notice contained in the notification”; and

(d) in paragraph 21 the words “or vary” and the words from “or, as” onwards were omitted.

[Sixth Schedule inserted by 2013 : 48 s. 29 effective 24 December 2013]

[Assent Date: 23 July 1979]

[This Act was brought into operation on 1 January 1980]

[Amended by:

BR 62 / 1980
BR 63 / 1980
BR 64 / 1980
BR 16 / 1984
BR 17 / 1984
1984 : 46
1989 : 56
BR 21 / 1992
BR 42 / 1998
2008 : 20
2011 : 31
BR 35 / 2013
2013 : 48
2014 : 36
BR 69 / 2016
BR 111 / 2017
BR 113 / 2017]