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SCHEDULES
AN ORDINANCE

Relating to Poisons and Narcotic Drugs


Dated this thirteenth day of December 1978.

ZELMAN COWEN
Governor-General

By His Excellency's Command,

RALPH J. HUNT
Minister of State for Health

POISONS AND NARCOTIC DRUGS ORDINANCE 1978

PART I—PRELIMINARY

1. This Ordinance may be cited as the Poisons and Narcotic Drugs Ordinance 1978.*

2. This Ordinance shall come into operation on such date as is fixed by the Minister of State for the Capital Territory by notice in the Gazette.

3. (1) In this Ordinance, unless the contrary intention appears—

"analyst" means a person holding office by virtue of an appointment under section 51;

"cannabis" means a cannabis plant, whether living or dead, and includes, in any form, any flowering or fruiting tops, leaves, seeds, stalks or any other part of a cannabis plant or cannabis plants and any mixture of parts of a cannabis plant or cannabis plants, but does not include cannabis resin or cannabis fibre;

"cannabis fibre" means goods that consist wholly or substantially of fibre obtained from a cannabis plant or cannabis plants but do not contain any other substance or thing obtained from a cannabis plant;

"cannabis plant" means a plant of the genus Cannabis;

* Notified in the Commonwealth of Australia Gazette on 19 December 1978.
“cannabis resin” means a substance that consists wholly or substantially of resin (whether crude, purified or in any other form) obtained from a cannabis plant or cannabis plants;

“Chairman” has the same meaning as in the Health Commission Ordinance 1975;

“Committee” means the Drugs Advisory Committee established by section 17;

“dentist” means a person registered or licensed under the Dentists Registration Ordinance 1931 or a law of a State or another Territory that provides for the registration or licensing of dentists;

“Medical Officer of Health” has the same meaning as in the Public Health Ordinance 1928;

“medical practitioner” means

(a) a person registered under the Medical Practitioners Registration Ordinance 1930; or

(b) a person registered or licensed under a law of a State or another Territory that provides for the registration or licensing of medical practitioners;

“pharmacist” means a person registered under the Pharmacy Ordinance 1931;

“veterinary surgeon” means a person registered or licensed under the Veterinary Surgeons Registration Ordinance 1965 or a law of a State or another Territory that provides for the registration or licensing of veterinary surgeons;

“scheduled substance” means a substance that is specified in a schedule to this Ordinance;

“supply” includes sell and sale.

(2) In this Ordinance, a reference that consists of a reference to a schedule to this Ordinance followed immediately by the word “substance” is a reference to a substance specified in that schedule.

(3) For the purposes of sub-sections (1) and (2) and the schedules to this Ordinance, where a substance is specified in a schedule to this Ordinance, there shall, unless the contrary intention appears, be taken to be specified in that schedule—

(a) every salt, active principle or derivative of the substance and every salt of such an active principle or derivative;

(b) except where the substance is opium, every alkaloid of the substance and every salt of such an alkaloid;

(c) except where the substance is levomethorphan or levorphanol—every stereoisomer of the substance and every salt of such a stereoisomer; and
(d) in the case of a substance specified in Schedule 8 or Schedule 12—every ester and ether of the substance and every salt of such an ester or ether.

(4) A preparation or admixture that contains a substance that is specified, or is to be taken to be specified, in a schedule to this Ordinance, other than Schedule 9, shall, unless the contrary intention appears, be taken to be a substance specified in that schedule.

(5) A person is an addict for the purposes of Part II if—

(a) as a result of the repeated administration of a Schedule 8 substance, the person is subject to an overpowering desire for the continued administration of that substance; and

(b) the cessation of the administration of that substance to the person is likely to lead to definite symptoms of mental or physical distress or disorder.

(6) A reference in this Ordinance to the prescribed traffickable quantity for a Schedule 8 substance or a Schedule 12 substance is a reference to a quantity of the substance having a mass equal to the mass specified in Column 2 of Schedule 9 opposite to the reference to that substance in Column 1 of Schedule 9.

PART II—ADDICTIVE SUBSTANCES
Division 1—Unauthorized Use, Possession and Supply

4. (1) In this section, "controlled substance" means a substance that is a Schedule 8 substance or a Schedule 12 substance.

(2) A person who supplies a controlled substance to another person is guilty of an offence.

(3) A person who has a controlled substance in his possession for the purpose of supplying the substance to another person or to other persons is guilty of an offence.

(4) In proceedings for an offence against sub-section (3), a person who has in his possession a quantity of a controlled substance, being a quantity that exceeds the prescribed traffickable quantity for that substance, shall, unless the contrary is proved or the person proves that he had lawful authority to have the substance in his possession, be taken to have the substance in his possession for the purpose of supplying the substance to another person or to other persons.

(5) An offence against sub-section (2) or (3) is punishable upon indictment.
(6) A person who commits an offence against sub-section (2) or (3) is punishable, on conviction—

(a) if the offence is committed in relation to a substance other than cannabis—by imprisonment for a term not exceeding 25 years or a fine not exceeding $100,000 or both such imprisonment and fine; and

(b) if the offence is committed in relation to cannabis—by imprisonment for a term not exceeding 10 years or a fine not exceeding $4,000 or both such imprisonment and fine.

(7) Where, on the trial of a person for an offence against sub-section (2) or (3), the jury is not satisfied that the person is guilty of the offence charged but is satisfied that he is guilty of an offence against sub-section 5 (1), the jury may find the person not guilty of the offence charged but guilty of an offence against sub-section 5 (1), and the person is liable to be dealt with by the Supreme Court accordingly.

(8) Where, in pursuance of sub-section (7), a person is convicted of an offence against sub-section 5 (1), the Supreme Court has, in addition to any other powers of the Supreme Court, the same powers as the Court of Petty Sessions would have had if—

(a) proceedings in relation to the offence had been instituted in the Court of Petty Sessions; and

(b) the Court of Petty Sessions had found the offence proved.

(9) In proceedings for an offence against sub-section (2), it is a defence that the defendant had lawful authority to supply the substance in relation to which the offence is alleged to have been committed.

(10) A reference in this section to the supply of a substance to a person does not include a reference to the administration of a substance to a person.

5. (1) A person who has a Schedule 8 substance or a Schedule 12 substance in his possession is guilty of an offence.

(2) A person who commits an offence against sub-section (1) is punishable, on conviction—

(a) if the offence is committed in relation to—

(i) a substance other than cannabis; or

(ii) a quantity of cannabis exceeding 25 grams in mass, by imprisonment for a term not exceeding 2 years or a fine not exceeding $2,000 or both such fine and imprisonment; and

(b) if the offence is committed in relation to a quantity of cannabis not exceeding 25 grams in mass—by a fine not exceeding $100.
(3) In proceedings for an offence against sub-section (1), it is a defence that the defendant had lawful authority to have in his possession the substance in relation to which the offence is alleged to have been committed.

6. (1) A person who administers a Schedule 8 substance to himself is guilty of an offence.

(2) A person who administers a Schedule 12 substance, other than cannabis, to himself is guilty of an offence.

(3) A person who uses cannabis is guilty of an offence.

(4) It is a defence to a prosecution for an offence against sub-section (1) for the defendant to prove that, in administering the substance to himself, he was acting in accordance with the directions of a medical practitioner.

7. (1) A person who administers a Schedule 12 substance to another person is guilty of an offence.

(2) A person, other than a medical practitioner or a dentist, who administers a Schedule 8 substance to another person is guilty of an offence.

(3) It is a defence to a prosecution for an offence against sub-section (2) for the defendant to prove that, when he administered the substance to the other person, he had reasonable grounds for believing that the administration of the substance to that person was in accordance with the directions of a medical practitioner.

(4) A medical practitioner or dentist who administers a Schedule 8 substance to another person is guilty of an offence unless—

(a) in the case of a medical practitioner—the substance is administered in the course of medical treatment; and

(b) in the case of a dentist—

(i) the substance is cocaine, pethidine, pentazocine or a salt of cocaine, pethidine or pentazocine; and

(ii) the substance is administered in the course of dental treatment.

8. (1) A person who commits an offence against sub-section 6 (3) is punishable, on conviction, by a fine not exceeding $100.

(2) A person who commits an offence against sub-section 10 (5) is punishable, on conviction, by a fine not exceeding $500.
(3) A person who commits an offence against a provision of this Part, being an offence for which a penalty is not provided by any other provision of this Part, is punishable, on conviction, by imprisonment for a term not exceeding 2 years or a fine not exceeding $2,000, or both such imprisonment and fine.

Division 2—Authority to Possess Addictive Substances

9. Subject to Division 6, for the purposes of this Ordinance—
   (a) a medical practitioner, a pharmacist or a veterinary surgeon has lawful authority to have a Schedule 8 substance in his possession if he has the substance in his possession for use in the practice of his profession;
   (b) a dentist has lawful authority to have in his possession a Schedule 8 substance, being cocaine, pethidine, pentazocine or a salt of cocaine, pethidine or pentazocine, if he has the substance in his possession for use in the practice of his profession;
   (c) a person who is—
      (i) an officer or employee of the Department of Health authorized by the Director-General of Health to have a Schedule 8 substance in his possession; or
      (ii) an officer or employee of the Capital Territory Health Commission authorized by the Chairman to have a Schedule 8 substance in his possession, has lawful authority to have the substance in his possession;
   (d) a person to whom a medical practitioner or a veterinary surgeon is authorized to supply a Schedule 8 substance, and any person who receives that substance on behalf of that first-mentioned person, each has lawful authority to have that substance in his possession;
   (e) a person to whom a pharmacist is authorized to supply a Schedule 8 substance in accordance with a prescription signed by a medical practitioner or a veterinary surgeon authorizing the supply of the substance to that person, and any person who receives that substance on behalf of that first-mentioned person, each has lawful authority to have that substance in his possession;
   (f) a person who is authorized under section 10 to have in his possession a Schedule 8 substance has lawful authority to have the substance in his possession;
   (g) a person who is the holder of a licence under a law in force in the Territory, being a licence that authorizes him to sell...
a Schedule 8 substance, has lawful authority to have the substance in his possession if he has the substance in his possession for the purpose of sale;

(h) a person who, in the course of a business conducted by him, is engaged to transport a Schedule 8 substance has lawful authority to have the substance in his possession for so long as is reasonably necessary for the delivery of the drug to the person to whom the drug was consigned;

(j) a person whose possession of a Schedule 8 substance is permitted or authorized by a law in force in the Territory has lawful authority to have the substance in his possession; and

(k) a person employed by a person who, under any of the preceding paragraphs of this section, has lawful authority to have a Schedule 8 substance in his possession has lawful authority to have a Schedule 8 substance in his possession at any time when it is necessary, for the proper performance of his duties, to have the substance in his possession or in his custody.

10. (1) The Chairman may, by instrument in writing, authorize a person who is engaged in scientific or medical research to have in his possession, for the purposes of that research, a Schedule 8 substance or a Schedule 12 substance specified in the instrument.

(2) The Chairman may, by instrument in writing, authorize a person who is a member of the staff of a prescribed educational institution to have in his possession, for use in the performance of his duties as a member of the staff of that institution, a Schedule 8 substance or a Schedule 12 substance specified in the instrument.

(3) An authorization under sub-section (1) or (2) is subject to such conditions and restrictions (if any) as the Chairman thinks fit to specify in the instrument.

(4) In this section, "prescribed educational institution" means—

(a) an educational institution prescribed for the purpose of sub-section (2); or

(b) an educational institution that is included in a class of educational institutions prescribed for the purpose of sub-section (2).

(5) Where an authorization under sub-section (1) is subject to a condition or restriction, the person to whom the authorization was granted is guilty of an offence if he fails to comply with that condition or restriction.
(6) For the purposes of this Part, a person who is authorized, under sub-section (1) or (2), to have a Schedule 12 substance in his possession has lawful authority to have that substance in his possession.

Division 3—Supply and Obtaining of Addictive Substances

11. (1) Subject to Division 6, for the purposes of this Ordinance, a person has lawful authority to supply a Schedule 8 substance to another person otherwise than by way of administration of the substance to the person if—

(a) the first-mentioned person is a medical practitioner and the other person is receiving medical treatment from that practitioner;

(b) the first-mentioned person is a veterinary surgeon and the substance is supplied by that veterinary surgeon to the other person for administration to an animal in the custody of that other person, being an animal that is receiving veterinary treatment from that veterinary surgeon;

(c) the first-mentioned person is a pharmacist, the other person is a person who has presented to that pharmacist a prescription signed by a medical practitioner or a veterinary surgeon authorizing the supply of the Schedule 8 substance to that other person or to a third person for whom that other person receives the substance and the quantity of the substance supplied is the quantity authorized by the prescription to be supplied;

(d) the first-mentioned person is a pharmacist, the other person is a medical practitioner or a veterinary surgeon and that other person has presented to the pharmacist a written order signed by that other person for the supply to him of that Schedule 8 substance;

(e) the first-mentioned person is a pharmacist, the other person is a dentist, the Schedule 8 substance is cocaine, penta-zocine, pethidine or a salt of cocaine, pethidine or penta-zocine and the dentist has presented to the pharmacist a written order signed by the dentist for the supply to him of that Schedule 8 substance;

(f) the first-mentioned person is the holder of a licence under a law in force in the Territory, being a licence that authorizes him to supply the Schedule 8 substance, or is an employee of the holder of such a licence, the other person is a person who, by virtue of paragraph (a), (b), (c), (f) or (g) of section 9, has lawful authority to have the Schedule 8 substance in his possession and there is
presented to the first-mentioned person a written order signed by the other person for the supply of the Schedule 8 substance to that other person; or

(g) the first-mentioned person is the holder of a licence under a law in force in the Territory, being a licence that authorizes him to sell the Schedule 8 substance, or is an employee of the holder of such a licence, the other person is an employee of a person who, by virtue of paragraph (a), (b), (c), (f) or (g) of section 9, has lawful authority to have the Schedule 8 substance in his possession, and there is presented to the first-mentioned person a written order signed by the employer of that other person for the supply of that Schedule 8 substance to that other person.

(2) Where a medical practitioner or a veterinary surgeon has requested a pharmacist to supply to the practitioner or surgeon a Schedule 8 substance for immediate use in the administration of emergency treatment to a person or animal, the pharmacist has lawful authority to supply the Schedule 8 substance to the medical practitioner or veterinary surgeon, as the case may be.

(3) Where a Schedule 8 substance has been supplied to a medical practitioner or veterinary surgeon in pursuance of sub-section (2), the medical practitioner or veterinary surgeon shall, within the period of 24 hours after the supply of the substance, deliver to the pharmacist a statement in writing setting out particulars of—

(a) the request made by the medical practitioner or veterinary surgeon, as the case may be;

(b) where the substance was administered to a person—the name of that person; and

(c) where the substance was administered to an animal—the name of the keeper of the animal.

(4) For the purposes of any law of the Territory relating to the retention or recording of prescriptions—

(a) a statement delivered in accordance with sub-section (3) shall be taken to be a prescription; and

(b) the Schedule 8 substance referred to in the statement shall be taken to have been delivered in accordance with a prescription.

(5) A person who fails to comply with sub-section (3) is guilty of an offence.
12. (1) A person who is a medical practitioner, a veterinary surgeon or a dentist shall not obtain a Schedule 8 substance for use in the practice of his profession from a person other than—

(a) a pharmacist; or

(b) a person who is the holder of a licence under a law in force in the Territory or a law of a State or another Territory, being a licence that authorizes him to supply that Schedule 8 substance to another person.

(2) A pharmacist shall not obtain a Schedule 8 substance for use in the practice of his profession from a person other than a person who is the holder of a licence under a law in force in the Territory or a law of a State or another Territory, being a licence that authorizes the holder to supply that Schedule 8 substance to another person.

(3) A person who contravenes sub-section (1) or (2) is guilty of an offence.

(4) It is a defence to a prosecution for an offence against sub-section (3) arising out of a contravention of sub-section (1) that, when the Schedule 8 substance was supplied to the defendant, the defendant had reasonable grounds for believing—

(a) that the person by whom the substance was so supplied was a pharmacist; or

(b) that that person was the holder of a licence under a law in force in the Territory or a law of a State or another Territory authorizing him to supply the substance to the defendant.

(5) It is a defence to a prosecution for an offence against sub-section (3) arising out of a contravention of sub-section (2) that, when the Schedule 8 substance was supplied to the defendant, the defendant had reasonable grounds for believing that the person by whom the substance was so supplied was the holder of a licence under a law in force in the Territory or a law of a State or another Territory authorizing him to supply the substance to the defendant.

Division 4—Restrictions on the Prescribing of Addictive Substances

13. (1) This section applies to every Schedule 8 substance, other than—

(a) amphetamine and its salts;

(b) dexamphetamine and its salts;

(c) methylamphetamine and its salts;

(d) methylphenidate and its salts; and

(e) phenmetrazine and its salts.
(2) Except in accordance with an approval given under this Part, a medical practitioner shall not supply to, or prescribe for, a person a substance to which this section applies if the substance is supplied or prescribed for therapeutic use by that person for a period exceeding 2 months or for a period that would, together with a period for which any substance to which this section applies has, to the knowledge of the medical practitioner, been used continuously by that person, exceed 2 months.

(3) Except in accordance with an approval given under this Part, a medical practitioner shall not prescribe for a person a substance to which this section applies unless the medical practitioner has reasonable grounds for believing—

(a) that the supply of the substance to the person is reasonably necessary for the treatment of a mental or physical condition from which the person is suffering; and

(b) that the person is not an addict.

(4) A medical practitioner who contravenes sub-section (2) or (3) is guilty of an offence.

14. (1) This section applies to—

(a) amphetamine and its salts;
(b) dexamphetamine and its salts;
(c) methylamphetamine and its salts;
(d) methylphenidate and its salts; and
(e) phenmetrazine and its salts.

(2) Except in accordance with an approval given under this Part, a medical practitioner shall not supply to, or prescribe for, a person a substance to which this section applies unless—

(a) the person is suffering from narcolepsy; or
(b) the person, being a person who has not attained the age of 13 years, is hyperkinetic as a result of brain damage.

(3) Except in accordance with an approval given under this Part, a medical practitioner shall not supply to, or prescribe for, a person a substance to which this section applies if any substance to which this section applies has, to the knowledge of the medical practitioner, been administered to the person for a period not less than 2 months or for periods that, in the aggregate, are not less than 2 months.

(4) A medical practitioner who contravenes sub-section (2) or (3) is guilty of an offence.
Division 5—Approvals

15. (1) An application for an approval under this Part—
(a) shall be in writing signed by the medical practitioner making the application;
(b) shall state—
(i) the name of the applicant and the address, or an address, at which the applicant carries on the practice of his profession;
(ii) the name of the substance to which the application relates;
(iii) the strength and form in which that substance is to be administered;
(iv) the dosage in which, and the frequency with which, that substance is to be administered;
(v) the name and place of residence of the person to whom that substance is to be administered; and
(vi) the condition from which that person is suffering that, in the opinion of the applicant, necessitates the administration of the substance to which the application relates; and
(c) in the case of an application relating to a substance to which section 13 applies, shall state whether, in the opinion of the applicant, the person referred to in sub-paragraph (b) (v) is an addict.

(2) An application for an approval under this Part shall—
(a) be enclosed in an envelope that is sealed and marked with the word “confidential”; and
(b) be given to the Medical Officer of Health.

16. (1) Subject to sub-section (2), the Medical Officer of Health may in his discretion—
(a) grant or refuse an application for an approval under this Part; or
(b) refer the application to the Committee.

(2) Where, in an application for an approval under this Part in relation to a substance to which section 13 applies, a medical practitioner states that, in his opinion, a person is an addict—
(a) if the application is made on the ground that the administration of the substance to the person is necessary for the treatment of an organic disease—the Medical Officer of Health may either grant the application or refer the application to the Committee; and
(b) in any other case—the Medical Officer of Health shall refer the application to the Committee.

17. (1) There shall be, for the purposes of this Part, a committee to be called the Drugs Advisory Committee.

(2) The Committee shall consist of 3 members who shall be appointed, as occasion requires, by the Minister.

(3) A person is not eligible for appointment under sub-section (2) unless he is a medical practitioner.

(4) The Minister shall so exercise his power under sub-section (2) as to ensure that, at all times—
   (a) at least one member of the Committee is a person who has had experience in the teaching or practice of psychiatry; and
   (b) one member is a person nominated on behalf of the members of the Australian Capital Territory Medical Association.

(5) A member of the Committee holds office for such period, not exceeding 3 years, as is specified in the instrument of his appointment, and is eligible for re-appointment.

(6) A member of the Committee may resign his office by writing under his hand given to the Minister.

18. (1) The Minister shall, as occasion requires, appoint one of the members of the Committee to be the Chairman of the Committee.

(2) The Chairman of the Committee may resign his office of Chairman by writing under his hand given to the Minister.

19. Where an application for an approval under this Part is referred to the Committee, the Chairman of the Committee shall make arrangements for the application to be considered by the Committee.

20. (1) Where an application for an approval under this Part has been referred to the Committee, the Committee may, in its discretion—
   (a) direct the Medical Officer of Health to grant the approval;
   (b) direct the Medical Officer of Health to grant the approval, but in terms different from the approval sought; or
   (c) direct the Medical Officer of Health not to grant the approval.

(2) The Chairman of the Committee shall notify the Medical Officer of Health, in writing, of each direction given by the Committee under sub-section (1).
Variation of approvals

21. (1) The Medical Officer of Health may, at any time and in his discretion, vary or revoke an approval granted under section 16.
    (2) The Committee may, at any time and in its discretion, direct the Medical Officer of Health to vary or revoke an approval granted under section 20.
    (3) The Chairman of the Committee shall notify the Medical Officer of Health, in writing, of each direction given by the Committee under sub-section (2).
    (4) The Medical Officer of Health shall comply with a direction given by the Committee under sub-section (2).

22. (1) An approval under this Part, and a variation or revocation of such an approval, shall be in writing signed by the Medical Officer of Health.
    (2) An approval under this Part—
        (a) shall specify the quantity of the substance to which the approval relates that may be supplied or prescribed;
        (b) shall specify the period during which that substance may be so supplied or prescribed; and
        (c) is subject to such conditions (if any) as the Medical Officer of Health or, in the case of an approval granted under section 20, the Committee thinks fit to impose.

23. (1) An approval under this Part is effective for the purposes of this Part from the time at which the approval is signed by the Medical Officer of Health.
    (2) A variation or revocation of an approval under this Part is effective for the purposes of this Part from the time at which the variation or revocation, as the case may be, is given to the person who was the applicant for the approval.

24. (1) A medical practitioner who, by prescription, directs the supply to a person of a substance to which section 14 applies shall—
        (a) if the prescription is issued in pursuance of an approval under this Part, endorse on the prescription “Section 14 Approval”; and
        (b) if the prescription is not issued in pursuance of such an approval—endorse on the prescription “Section 14”.
    (2) A registered pharmacist shall not supply to another person a substance to which section 14 applies unless—
        (a) there has been presented to the pharmacist a prescription endorsed in accordance with sub-section (1) and directing the supply of the substance; and
(b) where the prescription bears the endorsement referred to in paragraph (1) (a)—the prescription has attached to it the approval in pursuance of which it is given.

(3) A person who contravenes, or fails to comply with, sub-section (1) or (2) is guilty of an offence.

Division 6—Cancellation and Surrender of Authority to Possess or Supply Schedule 8 Substances

25. This Division applies to a person who is—
   (a) a medical practitioner;
   (b) a veterinary surgeon;
   (c) a pharmacist; or
   (d) a dentist.

26. (1) Where a person to whom this Division applies is convicted of an offence against this Part, the Court by which the person is convicted may, if it is satisfied that it would be in the interests of the convicted person or of the public to do so, direct that the person shall not, during the period specified in the direction—
   (a) supply or administer any Schedule 8 substance to another person;
   (b) prescribe any Schedule 8 substance for another person; or
   (c) have any Schedule 8 substance in his possession for use in the carrying on of his profession.

(2) Where a direction has been given under sub-section (1), the person to whom the direction relates does not, during the period specified in the direction, have lawful authority—
   (a) to supply or administer any Schedule 8 substance to another person; or
   (b) to have any Schedule 8 substance in his possession for use in the carrying on of his profession.

(3) Where a direction has been given under sub-section (1), the person to whom the direction relates shall not, during the period specified in the direction, prescribe any Schedule 8 substance for another person.

(4) Section 208 of the Court of Petty Sessions Ordinance 1930 applies in relation to a direction given by the Court of Petty Sessions under sub-section (1) of this section as if the direction were a penalty imposed by that Court in respect of the conviction of a person of an offence.

(5) A person who contravenes sub-section (3) is guilty of an offence.

27. (1) A person to whom this Division applies may, by notice in writing given to the Chairman, declare that he does not wish to have lawful authority to possess any Schedule 8 substance for use in the practice of his profession or to supply any such substance.

Authorised by the ACT Parliamentary Counsel—also accessible at www.legislation.act.gov.au
(2) A person who has made a declaration under sub-section (1) may revoke the declaration by notice in writing given to the Chairman.

(3) The revocation of a declaration made under sub-section (1) has effect on and from the day on which notice is given to the Chairman for the purpose of sub-section (2).

(4) While a declaration under sub-section (1) is in force, the person by whom the declaration was made does not have lawful authority—
   (a) to supply or administer any Schedule 8 substance to another person; or
   (b) have any Schedule 8 substance in his possession for use in the practice of his profession.

(5) While a declaration under sub-section (1) is in force, the person who made the declaration shall not prescribe any Schedule 8 substance for another person.

(6) For the purposes of this section, a notice may be given to the Chairman—
   (a) by leaving the notice at an office of the Chairman with a person apparently employed in that office; or
   (b) by post addressed to the Chairman at an office of the Chairman.

(7) A person who contravenes sub-section (5) is guilty of an offence.

**Division 7—Miscellaneous**

28. (1) This section applies to a person who has lawful authority to supply a Schedule 8 substance to another person.

(2) A person to whom this section applies shall keep a Register of Drugs of Addiction.

(3) Where a person to whom this section applies receives a Schedule 8 substance, he shall, not later than 24 hours after he receives the substance, enter in the Register of Drugs of Addiction—
   (a) the date on which the substance was received;
   (b) the name and quantity of the substance;
   (c) the form in which the substance was so received; and
   (d) the name and address of the person from whom the substance was received.
(4) Where a person to whom this section applies disposes of a Schedule 8 substance, he shall, not later than 24 hours after he disposes of the substance, enter in the Register of Drugs of Addiction—

(a) the date of such disposal;
(b) the name and quantity of the substance disposed of;
(c) the form of the substance;
(d) if the substance was supplied to a person, the name of that person and the address of his place of residence; and
(e) if the substance was not supplied to a person, particulars of the manner of its disposal.

(5) A person who fails to comply with a provision of this section is guilty of an offence.

29. (1) In this section, "licensed person" means a person who is the holder of a licence under a law in force in the Territory, being a licence that authorizes the person to supply Schedule 8 substances, or specified Schedule 8 substances, to another person.

(2) A person who knowingly presents, or causes to be presented, to a pharmacist a prescription for the supply of a Schedule 8 substance, being a prescription signed by a person who is not—

(a) a medical practitioner; or
(b) a veterinary surgeon,

is guilty of an offence.

(3) A person who knowingly presents, or causes to be presented, to a pharmacist a prescription for the supply of a Schedule 8 substance, being a prescription that has been altered without the authority of the person who signed the prescription, is guilty of an offence.

(4) A person who, without lawful authority, presents, or causes to be presented, to a pharmacist a prescription for the supply of a Schedule 8 substance to a person other than the first-mentioned person, is guilty of an offence.

(5) A person who, for the purpose of obtaining a Schedule 8 substance—

(a) presents, or causes to be presented, to a pharmacist a document purporting to be an order signed by a medical practitioner, a veterinary surgeon or a dentist for the supply of a Schedule 8 substance; and

(b) at the time at which he so presents the document, knows, or has reasonable grounds for suspecting, that the document is not signed by a medical practitioner, veterinary surgeon or dentist,

is guilty of an offence.
(6) A person who knowingly makes a false representation (whether by words or conduct, or both) to a medical practitioner for the purpose of obtaining from that medical practitioner a prescription for the supply of a Schedule 8 substance is guilty of an offence.

(7) A person who knowingly makes a false representation (whether by words or conduct, or both) to a veterinary surgeon for the purpose of obtaining from that veterinary surgeon a prescription for the supply of a Schedule 8 substance is guilty of an offence.

(8) A person who knowingly makes a false representation (whether by words or conduct or both) to a licensed person for the purpose of obtaining a Schedule 8 substance is guilty of an offence.

PART III—LABELLING AND PACKAGING

Division 1—Preliminary

Interpretation

30. (1) In this Part, unless the contrary intention appears—

"approved name" means—

(a) in relation to a scheduled substance that is a drug for the purposes of the Single Convention—the name used in respect of the substance in the Schedules to the Single Convention; and

(b) in relation to any other scheduled substance—

(i) if the substance is described by an English name in the British Pharmacopoeia, the Australian Pharmaceutical Formulary, the British Pharmaceutical Codex or the British Veterinary Codex—that English name;

(ii) if sub-paragraph (i) is not applicable to the substance but the substance has been given a name by the General Medical Council of the United Kingdom—the name so given;

(iii) if neither sub-paragraph (i) nor sub-paragraph (ii) is applicable to the substance but the substance has been given an English name by the British Standards Institution, the Standards Association of Australia or the International Organization for Standardization—the English name so given;

(iv) if none of the preceding sub-paragraphs is applicable to the substance but the World Health Organization has given a name to the substance as the International Non-proprietary name of the substance—that name; and
(v) if none of the preceding sub-paragraphs is applicable to the substance—the accepted scientific name of the substance or, if there is no such name, a name that is descriptive of the nature and origin of the substance;

"Australian Pharmaceutical Formulary" means the latest edition for the time being of the book called the Australian Pharmaceutical Formulary published by the Pharmaceutical Association of Australia or, if that edition has been added to or amended, that edition as added to or amended;

"British Pharmaceutical Codex" means—
(a) the latest edition for the time being of the book called the British Pharmaceutical Codex published by direction of the Council of the Pharmaceutical Society of Great Britain; or
(b) if that edition has been added to or amended—that edition as added to or amended;

"British Pharmacopoeia" means—
(a) the latest edition for the time being of the book called the British Pharmacopoeia published under the direction of the General Medical Council of the United Kingdom; or
(b) if that edition has been added to or amended—that edition as added to or amended;

"British Veterinary Codex" means—
(a) the latest edition for the time being of the book called the British Veterinary Codex published by direction of the Council of the Pharmaceutical Society of Great Britain; or
(b) if that edition has been added to or amended—that edition as added to or amended;

"immediate container" means the container, not being an immediate wrapper or a container intended for consumption with the substance, in which a scheduled substance is so packed that no other thing intervenes between the substance and the container;

"immediate wrapper" means the material, not being material intended for consumption, in which a scheduled substance is so wrapped that no other thing intervenes between the substance and that material;

"internal use", in respect of a scheduled substance other than—
(a) a substance prepared for topical use in the nose, eyes, ears, mouth or throat; or
(b) a douche for rectal, vaginal or urethral use, means administration orally, parenterally or by way of a body orifice;
“primary pack” means the outer package in which a scheduled substance is packed for sale by retail, excluding any wrapping, bag or carton in which the substance is placed at the time of sale;

“registered brand”, in relation to the manufacturer of a scheduled substance, means a trade mark registered under the Trade Marks Act 1955 and of which the manufacturer of the substance is the registered proprietor or a registered user within the meaning of that Act;

“selected container” means—

(a) a syringe intended to be discarded after being used once; or

(b) a container for substances for therapeutic use having a capacity not exceeding 10 millilitres;

“Single Convention” means the Convention entitled the Single Convention on Narcotic Drugs, 1961 that was adopted and opened for signature at New York on 30 March, 1961, being that Convention as in force in relation to the Commonwealth from time to time;

“therapeutic use” means use in or in connexion with—

(a) the preventing, diagnosing, curing or alleviating of a disease, ailment, defect or injury in persons or animals;

(b) the influencing, inhibiting or modifying of a physiological process in persons or animals; or

(c) the testing of the susceptibility of persons or animals to a disease or ailment;

“withholding period”, in relation to a substance prepared for use for agricultural purposes, means the period that must elapse between the use of the substance and the harvest or taking of plant or animal products that may be affected by the substance in order to ensure that the substance is not present in those plant or animal products.

(2) Where a provision of this Part requires, in relation to the supply of a substance, that the name of the manufacturer of the substance be stated on a label, the name to be so stated is—

(a) where the substance has been imported into Australia and is supplied in the immediate container or immediate wrapper in which it was imported—the name of the person by whom the substance was made or the person by whom, or whose behalf, the substance was so imported; and

(b) in any other case—the name of the person who last subjected the substance to any process or the person who packed the substance in the immediate wrapper or immediate container in which the substance is supplied.
31. (1) Subject to section 33, a word, expression or statement, other than the approved name of a scheduled substance, required by this Part to be written on a label, pack or container shall be written—

(a) in the English language;
(b) on the outside surface of the label, pack or container;
(c) in durable characters; and
(d) in such colour or colours as to afford a distinct contrast with the background colour.

(2) Subject to sub-section (3) and to section 33, a word, expression or statement required by this Part to be written on a label, pack or container shall be written in bold-faced letters not less than 1.5 millimetres high.

(3) A word, expression or statement required by this Part to be written on a label, pack or container in capital letters shall be so written in bold-faced sanserif capital letters.

(4) A person shall not supply a scheduled substance in a pack or container to which there is attached a label having written on it—

(a) any reference to this Ordinance;
(b) any comment on, or explanation of, a word or expression required by this Ordinance to be written on a label, being a comment or explanation that directly, or by implication, contradicts, qualifies or modifies that word or expression; or
(c) any statement that directly, or by implication, suggests that the scheduled substance has been recommended or approved by an authority of the Commonwealth, of a State or of a Territory.

(5) A person shall not supply a scheduled substance in an immediate container or primary pack to which a label is attached in such a way as to obscure—

(a) a word or expression required by this Part to be written or embossed on the container or pack; or
(b) any rib, groove, point or star that is required by this Part to be embossed on the outer surface of the container or pack.

32. Every label used in connexion with a scheduled substance shall be securely attached to the outside surface of the primary pack and immediate container.
33. (1) Where, by this Part, the expression "Dangerous Poison", the word "Poison", the word "Warning" or the word "Caution" is required to be written on a label, the expression or word shall be so written in accordance with this section.

(2) An expression or word referred to in sub-section (1) shall, together with any letter or numeral required by this Part to form part of the same line—

(a) be written in red on a white background;
(b) be surrounded by a rectangular red frame;
(c) form the first line of the label; and
(d) be written in sanserif capital letters of bold face.

(3) Subject to sub-section (6), no other word, letter, symbol or numeral shall be included in the same line as an expression or word specified in sub-section (1).

(4) The expression "Dangerous Poison" and the words "Poison" and "Caution" shall be written in letters of a height not less than—

(a) 1.5 millimetres; or
(b) one-half of the height of the largest letter or numeral on the label,

whichever is the greater.

(5) The word "Warning" shall be written in letters of a height not less than—

(a) 4.5 millimetres; or
(b) one-quarter of the height of the largest letter or numeral on the label,

whichever is the greater.

(6) Where a provision of this Part requires that the expression "Dangerous Poison", the word "Poison", the word "Warning" or the word "Caution", when written on a label, be followed by the letter "s" and a numeral, the letter and numeral shall form part of the same line as that expression or that word, as the case may be.

34. (1) This section applies to and in relation to the following phrases:

(a) Not to be taken;
(b) Use strictly as directed;
(c) Supply without prescription or possession without authority illegal;
(d) Supply without prescription illegal;
(e) Unauthorized supply illegal;
(f) Keep out of reach of children;
(g) If swallowed seek medical advice;
(h) Read safety directions before opening.
(2) Where a phrase set out in paragraph (a), (b), (c), (d) or (e) of sub-section (1) is required by this Part to be written on a label, the phrase shall—

(a) be so written on the label that the phrase or the first line of the phrase forms the line immediately below the words written on the label in accordance with section 33; and

(b) be written in sanserif capital letters of bold face.

(3) Where the phrase set out in paragraph (1)(f) is required by this Part to be written on a label, the phrase shall be written in red on a white background and shall—

(a) in the case of a Schedule 5 substance, form the line immedi­diately below the word “Warning”; 

(b) in the case of any other scheduled substance, form the line or lines immediately below that on which a phrase referred to in sub-section (2) is written; and

(c) be written in sanserif capital letters of bold face.

(4) Where a phrase set out in paragraph (1)(g) or (1)(h) is required by this Part to be written on a label, the phrase shall—

(a) form the line or lines immediately below the phrase “Keep out of reach of children”; and

(b) be written in sanserif capital letters of bold face.

35. Where the approved name of a substance is required by this Part to be written on a label or container, the name shall be so written—

(a) in the English language;

(b) on the outside surface of the label or container;

(c) in durable characters; and

(d) in sanserif capital letters of bold face in such a colour as to afford a distinct contrast with the background colour.

Division 2—Labelling

36. (1) A person shall not supply to another person by way of wholesale or by way of retail a preparation consisting of, or containing, a scheduled substance, being a substance that is contained in an immediate wrapper, unless—

(a) the immediate wrapper is enclosed in a primary pack;

(b) the primary pack bears a label stating the matters required by section 37 to be written on the label on a primary pack in which an immediate container is enclosed; and

(c) the immediate wrapper bears a label—

(i) stating the name and address, or bearing the regis­tered brand, of the manufacturer of the preparation;

(ii) stating the approved name of the substance; and
(iii) stating the quantity of the scheduled substance included in the preparation or the proportion of the total ingredients of the preparation represented by the scheduled substance.

(2) A post office, cable, telegraphic or code address is not an address for the purposes of sub-section (1).

37. (1) Subject to sub-sections (7) and (8), a person shall not supply to another person a preparation consisting of, or containing, a scheduled substance unless the immediate container and, if there is a distinct primary pack, that pack each bears a label that has written upon it the matter required by sub-sections (2) and (3) in relation to that substance.

(2) The matter required by this sub-section in relation to a scheduled substance described in Column 1 of the following table is the word and phrases, and the letter and numeral (if any), set out in Column 2 of that table opposite to the description of the substance in Column 1.

<table>
<thead>
<tr>
<th>Column 1</th>
<th>Column 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Schedule 1 substance prepared for internal use</td>
<td>Poison S1</td>
</tr>
<tr>
<td></td>
<td>Use strictly as directed</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
<tr>
<td>Schedule 1 substance prepared for any purpose other than internal use</td>
<td>Poison S1</td>
</tr>
<tr>
<td></td>
<td>Not to be taken</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
<tr>
<td>Schedule 2 substance prepared for internal use or for topical oral use</td>
<td>Caution S2</td>
</tr>
<tr>
<td></td>
<td>Use strictly as directed</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
<tr>
<td>Schedule 2 substance prepared for any purpose other than internal use or topical oral use</td>
<td>Poison S2</td>
</tr>
<tr>
<td></td>
<td>Not to be taken</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
<tr>
<td>Schedule 3 substance</td>
<td>Caution S3</td>
</tr>
<tr>
<td></td>
<td>Use strictly as directed</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
<tr>
<td>Schedule 4 substance</td>
<td>Caution S4</td>
</tr>
<tr>
<td></td>
<td>Supply without prescription illegal</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
<tr>
<td>Schedule 5 substance</td>
<td>Warning</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
<tr>
<td>Schedule 6 substance prepared for internal use in animals</td>
<td>Caution</td>
</tr>
<tr>
<td></td>
<td>Use strictly as directed</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
<tr>
<td>Schedule 6 substance prepared for any purpose other than internal use in animals</td>
<td>Poison</td>
</tr>
<tr>
<td></td>
<td>Not to be taken</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
<tr>
<td></td>
<td>Read safety directions before opening</td>
</tr>
<tr>
<td>Schedule 7 substance prepared for internal use</td>
<td>Caution S7</td>
</tr>
<tr>
<td></td>
<td>Unauthorized supply illegal</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
<tr>
<td>Schedule 7 substance prepared for any purpose other than internal use</td>
<td>Dangerous Poison S7</td>
</tr>
<tr>
<td></td>
<td>Not to be taken</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
<tr>
<td></td>
<td>Read safety directions before opening</td>
</tr>
<tr>
<td>Schedule 8 substance</td>
<td>Caution S8</td>
</tr>
<tr>
<td></td>
<td>Supply without prescription or possession with authority illegal</td>
</tr>
<tr>
<td></td>
<td>Keep out of reach of children</td>
</tr>
</tbody>
</table>
(3) The matter required by this sub-section in relation to a scheduled substance is—

(a) the approved name of the scheduled substance;

(b) either—

(i) the quantity of the substance included in the preparation; or

(ii) the proportion of the total ingredients of the preparation represented by the substance;

(c) where the preparation, not being a Schedule 4 substance or a Schedule 8 substance, is made up for a particular purpose, directions for the use of the preparation;

(d) the name and address of the manufacturer of the preparation;

(e) if the preparation contains a scheduled substance specified in Column 1 of an item in Schedule 10, the statement set out in Column 2 of that item;

(f) if the preparation contains a substance specified in Column 1 of Part II of Schedule 11, the directions set out in the paragraph or paragraphs of Part I of that Schedule specified in Column 2 of Part II of that Schedule opposite to the specification of the substance; and

(g) in the case of a preparation for use for agricultural purposes, the withholding period (if any) of the scheduled substance.

(4) Subject to sub-sections (7) and (8), a person shall not supply to another person a preparation consisting of, or containing, a scheduled substance prepared for use only in treating animals unless the immediate container and, if there is a distinct primary pack, that pack each bears a label that has written upon it, in addition to the matter required by sub-sections (2) and (3), the words “For animal treatment only”.

(5) Subject to sub-sections (7) and (8), a person shall not supply to another person a preparation consisting of, or containing, a scheduled substance prepared for use only as a pesticide unless the immediate container and, if there is a distinct primary pack, that pack each bears a label that has written upon it, in addition to the matter required by sub-sections (2) and (3)—

(a) the words “Not to be used for any other purpose” in capital letters not less than 2.5 millimetres high; and

(b) a statement of the withholding period (if any) in capital letters and forming a separate line or lines on the label.

(6) Nothing in this section requires that a label be applied to a container, wrapper or cover used solely for the purpose of transporting a scheduled substance and not forming part of a primary pack.
(7) A person does not contravene sub-section (1), (4) or (5) by supplying to another person a preparation consisting of, or containing, a scheduled substance in a quantity exceeding 25 litres in volume or 25 kilogrammes in weight if—

(a) the preparation is not supplied by retail; and

(b) the immediate container bears a label that has written upon it—

(i) the word "Poison" in capital letters;

(ii) the approved name of the scheduled substance; and

(iii) the name of the manufacturer of the preparation.

(8) Where—

(a) a person supplies a preparation consisting of, or containing, a scheduled substance to another person;

(b) the preparation is so supplied in a selected container; and

(c) that selected container is enclosed in a primary pack that bears a label having written upon it the matter required by the preceding provisions of this section,

it is sufficient compliance with sub-section (1), (4) or (5), as the case may be, in relation to the selected container, if the selected container bears a label having written upon it—

(c) the word or expression specified in the Second Column of the following table opposite to the description in the First Column of that table applicable to that substance written in capital letters;

(d) the name or registered brand of the manufacturer of the preparation;

(e) the approved name of the substance;

(f) the quantity of the scheduled substance included in the preparation or the proportion of the total ingredients of the preparation represented by the scheduled substance; and

(g) in the case of a preparation for use only in treating animals, the words "For animal treatment only" written in capital letters.

<table>
<thead>
<tr>
<th>First Column</th>
<th>Second Column</th>
</tr>
</thead>
<tbody>
<tr>
<td>Schedule 1 substance</td>
<td>Poison</td>
</tr>
<tr>
<td>Schedule 2 substance prepared for internal use</td>
<td>Caution</td>
</tr>
<tr>
<td>Schedule 2 substance not prepared for internal use</td>
<td>Poison</td>
</tr>
<tr>
<td>Schedule 3 substance</td>
<td>Caution</td>
</tr>
<tr>
<td>Schedule 4 substance</td>
<td>Caution</td>
</tr>
<tr>
<td>Schedule 5 substance</td>
<td>Warning</td>
</tr>
<tr>
<td>Schedule 6 substance prepared for internal use</td>
<td>Caution</td>
</tr>
<tr>
<td>Schedule 6 substance not prepared for internal use</td>
<td>Poison</td>
</tr>
<tr>
<td>Schedule 7 substance prepared for internal use</td>
<td>Caution</td>
</tr>
<tr>
<td>Schedule 7 substance not prepared for internal use</td>
<td>Dangerous Poison</td>
</tr>
<tr>
<td>Schedule 8 substance</td>
<td>Caution</td>
</tr>
</tbody>
</table>
38. (1) A reference in this Part to the quantity of a scheduled substance that is included in a preparation shall be read—

(a) in relation to a preparation in the form of a tablet, capsule, pastille, packaged single dose of powder or similar unit—as a reference to the quantity of the scheduled substance contained in each tablet, capsule, pastille, dose or unit, as the case may be;

(b) in relation to a solid preparation from which a single dose may be prepared or from which a single stated amount of liquid may be prepared for therapeutic use—as a reference to the quantity of the scheduled substance in the immediate container;

(c) in relation to a liquid preparation for internal use—as a reference to the volume of the normal dose of the liquid and the quantity of the scheduled substance in that volume of the liquid; and

(d) in relation to any other preparation—as a reference to the proportion of the total ingredients of the preparation represented by the scheduled substance.

(2) A reference in this Part to the proportion of the total ingredients of a preparation represented by a scheduled substance shall be read—

(a) in the case of a liquid scheduled substance in a liquid preparation, as a reference to either the weight or volume of the scheduled substance present in the preparation per unit of volume of the preparation;

(b) in the case of a liquid scheduled substance in a solid or semi-solid preparation, as a reference to either the weight or the volume of the scheduled substance present in the preparation per unit of weight of the preparation;

(c) in the case of a solid or semi-solid scheduled substance in a liquid preparation, as a reference to the weight of the scheduled substance present in the preparation per unit of volume of the preparation;

(d) in the case of a solid or semi-solid scheduled substance in a solid or semi-solid preparation, as a reference to the weight of the scheduled substance present in the preparation per unit of weight of the preparation;

(e) in the case of a gaseous scheduled substance in a liquid preparation, as a reference to the weight of the scheduled substance present in the preparation per unit of volume of the preparation;
(f) in the case of a gaseous scheduled substance in a solid or semi-solid preparation, as a reference to the weight of the scheduled substance present in the preparation per unit of weight of the preparation; and

(g) in the case of a gaseous scheduled substance in a gaseous preparation, the weight of the scheduled substance present in the preparation per unit of weight of the preparation.

(3) Where the symbol “%” or the expression “per cent” appears in a schedule to this Ordinance or on a label in relation to a scheduled substance, the symbol or expression shall, unless the contrary intention appears, denote—

(a) in the case of a liquid preparation—the percentage weight of the scheduled substance in a specified volume of the preparation; and

(b) in the case of any other preparation—the percentage weight of the scheduled substance in a quantity of the preparation of a specified weight.

Division 3—Containers for Scheduled Substances

39. A person shall not supply to another person by way of wholesale or by way of retail a scheduled substance in an immediate container unless the immediate container complies with the requirements of this Division.

40. (1) The immediate container in which a scheduled substance is supplied shall—

(a) be made of a material that is—

(i) impervious to the scheduled substance; and

(ii) incapable of reacting chemically with the scheduled substance; and

(b) be so constructed as to prevent breakage of the container, or leakage of its contents, during the ordinary operations of handling, storage or transport.

(2) The immediate container in which a scheduled substance is supplied shall—

(a) have sufficient capacity to prevent breakage of the container, or leakage of its contents, by reason of expansion of those contents during handling, storage or transport of the container;

(b) be securely closed; and

(c) unless it contains a preparation packed for use on one occasion only, be capable of being closed securely after each occasion on which it has been opened.
41. (1) In this section, "hazardous substance" means—
   (a) a liquid hydrocarbon that distils at a temperature lower than 300° Celsius when tested according to method D86-61 of the American Society for Testing and Materials;
   (b) kerosene;
   (c) methylated spirit;
   (d) mineral turpentine;
   (e) oil of turpentine;
   (f) petrol; or
   (g) white spirit.

   (2) The immediate container in which a Schedule 5 substance, other than a hazardous substance, is supplied—
   (a) shall be so made as to be readily distinguishable from any container in which food or drink is supplied; and
   (b) shall have embossed or indelibly written on its side the words "Not to be used as a food container" or the words "Not to be taken".

   (3) The immediate container in which a hazardous substance is supplied shall comply with the requirements of section 42 as if the substance were a scheduled substance other than a Schedule 5 substance.

42. (1) A bottle or jar in which a scheduled substance, other than a Schedule 5 substance, is supplied, being a bottle or jar having a capacity not exceeding 2 litres—
   (a) shall, in the case of a glass bottle or jar, be made of colourless or brown glass;
   (b) shall have embossed on its outer surface the word "Poison" and the words "Not to be taken";
   (c) shall have embossed on its outer surface prominent vertical ribs or grooves or prominent points or stars of sufficient number to render the bottle or jar distinguishable by sight and by touch from bottles and jars ordinarily used as containers for any food, drink or condiment or for medicine for internal use; and
   (d) shall have on its outer surface a panel or panels free from ribs, grooves, points or stars for the affixing of a label or labels.

   (2) A container in which a scheduled substance, other than a Schedule 5 substance, is supplied, being a container having a capacity exceeding 2 litres shall either—
   (a) have embossed on the side of the container the word "Poison"; or
43. (1) Sections 41 and 42 do not apply to, or in relation to, the immediate container in which a scheduled substance prepared for internal use in humans or animals is supplied.

(2) Sections 41 and 42 do not apply to, or in relation to, the immediate container in which a scheduled substance is supplied if—
   (a) the substance is prepared for use as drops or a spray in the eyes, ears or nose; and
   (b) the quantity of the substance supplied in the container does not exceed 15 millilitres.

(3) The immediate container in which a scheduled substance prepared for use as drops in the eyes is supplied—
   (a) shall be so made as to be capable of being sterilized;
   (b) shall be fitted with a locking cap and a screw or bayonet fitting capable of delivering drops; and
   (c) shall, if the capacity of the container exceeds 15 millilitres, have embossed on its outer surface prominent vertical ribs or grooves or prominent points or stars of sufficient number to render the container distinguishable by sight and by touch from bottles and jars ordinarily used as containers for food, drink, condiments or medicines for internal use.

(4) Sections 41 and 42 do not apply to, or in relation to, the immediate container in which a scheduled substance is supplied where—
   (a) the container and its contents are prepared for use in a machine used in an automatic photographic or photocopying process; and
   (b) the container is designed to be fitted into the machine.

44. A person shall not supply to another person any food, drink or condiment or a preparation for internal use in a container that—
   (a) complies with the requirements of this Division in relation to the immediate container in which a scheduled substance prepared for external use may be supplied; or
   (b) is not readily distinguishable by sight and by touch from such a container.

45. A person shall not use a container that has the name of a scheduled substance embossed or otherwise permanently marked on it for the purpose of containing any substance other than that schedule substance.
Division 4—Miscellaneous

46. Divisions 2 and 3 do not apply to, or in relation to, the supply of a medicine, drug or preparation dispensed by a pharmacist in accordance with a prescription signed by a medical practitioner, veterinary surgeon or dentist if the container in which the medicine, drug or preparation is supplied has affixed to it a label on which are written—

(a) the words "Keep out of reach of children" in red letters on a white background;
(b) in the case of a prescription signed by a medical practitioner or dentist—the name of the person for whom the medicine, drug or preparation was prescribed;
(c) in the case of a prescription signed by a veterinary surgeon—the name of the keeper of the animal for whose treatment the medicine, drug or preparation was prescribed;
(d) the directions given in the prescription in relation to the administration of the medicine, drug or preparation;
(e) the name and address of the person by whom the medicine, drug or preparation is supplied;
(f) in the case of a medicine, drug or preparation prescribed for use otherwise than by way of internal use—the words "Caution—Not to be taken" in red letters; and
(g) unless the prescription contains a direction to the contrary or requires that the medicine, drug or preparation be compounded in accordance with a formula stated in the prescription—the approved name of the medicine, drug or preparation.

47. A person who contravenes, or fails to comply with, a provision of this Part is guilty of an offence and punishable, on conviction—

(a) in the case of a person not being a body corporate—by a fine not exceeding $1,000 or imprisonment for a term not exceeding 6 months or both such imprisonment and fine; and
(b) in the case of a body corporate—by a fine not exceeding $2,000.

PART IV—MISCELLANEOUS

48. Nothing in this Ordinance applies to, or in relation to, the possession, supply or use of any of the goods specified in Schedule 13, notwithstanding that the goods contain a scheduled substance.

49. (1) In this section, "Tribunal" means the Administrative Appeals Tribunal established by the Administrative Appeals Tribunal Act 1975.
(2) Application may be made to the Tribunal for a review of a decision of the Chairman—
(a) refusing to give an authorization under section 10;
(b) specifying a condition or restriction to which an authorization under that section is subject; or
(c) revoking, amending or varying an authorization under section 10.

50. (1) In this section, “prescribed substance” means a Schedule 8 or Schedule 12 substance in relation to which there are reasonable grounds for suspecting that an offence has been committed against a provision of Part II.

(2) If a magistrate is satisfied by information on oath that there is reasonable ground for suspecting that there is on any premises or at any place a prescribed substance, he may grant a search warrant authorizing a member of the Police Force of the Territory named in the warrant, with such assistance as he thinks necessary, to enter, if need be by force, at any time or times within the period of 28 days from and including the date of the warrant, upon the premises or place, to search the premises or place and any person found on the premises or place and to seize any prescribed substance found on the premises or place or upon any person on the premises or place.

(3) A person who seizes a prescribed substance in pursuance of a warrant under sub-section (2) shall deliver the substance to the Chairman.

(4) Where a prescribed substance is delivered to the Chairman in accordance with sub-section (3), the Chairman shall cause the substance to be destroyed.

51. The Minister may, by instrument in writing, appoint a person to be an analyst for the purposes of this Ordinance.

52. (1) In proceedings in the Court of Petty Sessions for an offence against this Ordinance, a certificate signed by an analyst and stating the result of an analysis of a substance carried out at the request of a member of the Police Force is evidence of the matters stated in the certificate.

(2) For the purpose of sub-section (1), a document that purports to be signed by an analyst shall, unless the contrary is proved, be taken to have been so signed.

53. (1) Where a court convicts a person of an offence against a provision of Part II, being an offence constituted by the possession of a Schedule 8 substance or a Schedule 12 substance, the court may declare the substance forfeit to the Commonwealth.
(2) Where a substance is forfeit to the Commonwealth by virtue of a declaration under sub-section (1), the substance shall, as soon as is reasonably practicable, be destroyed in such manner as the Chairman directs.

54. (1) This section applies to—
   (a) an analyst; and
   (b) a member of the Police Force.

(2) For the purposes of Part II, a person to whom this section applies has lawful authority to have a Schedule 8 substance or a Schedule 12 substance in his possession for the purposes of the performance of his official duties.

(3) For the purposes of Part II, a person to whom this section applies has lawful authority to deliver to another person to whom this section applies a Schedule 8 substance or a Schedule 12 substance received by the first-mentioned person in the course of the performance of his duties.

55. The Minister may make regulations, not inconsistent with this Ordinance, prescribing all matters which by this Ordinance are required or permitted to be prescribed or which are necessary or convenient to be prescribed for carrying out or giving effect to this Ordinance.
SCHEDULE 1  Section 3

Aconite (root ofaconitum napellus)
Antimony, compounds of (other than organic compounds), except in preparations containing the equivalent of 1 per cent or less of antimony trioxide and except antimony chlorides in polishes
Atropine, except—
(a) atropine in preparations specified in Schedule 2;
(b) atropine methonitrate.
Belladonna Herb, except in preparations containing 0.25 per cent or less of the alkaloids of belladonna calculated as hyoscyamine
Bromine, excluding its salts and derivatives
Brucine, except in preparations containing 0.2 per cent or less of brucine
Colchicine, except in preparations containing 0.5 per cent or less of colchicine
Coniine, except in preparations containing 0.1 per cent or less of coniine
Cotamine
Croton Oil
Homatropine, except in preparations containing 0.25 per cent or less of homatropine
Hydrocyanic Acid and Cyanides for therapeutic use, except in preparations containing the equivalent of 0.15 per cent or less of hydrocyanic acid
Hyoscine, except in preparations containing 0.25 per cent or less of hyoscine and except hyoscine butylbromide
Hyoscyamine, except in preparations containing 0.25 per cent or less of hyoscyamine
Hyoscyamus, except in preparations containing 0.25 per cent or less of the alkaloids of hyoscyamus calculated as hyoscyamine
Lobelia, except in preparations containing 0.5 per cent or less of the alkaloids of lobelia and except preparations for smoking or burning
Mercuric Chloride, except—
(a) in preparations containing 0.5 per cent or less of mercuric chloride;
(b) in batteries;
(c) when prepared for use for agricultural, pastoral, horticultural or industrial purposes.
Mercuric Iodide, except in preparations containing 2 per cent or less of mercuric iodide and except when prepared for use for agricultural, pastoral, horticultural or industrial purposes
Mercuric Nitrate, except in preparations containing the equivalent of 3 per cent or less of mercury (Hg), in such form
Mercuric Potassium Iodide, except in preparations containing the equivalent of 2 per cent or less of mercuric iodide, in such form
Mercuric Thiocyanate, except when prepared for use for photographic purposes
Mercury, organic compounds of, except—
(a) in preparations containing the equivalent of 0.5 per cent or less of mercury (Hg);
(b) for therapeutic use;
(c) when prepared for use for agricultural, pastoral or horticultural purposes;
(d) compounds specified in Schedule 7.
Nux Vomica
Phosphorus, yellow, excluding its salts and derivatives, except in preparations containing 0.5 per cent or less of free phosphorus
Savin Oil
Stramonium, except in preparations containing 0.25 per cent or less of alkaloids calculated as hyoscyamine, and except preparations for smoking or burning
Strychnine, except strychnine in preparations containing 1 per cent or less of strychnine when prepared for use in the destruction of vermin
Tansy Oil
Veratrum, except for therapeutic use
Acetic Acid (excluding its salts and derivatives) for therapeutic use, except in preparations containing 80 per cent or less of acetic acid.

Acetyldihydrocodeine, when compounded with one or more other medicaments, in preparations containing 1 per cent or less of acetyldihydrocodeine.

Antimony, compounds of (other than organic compounds), in preparations containing the equivalent of 1 per cent or less of antimony trioxide, except antimony chlorides in polishes.

Atropine, except atropine methonitrate, in preparations containing 0.25 per cent or less of atropine, and atropine sulphate 0.5 mg tablets in packs of six, when labelled for treatment of organo-phosphorus poisoning.

Belladonna Herb in preparations containing 0.25 per cent or less of the alkaloids of belladonna, calculated as hyoscyamine.

Brucine in preparations containing 0.2 per cent or less of brucine, except when used in concentrations of 0.02 per cent or less for the denaturation of alcohol.

Camphorated Oil.

Cantharidin in preparations containing 0.01 per cent or less of cantharidin.

Chloroform (excluding its derivatives) except—
(a) in preparations containing 10 per cent or less of chloroform;
(b) when specifically prepared and packed as a therapeutic agent for the induction of inhalation anaesthesia.

Codeine, when compounded with one or more other medicaments, in preparations containing 1 per cent or less of codeine.

Colchicine in preparations containing 0.5 per cent or less of colchicine.

Coniine in preparations containing 0.1 per cent or less of coniine.

Dextropropoxyphene in preparations containing 1 per cent or less of dextropropoxyphene.

Dextrophan in preparations containing 1 per cent or less of dextrophan.

Diamines (including phenylene, toluene and all other alkylated benzene diamine derivatives), except to the extent that they are included in Schedule 6.

Dihydromorphine when compounded with one or more other medicaments, in preparations containing 1 per cent or less of dihydromorphine.

Ether (excluding its derivatives, ether solvent and ether preparations for use in internal combustion engines), except—
(a) in preparations containing 10 per cent or less of ether;
(b) when specifically prepared and packed as a therapeutic agent for the induction of inhalation anaesthesia.

Ethoheptazine in preparations containing 1 per cent or less of ethoheptazine.

Ethylmorphine when compounded with one or more other medicaments, in preparations containing 1 per cent or less of ethylmorphine.

Ferrous Sulphate and other iron preparations for human internal use, except in preparations containing 5 per cent or less of iron.

Fluoride, metallic, including ammonium fluoride, when prepared for therapeutic use, except—
(a) in dentifrices containing 0.5 per cent or less of fluoride ion;
(b) in substances containing 15 parts per million or less of fluoride ion.

Gelsemium.

Homatropine in preparations containing 0.25 per cent or less of homatropine.

Hydrocyanic Acid and Cyanides in preparations containing the equivalent of 0.15 per cent or less of hydrocyanic acid.

Hyoscine in preparations containing 0.25 per cent or less of hyoscine, except hyoscine butylbromide.

Hyoscyamine in preparations containing 0.25 per cent or less of hyoscine.

Hyoscyamus in preparations containing 0.25 per cent or less of the alkaloids of hyoscyamus, calculated as hyoscyamine.

Iodine (excluding its salts and derivatives) except in preparations containing 2.5 per cent or less of iodine.

Iodophors containing more than 2.5 per cent available iodine.

Lead Salts and compounds of lead when prepared for medical or cosmetic use, except in preparations for hair dressing containing 1 per cent or less of lead.

Lobelia in preparations containing 0.5 per cent or less of the alkaloids of lobelia, except preparations for smoking or burning.

Metendazole for human therapeutic use.
SCHEDULE 2—continued

Mercuric Chloride in preparations containing 0.5 per cent or less of mercuric chloride, except—
(a) in batteries;
(b) when prepared for use for agricultural, pastoral, horticultural or industrial purposes

Mercuric Iodide in preparations containing 2 per cent or less of mercuric iodide, except when prepared for agricultural, pastoral or horticultural purposes

Mercuric Nitrate in preparations containing the equivalent of 3 per cent or less of mercury (Hg), in such form

Mercuric Oxide and all oxides of mercury

Mercuric Potassium Iodide in preparations containing the equivalent of 2 per cent or less of mercuric iodide, in such form

Mercury (Metallic), (excluding its salts and derivatives), except in scientific instruments

Mercury, organic compounds of, in preparations containing the equivalent of 0.5 per cent or less of mercury (Hg), except—
(a) when prepared for use for agricultural, pastoral or horticultural purposes;
(b) as a preservative in substances containing 0.01 per cent or less of mercury;
(c) compounds specified in Schedule 7

Niclosamide for human therapeutic use

Nicocodine when compounded with one or more other medicaments, in preparations containing 1 per cent or less of nicocodine

Nicodicodine when compounded with one or more other medicaments, in preparations containing 1 per cent or less of nicodicodine

Norcodeine when compounded with one or more other medicaments, in preparations containing 1 per cent or less of norcodeine

Phenol, any homologue of phenol boiling below 220°C and creosote, for therapeutic use except in preparations containing 3 per cent or less by weight of such substances or homologues

Pholcodine when compounded with one or more other medicaments, in preparations containing 1 per cent or less of pholcodine

Potassium Chlorate except in preparations containing 10 per cent or less of potassium chlorate

Silver Nitrate

Staphisagria except in preparations containing 0.2 per cent or less of staphisagria

Stramonium in preparations containing 0.25 per cent or less of the alkaloids calculated as hyoscyamine, except preparations for smoking or burning

Zinc Pyrithione except in preparations containing 2 per cent or less of zinc pyrithione
SCHEDULE 3 Section 3

Adrenaline, natural or synthetic, in preparations containing more than 0.01 per cent of adrenaline but not more than 1 per cent of adrenaline

Amyl Nitrite

Anaesthetics (Local), the following only—
(a) benzocaine;
(b) butyl aminobenzoate;
(c) orthocaine;
(d) benzamine lactate;
(e) lignocaine,
when included in—
(f) lozenges, pastilles, tablets and capsules containing 30 mg or less of such substances;
(g) suppositories or bougies containing 200 mg or less of such substances in each;
(h) preparations for external use, other than eyedrops, containing 10 per cent or less of such substances

Antihistamine substances—
(a) in preparations labelled and packed for the treatment of motion sickness in packs of 10 doses or less;
(b) in oral liquid cough preparations containing 0.3 per cent or less of such substances;
and
(c) in preparations labelled and packed as nasal preparations or eyedrops

Apomorphine

Bufexamac in preparations containing 5 per cent or less of bufexamac

Chloral Hydrate in preparations containing 5 per cent or less of chloral hydrate, except alpha-chloralose in preparations containing 5 per cent or less of alpha-chloralose and prepared for use as rodenticides

Chlorbutol in oral preparations containing 250 mg or less of chlorbutol per adult dosage unit

Cinnamedrine

Dextromethorphan in preparations containing 1 per cent or less of dextromethorphan when compounded with one or more other medicaments in such a way that the dextromethorphan contained therein cannot readily be extracted

Dicophane (DDT) in preparations for human therapeutic use

Dicyclomine in preparations containing 0.1 per cent or less of dicyclomine

Dimethisoquin in preparations for topical use

Diphenamid Methylsulphate in preparations for topical use

Ephedrine and Pseudoephedrine except—
(a) in preparations for internal use containing 0.5 per cent or less of ephedrine and pseudoephedrine;
or
(b) in other preparations containing 1 per cent or less of ephedrine and pseudoephedrine

Erythritol Tetrarnitrate and other nitric esters of polyhydric alcohols

Etadrenaline

Glyceryl Trinitrate

Guaiphenesin in preparations containing 120 mg or less of guaiphenesin per adult dosage unit

Hexachlorophane in preparations for skin cleansing purposes containing 3 per cent or less of hexachlorophane, except—
(a) in preparations for use on infants;
(b) in preparations for the treatment of animals;
(c) in preparations containing 0.1 per cent or less of hexachlorophane as a preservative

8-Hydroxyquinoline and its derivatives, for human therapeutic use, except—
(a) non-halogenated derivatives containing 1 per cent or less of the derivative for external use;
(b) 8-Hydroxyquinoline and its derivatives in forms in which they are specified in Schedule 4

Insulin and preparations containing the specific hypoglycaemic principle of the pancreas

Isoprenaline in preparations containing 1 per cent or less of isoprenaline, except when contained in metered aerosols delivering more than 80 micrograms per metered dose

Maladin in preparations for external human therapeutic use containing 2 per cent or less of maldison

Mercurous Chloride (Calomel) in preparations for internal use, except when contained in teething powders or preparations for infants

Methoxamine, except—
(a) in preparations for internal use containing 0.5 per cent or less of methoxamine; or
(b) in other preparations containing 1 per cent or less of methoxamine
SCHEDULE 3—continued

Methoxyphenamine
Methylephedrine
Naphazoline
Noradrenaline, in preparations containing more than 0.01 per cent of noradrenaline but not more than 1 per cent of noradrenaline
Noscapine
Octyl Nitrite
Oxethazaine in preparations for internal use only
Oxymetazoline
Papaverine
Phedrazine
Phenamazoline
Phenazone for external use
Phenylephrine, except—
   (a) in preparations for internal use containing 0.5 per cent or less of phenylephrine; or
   (b) in other preparations containing 1 per cent or less of phenylephrine
Propylhexedrine in appliances for inhalation in which the substance is absorbed upon an inert solid material
Propyphenazone
Pyrantel for human therapeutic use
Rinidol
Santonin
Sodium Nitrite for therapeutic use
Tetrahydrozoline
Tramazoline
Triclofos in preparations containing 5 per cent or less of triclofos
Trimizoline
Tymazoline
Viprynium
Xylometazoline
Acetanilide and alkyl acetanilides, for human therapeutic use
Acetazolamide
Acetohexamide
Acetylcholine and other choline esters
Acetylcysteine
Acetyldihydrocodeine when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of acetyldihydrocodeine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of acetyldihydrocodeine;
except in preparations specified in Schedule 2
Acetylmethyldimethyloximidophenylhydrazine
Adiphenine
Adrenaline, natural or synthetic, except in preparations containing 1 per cent or less of adrenaline
Alcuronium
Alphadolone
Alpha-receptor Blocking Agents, including phenotamine and phenoxybenzamine
Alphaxolone
Amantadine
Ambenonium
Ambuceteamide
Ambutonium
Amiloride
Aminometradine
Aminorex
Amiphenazole
Amisometradine
Amitriptyline and other compounds structurally derived there from by substitution in the side chain
Amoxycillin
Anabolic steroidal agents
Anaesthetics, (Local), being synthetic cocaine substitutes, except local anaesthetics specified in Schedule 3
Angiotensin Amide
Antibiotics including penicillin, penicillanic acid, streptomycin, chloramphenicol, tetracycline and any other antibiotic substances however derived and their chemical derivatives, except—
(a) antibiotic substances specified in Schedule 6;
(b) animal feedstuffs for growth promotion containing bacitracin and its salts, erythromycin, flavophospholipol, oleandomycin and its salts, tylosin and its salts and virginiamycin and its salts in concentrations of 50 parts per million or less of the total active antibiotic principle;
(c) milk replacers for calves and starter rations for pigs containing bacitracin and its salts, erythromycin and tylosin and its salts in concentrations of 100 parts per million or less of the total active antibiotic principle
Antifolic Acid substances, including aminopterin, teropterin and orthopterin
Antihistamine substances except in preparations specified in Schedule 3
Antimalarial substances including amodiaquine, chloroquine, mepacrine, pamaquine primaquine, pyrimethamine, proguanil and sontoquine, but not including quinine
Antimony, organic compounds of, for therapeutic use
Antitubercular substances, including isoniazid and its derivatives, para-aminosalicylic acid and thiacetazone
Aprotinin
Arecoline
Arsenic, organic compounds of, for human therapeutic use
Atropine Methonitrate
Azaperone
Azapetine
Azatadine
Baclofen
Barbituric Acid and its derivatives
Beclamide
Bemegride
Benactyzine and other substances structurally derived from diphenylmethane with ataractic properties when used for therapeutic purposes
Benzhexol
Benzilationum
Benzphetamine and other substances structurally derived from betaminopropylbenzene or beta-aminoisopropylbenzene by substitution in the side chain or by ring closure therein (or by both such substitution and such closure) except—
(a) substances specified in Schedule 3 or Schedule 8;
(b) ephedrine, pseudoephedrine and phenylephrine in preparations not specified in Schedule 3
Benztropine
Benzydamine
Betahistine
Beta-receptor Blocking Agents, including alprenolol and propranolol
Bethanidine
Biperiden
Bismuth subgallate for oral use in humans
Bleomycin
Boron Compounds for human therapeutic or cosmetic use, except in preparations for external use containing 1 per cent or less as boron
Bretylium
Bromhexine
Bromides, inorganic, for therapeutic use
Bromoform for therapeutic use
Bromvaletone
Bufexamac, except in preparations specified in Schedule 3
Bunamide
Busulphan
Butylchloral Hydrate
Calcitonin
Calcium Carbimide
Camphotamide
Cantharidin, except in preparations containing 0.01 per cent or less of cantharidin
Captodiame
Capuride
Caramiphen
Carbachol
Carbamazepine
Carbazochrome
Carbenoxolone for human therapeutic use, except preparations containing 2 per cent or less of carbenoxolone
Carbidopa
Carbimazole
Carbocromen
Carbromal
Cardiac Glycosides not specified in Schedules 1-3 or Schedules 5-8
Cephaolizin
Chiniofon and other halogenated 8-hydroxyquinoline derivatives for internal human use
Chloral Formamidine
Chloral Hydrate except in preparations containing 5 per cent or less of the equivalent of chloral hydrate
Chlorazanil
Chlorbutol in oral preparations, except in such preparations containing 250mg or less of chlorbutol per adult dosage unit
Chlordiazepoxide and other substances structurally derived from benzodiazepine with ataractic properties when used for therapeutic purposes
Chlormerodrin
SCHEDULE 4—continued

Chlormethiazole
Chlormezanone
Chloroform when specifically prepared and packed as a therapeutic agent for the induction of inhalation anaesthesia
Chlorthiazide and other substances structurally derived from benzothiadiazine for therapeutic use
Chlorphentermine
Chlorpromazine and other substances structurally derived from phenothiazine with ataractic properties when used for therapeutic purposes
Chlorpropamide
Chlorprothixene
Chlorzoxazone
Clidinium
Clofibrate
Clonazepam
Clonidine
Clorazepate
Clorexolone
Clozapine

Codeine when compounded with one or more other medicaments—
   (a) in divided preparations containing not more than 100 mg of codeine per dosage unit; or
   (b) in undivided preparations with a concentration of not more than 2.5 per cent of codeine;
      except in preparations specified in Schedule 2

Colaspase
Cortisone and steriod suprarenal cortical hormones, either natural or synthetic
Coumarin derivatives and phenylindanedione derivatives for therapeutic use
Curare, Tubocurarine, d-Tubocurarine, d-Tubocurarine dimethylether, and all synthetic quaternary ammonium compounds and other compounds having curarising properties
Cyclandelate
Cyclopentolate
Cyclopropane when specifically prepared and packed as a therapeutic agent for the induction of inhalation anaesthesia
Cycrinine

Dapsone and all derivatives of 4, 4-diaminodiphenylsulphone
Deanol
Demecarium Bromide
Desipramine
Dextromethorphan, except in preparations specified in Schedule 3
Dextropropoxyphene, except in preparations containing 1 per cent or less of dextropropoxyphene
Dextrorphan, except in preparations containing 1 per cent or less of dextrorphan
Dibenzepin
Dichloralphenazone
Dichlorphenamide
Dicyclomine, except in preparations containing 0.1 per cent or less of dicyclomine
Diethazine
Diethylcarbamazine for human therapeutic use
Diethylpropion
Difenoxin in preparations containing, per dosage unit, 0.5 mg or less of difenoxin (base) and a quantity of atropine sulphate equivalent to at least 5 per cent of the dose of difenoxin
Digitalis and its glycosides
Dihydrallazine
Dihydrocodeine when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of dihydrocodeine per
dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of
dihydrocodeine;
except in preparations specified in Schedule 2
Diisopropylamine Dichloroacetate for therapeutic use
Dimethoxanate
Dimethyl Sulphoxide for therapeutic use
Dinitocresols for therapeutic use
Dinitronaphthols for therapeutic use
Dinitrophenols for therapeutic use
Dinitrothymols for therapeutic use
Diphenoxylate in preparations containing per dosage unit 2.5 mg or less of diphenoxylate
(base), and a quantity of atropine sulphate equivalent to at least 1 per cent of the dose
of diphenoxylate
Dipyridamole
Disulfiram, except when used for industrial purposes
Dithiazone, except in preparations containing 2 per cent or less of dithiazone for the
treatment of animals
Droperidol
Emetine, except in preparations containing 0.2 per cent or less of emetine
Enflurane when specifically prepared and packed as a therapeutic agent for the induction of
inhalation anaesthesia
Epicellin
Ergot
Ethacrynic Acid
Ethamivan
Ether when specifically prepared and packed as a therapeutic agent for the induction of
inhalation anaesthesia
Ethoglucid
Ethoheptazine, except in preparations containing 1 per cent or less of ethoheptazine
Ethopropazine
Ethozolamide
Ethyl Chloride when specifically prepared and packed as a therapeutic agent for the induction
of inhalation anaesthesia
Ethylene when specifically prepared and packed as a therapeutic agent for the induction of
inhalation anaesthesia
Ethylmorphine when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of ethylmorphine per
dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of
ethylmorphine;
except in preparations specified in Schedule 2
Fencamfamin
Fenfluramine
Fenoterol
Fenpipramide
Fenpiprane
Flucytosine
Flufenamic Acid
Fluroxane when specifically prepared and packed as a therapeutic agent for the induction of
inhalation anaesthesia
Fluspirilene
Frusemide
Galanthamine
Gallamine
SCHEDULE 4—continued

Glibornuride
Glucagon
Glutethimide
Glycopyrronium
Glymidine
Guanacline
Guanethidine
Haloperidol and other substances structurally derived from butyrophenone with ataractic properties when used for therapeutic purposes
Halothane when specifically prepared and packed as a therapeutic agent for the induction of inhalation anaesthesia
Heparin
Hexachlorophane in preparations for use on infants and in all other preparations, except—
(a) preparations specified in Schedule 3 and 6;
(b) preparations, other than preparations for use on infants, containing 0.1 per cent or less of hexachlorophane as a preservative
Hexamethonium
Hexocyclium
Hydralazine
Hydroquinone for human therapeutic use, except in preparations containing 2 per cent or less of hydroquinone
1-Hydroxy-pyrido (3, 2, a) -5-phenoxazone-3-carboxylic Acid
Hydroxyurea
Hydroxyzine
Hygromycin, except—
(a) in preparations specified in Schedule 6;
(b) in preparations containing 20 parts per million or less of hygromycin
Hyoscine Butylbromide
Ibufenac
Ibuprofen
Idoxuridine
Imipramine
Indomethacin
Inositol Nicotinate, for internal use
Iron compounds, injectable preparations for human therapeutic use
Isocarboxazid
Isotocarine
Isometheptene
Isoprenaline, except in preparations specified in Schedule 3
Isopropamide
Khellin
Laudexhm Methys Iphate
Leptazol
Levamisole for human therapeutic use
Levodopa
Lidoflazinc
Lithium salts for therapeutic use, except in preparations containing 0.01 per cent or less of lithium
Lorazepam
Mafenide
Maldison for human therapeutic use, except in preparations specified in Schedule 3
Mazindol
Mebverine
Mecamylamine
Meclofenoxate
Medazepam
Mefenamic Acid
Mefruside
Mepenzolate
Mephenesin and its derivatives, except guaiphenesin in preparations specified in Schedule 3
Mephentermine
Meprobamate
Mercaptopurine and other substances structurally derived therefrom with cytotoxic properties when used for therapeutic purposes
Mercurous Chloride (Calomel) when contained in teething powders or preparations for infants
Mercury organic compounds of, for therapeutic use, except preparations for topical use containing 0.5 per cent or less of mercury
Metaraminol
Metformin
Methanthelinium
Methazolamide
Methimazole
Methixene
Methocarbamol
Methoxalen
Methoxyflurane when specifically prepared and packed as a therapeutic agent for the induction of inhalation anaesthesia
Methylbuprone
Methypentynol and other substituted alkynes for internal use
Methyprylon
Metoclopramide
Metolazone
Metronidazole
Metyrapone
Minocycline
Mithramycin
Mitobronitol
Monensin, except in animal feed containing 120 parts per million or less of monensin
Monoamine Oxidase Inhibitors, including iproniazid, isocarboxazid, nialamide, phenelzine, p-chlorpseudopropazine and other preparations for which monoamine oxidase inhibition is claimed, except triparanol
Monobenzone for human therapeutic use, except in preparations containing 2 per cent or less of monobenzone
Moperon
Morphine Antagonists including nalorphine, naloxone and levallorphan
Mustine and other substances structurally derived therefrom with cytotoxic properties when used for therapeutic purposes
Nalidixic acid
Neostigmine
Nicocodine when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of nicocodine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of nicocodine;
except in preparations specified in Schedule 2
Nicodocodine when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of nicodocodine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of nicodocodine;
except in preparations specified in Schedule 2
Nicotinyl Alcohol for internal use
Nifenazonc
Nikethamide
Niridazole
Nitrazepam
Nitroscyanate
Nitrous oxide when specifically prepared and packed as a therapeutic agent for the induction of inhalation anaesthesia
Nitrofurantoin and its derivatives for human therapeutic use
Noradrenaline, except in preparations containing 1 per cent or less of noradrenaline
SCHEDULE 4—continued

Norcodeine when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of norcodeine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of norcodeine;
except in preparations specified in Schedule 2
Nortriptyline
Octamylamine
Octatropine
Orciprenaline

Organo-phosphorus Compounds with anticholinesterase activity for human therapeutic use,
except organo-phosphorus compounds specified in Schedule 3
Ornipressin
Orphenadrine
Oxandrolone
Oxazepam
Oxolamine
Oxprenolol
Oxyphenbutazone
Oxyphencyclidine
Oxyphenonium

Pancuronium
Paraldehyde
Pemoline
Pempidine
Pentamethonium
Pentathionate
Pentolinium
Perhexilene
Phenacemide and other substances structurally derived from acetylurea with anticonvulsant properties when used for therapeutic purposes
Phenacetin
Phenazine for internal use
Phenazopyridine
Phenformin
Phenglutarimide
Phenoxybenzamine
Phensuximide and other substances structurally derived from succinamide with anticonvulsant properties when used for therapeutic purposes
Phentermine
Phenthimentonium
Phenyapin
Phenylbutazone
Phenylobromide
Phenylpropanolamine
Phenyltoin and other substances structurally derived from hydantoin with anticonvulsant properties when used for therapeutic purposes
Pholcodine when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of pholcodine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of pholcodine;
except in preparations specified in Schedule 2
Physostigmine
Picrotoxin
Pilocarpine, except in preparations containing 0.025 per cent or less of pilocarpine
Pimozide
Pindolol
Pipenzolate
Piperaldolate
Pipobroman
Pipradrol
SCHEDULE 4—continued

Pituitary, its extracts, its active principles or their synthetic substitutes, except the extracts, active principles and substitutes specified in Schedule 7

Pizotifen
Polymethylene Bistrimethyl Ammonium Compounds
Potassium Perchlorate for therapeutic use
Praziquantel
Pregnenolone Acetate, except in preparations for topical use
Prenylamine
Primidone
Probenecid
Procarbazine
Prochlorperazine
Procyclidine
Prolintane
Propanidid
Propantheline, except in preparations for topical use
Propylhexedrine, except when included in Schedule 3
Prothionamide
Pyridostigmine
Quinethazone
Quinidine
Rauwolfia Serpentina
Salbutamol
Selenium, compounds of, except compounds specified in Schedule 5 or 6
Sex Hormones, natural or synthetic, their substitutes in all preparations, including cosmetics, except—
(a) their derivatives and their substitutes without sex hormonal activity;
(b) testosterone propionate and testosterone dipropionate in preparations specified in Schedule 6
Sodium Cromoglycate
Sparteine
Sulphonamethoxazole and its glycosides
Sulphanilamide and its derivatives, except—
(a) the substances specified in Schedule 6 or 7;
(b) sulphaquinoxaline when incorporated in baits for the destruction of vermin and in animal feedstuffs containing 200 parts per million or less of sulphaquinoxaline

Sulphinpyrazone
Sulphonals and alkyl sulphonals
Sulthiame
Suxamethonium
Tacrine
Terbutaline
Thiambutosine
Thia ouabaine
Thioguanine and other substances structurally derived therefrom with cytotoxic properties when prepared for use for therapeutic purposes
Thioureic acid
Thiouracil and substances structurally derived therefrom with antithyroid properties when prepared for use for therapeutic purposes
Thiouracil for therapeutic use
Thyroid and its extract, and its active principles
Tiemonium
Tigloidine
Timidazole
Tipepidine
Tolazamide
Tolazoline for internal use
Tolbutamide
Tretamine
Triamterene
Triaziquone
Trichloroethylene when specifically prepared and packed as a therapeutic agent for the induction of inhalation anaesthesia
Triclofos, except preparations containing 5 per cent or less of triclofos
Triclamol
Tridihexethyl
Trifluoperidol
Trimetaphan
Trimethoprim
Trimipramine and other compounds structurally derived therefrom by substitution in the side chain
Trimethine
Trioxysalen
Troxidone and other substances structurally derived from oxazolidinone with anticonvulsant properties when prepared for use for therapeutic purposes
Urethane (excluding its derivatives), when prepared for use for therapeutic purposes
Urethanes and Ureides having or purporting to have soporific, hypnotic or narcotic properties, not specifically included in this or any other schedule
Vaccines, sera, toxoids, and antigens prepared for human parenteral use
Vaccines, veterinary live virus
Valnoctamide
Veratrum for therapeutic use
Vinca Alkaloids
Vinyl Ether when specifically prepared and packed as a therapeutic agent for the induction of inhalation anaesthesia
Visnadine
Vitamin A in preparations containing more than 10,000 I.U. per recommended daily dosage for human use
Vitamin D in preparations containing more than 25 micrograms per recommended daily dosage for human use
Xanthine Oxidase Inhibitors including allopurinol
Xanthinol Nicotinate
Xylazine
Yohimbine
Acetic Acid (excluding its salts and derivatives) in preparations containing not more than 80 per cent and not less than 30 per cent of acetic acid, except preparations for therapeutic use.

Acetic Anhydride (excluding its salts and derivatives) in preparations containing not more than 80 per cent and not less than 30 per cent of acetic anhydride, except preparations for therapeutic use.

Acetone when packed in containers of 20 litres or less, except—
(a) in preparations containing 25 per cent or less of acetone;
(b) when packed in containers of 60 mls or less.

Aklomide
Alachlor
Amitrole

Ammonia (excluding its salts and derivatives other than ammonium hydroxide) in preparations containing 5 per cent or less of free ammonia, except—
(a) medicinal preparations for internal use;
(b) in appliances for inhalation in which the substance is absorbed upon an inert solid material;
(c) in preparations containing 0.5 per cent or less of free ammonia.

Ammonium Thiocyanate

Arsenic, organic compounds of, in preparations containing 3 per cent or less of arsenic, when prepared for use as herbicides or defoliants.

Barium Silicofluoride when coated on paper in an amount not exceeding 8 mg per sq. cm.

Bentazone
Benzoyl Peroxide

Borax Hexachloride preparations containing 10 per cent or less of benzene hexachloride.

Boron Compounds when included in soaps, detergents, laundry preparations, cleaning agents and bleaches, except preparations containing 1 per cent or less as boron.

Cadmium Sulphide in preparations containing 2.5 per cent or less of cadmium sulphide for human therapeutic use.

Camphor, except—
(a) in preparations containing 10 per cent or less of camphor;
(b) in camphorated oil.

Captafol
Carbaryl—
(a) when impregnated in plastic resin strip material containing 20 per cent or less of carbaryl;
(b) in preparations containing 10 per cent or less of carbaryl.

Chlordecone in preparations containing 5 per cent or less of chlordecone.

Chlorethalin
Chlorfenac
Chlorfenuron

Chlorinating compounds and bleaches containing more than 4 per cent of available chlorine, except compounds and bleaches specified in Schedule 7 or specified elsewhere in this Schedule.

Chlornidine
Chlororesol
Chloropropylate

Copper Sulphate as such

4-CPA
Cyanatryn
Cyanacrylacid Esters
Cyanuric Acid (excluding its salts and derivatives)
Cyclohexanone Peroxide

3-Cyclohexyl-(6(dimethylamino)-1-methyl-1, 3, 5, 3-triazine 2, 4-(1H, 3H)-dione

2, 4-D
2, 4-DB
2, 4-DES

Dicamba
Dichlone

Dichloroisocyanurates and in preparations containing more than 4 per cent available chlorine.
Dichlorvos when impregnated in plastic resin strip material containing 20 per cent or less dichlorvos and when in aerosol preparations containing 1 per cent or less dichlorvos.
Dicloran
Dicofol
Dicophane in preparations containing 10 per cent or less of dicophane, except preparations for human therapeutic use
Dimethirimol
Dinitramine
Diphenamid
Dodine
Epoxy Resins (Liquid) and all amines and organic anhydrides used as curing agents for epoxy resins
EPTC
Ethephon (excluding its salts and derivatives)
Ether Preparations for use in internal combustion engines
Ethofumesate
Ethoxyquin, except in preparations containing 10 per cent or less of ethoxyquin
N-((1-Ethylpropyl)-3, 4-dimethyl-2, 6-dinitroaniline
Eucalyptus Oil, except in preparations containing 25 per cent or less of eucalyptus oil
Fenbutatin-oxide
Fenoprop
Fenson
Fenthion in preparations containing 20 per cent or less of fenthion when packed in single use containers having a capacity of 0.3ml or less
Flamprop-methyl
Formic Acid (excluding its salts and derivatives)
Fospirate when impregnated in plastic resin strip material containing 20 per cent or less of fospirate
Glyphosate
Hydrocarbons (Liquid) distilling at a temperature lower than 300°C when tested according to method D86-61 of the American Society for Testing and Materials, except liquid hydrocarbons—
(a) specified in Schedule 6 or 7;
(b) packed in containers having a capacity of more than 20 litres;
(c) in substances containing 25 per cent or less of such liquid hydrocarbons;
(d) in solid or semi-solid cleaning and polishing preparations;
(e) in preparations packed in pressurised aerosol containers; or
(f) in adhesives packed in containers each containing 50 grammes or less of adhesive
Hydrochloric Acid (excluding its salts and derivatives) in preparations containing 10 per cent or less of hydrochloric acid (HCl), except—
(a) in preparations containing 0.5 per cent or less of hydrochloric acid (HCl); and
(b) in preparations for therapeutic use
Hydrogen Peroxide (excluding its salts and derivatives), except in preparations containing 6 per cent weight-in-volume (20 vol) or less of hydrogen peroxide
Iodosphenphos
2-Isobutylamino-4-ethylamino-6-methoxy-1, 3, 5-triazine
Kerosene, when packed in containers of 20 litres or less, except in preparations containing 25 per cent or less of kerosene
Levamisole in preparations containing 15 per cent or less of levamisole for the treatment of animals
Lindane in preparations containing 10 per cent or less of lindane
Maldison in preparations containing 10 per cent or less of maldison, except preparations for human therapeutic use
Mancozeb
Maneb
MCPA
MCPB
Me-oprop
Metaldehyde in preparations containing 2 per cent or less of metaldehyde
Methabenzthiazuron
Methazole
Methiocarb in pelleted preparations containing 2 per cent or less of methiocarb when labelled and packed for the control of snails and slugs
Methoxychlor
Methylated Spirit when packed in containers of 20 litres or less, except in preparations containing 25 per cent or less of methylated spirit

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SCHEDULE 5—continued

Methylene Chloride, except when used in aerosols
Methyl Ethyl Ketone when packed in containers of 20 litres or less, except in preparations containing 25 per cent or less of methyl ethyl ketone
Methyl Ethyl Ketone Peroxide
Methyl-iso-butyl Ketone when packed in containers of 20 litres or less, except in preparations containing 25 per cent or less of methyl-iso-butyl ketone
Metiram
Metribuzin
Metizine
Mineral Turpentine when packed in containers of 20 litres or less, except in preparations containing 25 per cent or less of mineral turpentine
NAA
Naled when impregnated in plastic resin strip material containing 20 per cent or less of naled
Naphthalene as such
Nitric Acid (excluding its salts and derivatives) in preparations containing 100 grams per kilogram or less of nitric acid, except preparations containing 5 grams per kilogram or less of nitric acid
Norbormide
Organo-tin Compounds not elsewhere specified in this Schedule in preparations containing 1 per cent or less of such compounds
Oxycarboxin
Oxythioquinox
Paradichlorobenzene
Pebulate
Petrol when packed in containers of 20 litres or less, except preparations containing 25 per cent or less of petrol
Potassium Hydroxide (excluding its salts and derivatives) in preparations containing 5 per cent or less of potassium hydroxide, except—
(a) in preparations containing 0.5 per cent or less of potassium hydroxide;
(b) in accumulators and batteries
Prometryn
Propanil
Propionic Acid (excluding its salts and derivatives) in preparations containing not more than 80 per cent and not less than 30 per cent of propionic acid, except preparations for therapeutic use
Propoxur in dust preparations containing 3 per cent or less of propoxur
Prynachlor
Pyrethrins and related compounds, except in preparations containing 10 per cent or less of such compounds
N-3-Pyridylmethyl-N'-p-nitrophenylurea in preparations containing 10 per cent or less of N-3-pyridylmethyl-N'-p-nitrophenylurea
Quaternary ammonium compounds and in preparations containing more than 10 per cent quaternary ammonium compounds, except when specified in any of Schedules 1 to 4 or 6 to 8
Quintozene
Salicylanilide
Selenium Sulphide in preparations containing 2.5 per cent or less of selenium sulphide for therapeutic use
Sodium Chlorate in preparations containing 10 per cent or less of sodium chlorate
Sodium Hydrogen Sulphate
Sodium Hydroxide (excluding its salts and derivatives) in preparations containing 5 per cent or less of sodium hydroxide, except in preparations containing 0.5 per cent or less of sodium hydroxide
Sodium Nitrite, except—
(a) in preparations containing 1 per cent or less of sodium nitrite;
(b) in preparations for therapeutic use
Styrene (excluding its derivatives) when packed in containers of 20 litres or less
2, 3, 6-TBA
TDE in preparations containing 10 per cent or less TDE
Terbuthylazine
Terbutryn
2-Tert-butylamino-4-ethylamino-6-methoxy-1, 3, 5-triazine
Tetrachlorvinphos
Tri-allate
1, 1, 1-Trichloroethane when packed in containers of 20 litres or less, except—
(a) in preparations containing 25 per cent or less of 1, 1, 1-trichloroethane;
(b) when packed in aerosols for purposes other than therapeutic use;
(c) when packed in containers of 50 mls or less
Trichloroisocyanuric Acid when compressed in block form for use in swimming pools
Trietazine
Turpentine Oil when packed in containers of 20 litres or less, except in preparations containing
25 per cent or less of turpentine oil
Vernolate
White Spirit when packed in containers of 20 litres or less, except in preparations containing
25 per cent or less of white spirit
Zinc Pyrithione in preparations containing 2 per cent or less of zinc pyrithione
Zineb
Ziram
Acephate
Acetic Acid (excluding its salts and derivatives), except—
   (a) in preparations containing 80 per cent or less of acetic acid, and
   (b) in preparations for therapeutic use
Acetic Anhydride (excluding its salts and derivatives), except—
   (a) in preparations containing 80 per cent or less of acetic anhydride;
   (b) in preparations for therapeutic use
Allidochlor
Alpha-chloralose in preparations containing 5 per cent or less of alphachloralose when prepared for use as a rodenticide
Ametryn
Amidithion
Amines (Aromatic), including phenylene diamine, toluene diamine and all other aromatic amines, when prepared in hairdyes
2-Aminobutane
Aminocarb in preparations containing 25 per cent or less of aminocarb
2-Amino-5-diethylamine toluene
2-Amino-5-N-ethyl-N-(B hydroxyethyl) amino toluene
2-Amino-5-N-ethyl-N-(B methane sulphonamide ethyl) amino toluene
2-Amino-5-N-ethyl-N-B methoxyethyl amino toluene di-p-toluene
Amitraz
Ammonia (excluding its salts and derivatives other than ammonium hydroxide), except—
   (a) in preparations containing 5 per cent or less of free ammonia;
   (b) in medicinal preparations for internal use;
   (c) in appliances for inhalation in which the substance is absorbed in an inert solid material
Aniline (excluding its salts and derivatives), except in preparations containing 1 per cent or less of aniline
Arsenical Preparations, being ant poisons containing 0.5 per cent or less of arsenic trioxide
Arsenic, compounds of, except—
   (a) compounds specified in Schedules 5 and 7;
   (b) compounds in animal feedstuffs containing 75 parts per million or less of arsenic (As);
   (c) compounds prepared for human therapeutic use
Arsenic, organic compounds of, when prepared for use as herbicides or defoliants, except preparations specified in Schedule 5
Azobenzene
Bacitracin in animal feedstuff premixes for growth promotion purposes containing concentrations greater than 50 parts per million but not more than 20,000 parts per million of the total antibiotic principle
Barban
Barium, salts of, except—
   (a) barium sulphate;
   (b) paint containing barium metaborate;
   (c) salts of barium specified in Schedule 5
Bensulide
Bendiocarb in wettable powders containing 80 per cent or less of bendiocarb and when packed and labelled for the control of cockroaches in containers or primary packs containing not less than 100 g of bendiocarb
Benquinox
Benzy1 Penicillin, including procaine penicillin, in preparations—
   (a) for intramammary infusion in animals when suitably coloured with Brilliant Blue FCF or other approved colour as a marker and when packed in applicator devices specially designed for the purpose;
   (b) in animal feedstuff premixes for growth promotion purposes containing concentrations greater than 50 parts per million but not more than 20,000 parts per million of the total antibiotic principle
Beryllium
Benzene Hexachloride, except in preparations containing 10 per cent or less of benzene hexachloride
Binapacryl
Bithionol in preparations for treatment of animals
SCHEDULE 6—continued

Boron Compounds, except—
(a) compounds prepared for human therapeutic or cosmetic use;
(b) preparations containing 1 per cent or less as boron;
(c) compounds specified in any of Schedules 1 to 5, 7 and 8.

Bromoform, except when prepared for therapeutic use
Bromophos-ethyl
Bromoxynil
Brotianide
Butacarb
2-Butoxy-2'-thiocyano-diethyl Ether
Butynorate
Cadmium, compounds of, except compounds specified in Schedule 5
Cambendazole
Camphenechlor
Carbaryl, except in a form or preparation specified in Schedule 5
Carbon Disulphide
Chloramphenicol prepared for use on animals for the topical treatment of foot rot and for ocular use
Chlordane
Chlordecone, except in preparations containing 5 per cent or less of chlordecone
Chlorfenethol
Chlormequat
Chlorodimeform
Chloromethiuron
Chlorophacinone
Chloropicrin in preparations containing 5 per cent or less of chloropicrin
Chlorpyrifos
Chlortetracycline in preparations—
(a) for topical application to animals for ocular use only;
(b) for intramammary infusion in animals when suitably coloured with Brilliant Blue FCF or other approved colour as a marker and when packed in applicator devices specially designed for the purpose

Chlorthiamid
Chromates and Dichromates
Chromium Trioxide (excluding its salts and derivatives)
Coumaphos in preparations containing 5 per cent or less of coumaphos
Coumarin Derivatives and phenylindanedione derivatives not specified in any of Schedules 1 to 5, 7 and 8
Coumatetralyl
Crotoxyphos
Crufomate
Cyanazine
Cyhexatin
Dazomet
Demeton-O-methyl and Demeton-S-methyl in preparations containing 50 per cent or less of one or both demeton-O-methyl and demeton-S-methyl
Di-allate
Diazinon
1, 2-Dibromo-3-chloropropane
Dichlofenthion
Dichloroethyl Ether
1, 2-Dichloropropane
1, 3-Dichloropropene
Dichlorvos in preparations containing 50 per cent or less dichlorvos, except preparations specified in Schedule 5
Dicofophane and in preparations containing more than 10 per cent of dicofophane, except preparations for human therapeutic use
Diethylene Dioxide
N, N-Diethyl-p-phenylene diamine
Difenoquat
Dimethanonaphthalene and all substitution and/or addition products of, including aldrin and dieldrin
SCHEDULE 6—continued

Dimethoate
1, 3-Di (methoxycarbonyl)-1-propen-2-yl-dimethyl Phosphate in preparations containing 25 per cent or less of 1, 3-di (methoxycarbonyl)-1-propen-2-yl-dimethyl phosphate

Dimethyl Formamide

Dimethyl Sulphoxide, except when prepared for therapeutic use

Dimetilan in preparations containing 25 per cent or less of dimetilan

Dimetridazole

Dinitrocresols, Dinitrophenols and their homologues in preparations containing 5 per cent or less of such compounds, except preparations for therapeutic use

Dinocap

Dioxacarb

Diphenoxine

Diquat

Disulfiram, except when prepared for therapeutic use

Disulfoton in granular preparations containing 5 per cent or less of disulfoton

Dithianon

Dithiazanine in preparations containing 2 per cent or less of dithiazanine for the treatment of animals

Dithiocarbamates in preparations for agricultural, pastoral or horticultural purposes, except preparations specified in Schedule 5

DSMA in herbicides, except preparations specified in Schedule 5

Endosulfan

Endothal

Epichlorohydrin, except in preparations containing 1 per cent or less of epichlorohydrin

Erythromycin in preparations—
(a) for intramammary infusion in animals when suitably coloured with Brilliant Blue FCF or other approved colour as a marker and when packed in applicator devices specially designed for the purpose;
(b) in animal feedstuff premixes for growth promotion purposes containing concentrations greater than 50 parts per million but not more than 20,000 parts per million of the total antibiotic principle

Ether Solvent

Ethiofencarb

Ethoate-methyl

Ethoprophos in granular preparations containing 10 per cent or less of ethoprophos

5-Ethoxy-3-trichloromethyl-1, 2, 4-thiadiazole

Ethyl Bromide

Ethylene Chlorohydrin

Ethylene Dibromide

Ethylene Dichloride

Ethylene Oxide

Famphur in preparations containing 20 per cent or less of famphur

Fenamidosulf in preparations containing 10 per cent or less of fenamidosulf when labelled and packed as dry seed dressings

Fenamiphos in granular preparations containing 5 per cent or less of fenamiphos

Fenazaflor

Fenchlorphos

Fenitrothion

Fenthiion except when included in Schedule 5

Ferbam

Ferrocyanides and Ferricyanides, except in preparations containing 1 per cent or less of such substances

Flavophospholipol in animal feedstuff premixes for growth promotion purposes containing concentrations greater than 50 parts per million but not more than 20,000 parts per million of the total antibiotic principle

Formaldehyde (excluding its derivatives), except in preparations containing 5 per cent or less of formaldehyde

Formothion

Fospirate, except in a form specified in Schedule 5

HCB

Heptachlor

Hexachlorophane in preparations for the treatment of animals
Hydrazine

Hydrochloric Acid (excluding its salts and derivatives), except in preparations containing 10 per cent or less by weight of hydrochloric acid (HCl)

Hydrofluoric acid and Hydrosilicofluoric Acid and other fluorine compounds, except—
(a) in preparations for human therapeutic use;
(b) in dentifrices containing 0.5 per cent or less of fluoride ion;
(c) in preparations containing 3 per cent or less of sodium fluoride or sodium silico-fluoride prepared for use as preservatives;
(d) in compounds specified in Schedule 7;
(e) in substances containing 15 parts per million or less of fluoride ion

Hygromycin in animal feedstuff premixes for use as an anthelmintic containing concentrations greater than 20 parts per million but not more than 20,000 parts per million of hygromycin

Iodine (excluding its salts and derivatives) in liquid preparations containing 2.5 per cent or less of iodine

Iodophors containing 2.5 per cent or less of available iodine

Ioxynil

Iron Compounds, in preparations for the treatment of animals

Isocyanates, free organic

Laurylisouquinolinium Bromide

Lead Compounds, except—
(a) in preparations for therapeutic or cosmetic use;
(b) in pencils containing 0.5 per cent or less of lead in the surface coating or 0.25 per cent or less of lead in the cores;
(c) in finger colours, show card colours, poster paints, school pastels or crayons containing 0.025 per cent or less of lead

Lindane, except in preparations containing 10 per cent or less of lindane

Maldison, except—
(a) in preparations for human therapeutic use;
(b) in preparations specified in Schedule 5

Mebendazole in preparations for the treatment of animals

Meclofenamic Acid in preparations for the treatment of animals

Menazon

Mercuric Iodide in preparations for use for agricultural, industrial, pastoral or horticultural purposes

Mercuric Thiocyanate in preparations for use for photographic purposes

Mercurous Chloride, except in preparations for internal use

Mercury, organic compounds of, in preparations for use for agricultural, pastoral or horticultural purposes, except preparations specified in Schedule 7

Metaldehyde, except in preparations containing 2 per cent or less of metaldehyde

Metham-sodium

Methyl Alcohol (excluding its salts and derivatives), except in methylated spirit

Methyl Chloride

Methyl Isothiocyanate

1-(B Methyl sulphonamide ethyl)-2-amino-3-N, N-diethylamino benzene

Molinate

Naled, except in a form specified in Schedule 5

Naphthalophos when specifically prepared and packed for use as a sheep drench

Neomycin in preparations for topical application to animals for ocular use only

Niclosamide when prepared for the treatment of animals

Nicotine in preparations containing 3 per cent or less of nicotine when labelled and packed for the treatment of animals

Nimidane in preparations containing 25 per cent or less of nimidane

Nithiamide, except in preparations containing 20 per cent or less of nithiamide

Nitric Acid (excluding its salts and derivatives), except in preparations containing 100 grams per kilogram or less of nitric acid

Nitrobenzene, except—
(a) in solid or semi-solid polishes;
(b) in soaps containing 1 per cent or less of nitrobenzene;
(c) in preparations containing 0.1 per cent or less of nitrobenzene

Nitrophenols, ortho, meta and para
Nitroxynil

Oleandomycin in animal feedstuff premixes for growth promotion purposes containing concentrations greater than 50 parts per million but not more than 20,000 parts per million of the total antibiotic principle

Omethoate in preparations containing 50 per cent or less of omethoate

Organo-tin Compounds not specified in any of Schedules 1 to 5, 7 and 8

Orthodichlorobenzene

Oxalic Acid (excluding its salts and derivatives) and soluble oxalates

Oxantel Pamoate for the treatment of animals

Oxyclozanide

Oxycodone in preparations—
  (a) for topical application to animals for ocular use only;
  (b) for intramammary infusion in animals when suitably coloured with Brilliant Blue FCF or other approved colour as a marker and when packed in applicator devices specially designed for the purpose

Paraquat in granular preparations containing 3 per cent or less of paraquat

Parbendazole

Pentachlorophenol, except in preparations containing 0.5 per cent or less of pentachlorophenol

Permanganates

Phenkapton in preparations containing 50 per cent or less of phenkapton

Phenol, creosote and any homologue of phenol boiling below 220°C, except—
  (a) in preparations containing 3 per cent or less by weight of such substances or homologues;
  (b) in preparations for therapeutic use

Phenoxyphene and Phenethicillin in preparations for intramammary infusion in animals when suitably coloured with Brilliant Blue FCF or other approved colour as a marker and when packed in applicator devices specially designed for the purpose

Phtosalone

Phosmet

Phosphides, Metallic

Phosphoric Acid (excluding its salts and derivatives), except in preparations containing 5 grams per kilogram or less of phosphoric acid (H₃PO₄)

Phosphorus, yellow (excluding its salts and derivatives) in preparations containing 0.5 per cent or less of free phosphorus

Picric Acid (excluding its derivatives), except in preparations containing 5 per cent or less of picric acid

Pindone

Piperophos

Pirimicarb

Pirimiphos-methyl

Potassium Bromate, except in preparations containing 0.5 per cent or less of potassium bromate

Potassium Cyanate

Potassium Hydroxide (excluding its salts and derivatives), except in preparations containing 5 per cent or less of potassium hydroxide

Promecarb in preparations containing 50 per cent or less of promecarb

Propachlor

Propargite

Propionic Acid (excluding its salts and derivatives), except—
  (a) in preparations containing 80 per cent or less of propionic acid;
  (b) for therapeutic use

Propoxur, except in dust preparations containing 3 per cent or less of propoxur

Pyrazophos

N-3-Pyridylmethyl-N'-p-nitrophenylurea except in preparations specified in Schedule 5

Rafoxanide

Selenium, compounds of, in preparations containing 2.5 per cent or less—
  (a) when packed and labelled for the blueing of gun barrels;
  (b) when packed and labelled for photographic purposes

Sodium Bromate, except in preparations containing 0.5 per cent or less of sodium bromate

Sodium Chlorate, except in preparations containing 10 per cent or less of sodium chlorate
SCHEDULE 6—continued

Sodium Hydroxide (excluding its salts and derivatives), except in preparations containing 5 per cent or less of sodium hydroxide

Streptomycin in preparations for intramammary infusion in animals when suitably coloured with Brilliant Blue FCF or other approved colour as a marker and when packed in applicator devices specially designed for the purpose

Strychnine in preparations containing 1 per cent or less of strychnine when prepared for the destruction of vermin

Sulfathiazole

Sulfanilamide in preparations for the treatment of animals, except animal feedstuffs containing 200 parts per million or less of sulphaquinoxaline

Sulphuric Acid (excluding its salts and derivatives), except—

(a) in accumulators, batteries and fire extinguishers;

(b) in preparations containing 5 grams per kilogram or less of sulphuric acid ($\text{H}_2\text{SO}_4$)

2, 4, 5-T

TCA (excluding its salts and derivatives)

TCMTB (2-[thiocyanomethylthio] benzothiazole)

TDE, except in preparations containing 10 per cent or less of TDE

Temephos

Terpenes, chlorinated

Testosterone Propionate and Testosterone Dipropionate in preparations for the treatment of animals

Tetrachloroethylene, except—

(a) when prepared for use for the treatment of humans and for the treatment of animals;

(b) when packed in containers of 50 mls or less

Tetracycline in preparations—

(a) for topical application to animals for ocular use only;

(b) for intramammary infusion in animals when suitably coloured with Brilliant Blue FCF or other approved colour as a marker and when packed in applicator devices specially designed for the purpose

Tetradifon

Tetramisole including Levamisole, in preparations for the treatment of animals, other than preparations specified in Schedule 5

Thiazafluron

Thiometon

Thiourea, except when prepared for therapeutic use

Thiram

Toluene and Xylene (excluding their derivatives), when packed in containers of 20 litres or less, except—

(a) in preparations containing 50 per cent or less of one or both toluene and xylene when tested according to method D1019-67 of the American Society for Testing and Materials;

(b) when packed in containers of 50 mls or less

Triadimefon

S,S,S-Tributylphosphorothioate

Trichloroethylene, except—

(a) when specifically prepared for medicinal purposes;

(b) when packed in containers of 50 ml or less

Trichlorophenol

Trichlorphon

Triethyl Phosphate

Tylosin and its salts in animal feedstuff premixes for growth promotion purposes containing concentrations greater than 50 parts per million but not more than 20,000 parts per million of the total antibiotic principle

Vamidothion

Virginiamycin in animal feedstuff premixes for growth promotion purposes containing concentrations greater than 50 parts per million but not more than 20,000 parts per million of the total antibiotic principle

Warfarin, except when prepared for therapeutic use

Zinc Chloride, except in preparations containing 5 per cent or less of zinc chloride
Zinc p-Phenolsulphonate, except in preparations containing 5 per cent or less of zinc p-phenolsulphonate
Zinc Sulphate, except in preparations containing 5 per cent or less of zinc sulphate
SCHEDULE 7 Section 3

Acrolein
Aldicarb
Allyl Alcohol
Aminocarb, except in preparations containing 25 per cent or less of aminocarb
4-Aminopyridine
Amion
ANTU
Arsenical Preparations, being ant poisons or prepared for use as liquid weedkillers or as liquid preparations for the destruction of termites, except compounds specified in Schedule 5 or 6
Azinphos-ethyl
Azinphos-methyl
Bendiocarb, except in preparations specified in Schedule 6
Benzene (excluding its derivatives), except—
   (a) preparations containing 1 per cent v/v or less of benzene;
   (b) petrol containing 5 per cent v/v or less of benzene
Beta-hydroxyethylhydrazine
Bromocriptine
Carbofuran
Carbon Tetrachloride
Carbophenothion
Chlorfenavinphos
Chlorine (excluding its salts and derivatives)
Chloropicrin, except in preparations containing not more than 5 per cent of chloropicrin
5-Chloro-3-methyl-4-nitropyrazole
Clomiphene and other products specifically prepared to stimulate ovulation
Coumaphos, except in preparations containing 5 per cent or less of coumaphos
Cyclofenil
Demeton
Demeton-O-methyl and Demeton-S-methyl, except in preparations specified in Schedule 6
Dichlorvos, except in preparations specified in Schedule 5 or 6
Dicrotophos
Dimefox
1, 3-Di(methoxycarbonyl)-1-propen-2-yl-dimethyl Phosphate, except in preparations specified in Schedule 6
Dimetilan, except in preparations specified in Schedule 6
Dinitrocresols, Dinitrophenols and their homologues, except—
   (a) in preparations containing not more than 5 per cent of those compounds or their homologues;
   (b) preparations for therapeutic use
Dioxathion
Disulphoton, except in granular preparations containing 5 per cent or less of disulphoton
Ethion
Ethoprophos, except in preparations specified in Schedule 6
Ethoxyethyl Mercury Chloride
Ethyl Mercury Chloride
Famphur, except in preparations containing not more than 20 per cent of famphur
Fenaminosulf, except in preparations specified in Schedule 6
Fenamiphos, except in preparations specified in Schedule 6
Fensulphothion
Fenthion-ethyl
Fluoroacetic Acid
Formetanate
Hydrocyanic Acid and Cyanides, except—
   (a) in preparations containing the equivalent of 0.15 per cent or less of hydrocyanic acid;
   (b) in preparations for therapeutic use
Isocarbophos
Leptophos
Mazidox
Mecarbam
Mercuric Chloride in preparations for use for agricultural, industrial, pastoral or horticultural purposes
Methamidophos
Methidathion
Methomyl
Methyl Bromide
Mevinphos
Mipafox
Monocrotophos
Naphthalophos, except in preparations specified in Schedule 6
Nicotine, except—
  (a) in preparations specified in Schedule 6;
  (b) in tobacco
Nimidane, except in preparations specified in Schedule 6
Nitrilotriacetic Acid and all preparations containing nitrilotriacetic acid when labelled and
packed for use as detergents or washing preparations
Omethoate, except in preparations specified in Schedule 6
Paraquat, except in preparations specified in Schedule 6
Parathion
Parathion-methyl
Phenkapton, except in preparations containing 50 per cent or less of phenkapton
Phorate
Phosfolan
Phosphamidon
Polychlorinated Biphenyls
Promecarb, except in preparations containing 50 per cent or less of promecarb
Prostaglandins
Schradan
Silver Sulphadiazine
Sulfotep
TEPP
Tetrachloroethane
Thallium
o-Tolidine
Trichloroisocyanuric Acid and its salts, except—
  (a) in preparations containing 4 per cent or less of available chlorine;
  (b) in a form specified in Schedule 5
Vinyl Chloride
SCHEDULE 8 Section 3

Acetorphine
Acetyldihydrocodeine, except when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of acetyldihydrocodeine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of acetyldihydrocodeine

Acetylmethadol
Allylprodine
Alphacetylmethadol
Alphameprodine
Alphaprodine
Amphetamine
Anileridine
Benzethidine
Benzylmorphine
Betacetylmethadol
Betameprodine
Betamethadol
Betaprodine
Bezitramide
Clonitazene
Cocaine
Coca Leaf

Codeine, except when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of codeine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of codeine

Codeine-N-oxide
Codoxime

Concentrate of Poppy Straw (the material arising when poppy straw has entered into a process for concentration of its alkaloids)

Dexamphetamine
Dextromoramide
Diampromide
Diethylthiambutene

Difenoxin, other than preparations containing per dosage unit, not more than 0.5 mg of difenoxin (base) and a quantity of atropine sulphate equivalent to at least 5 per cent of the dose of difenoxin

Dihydrocodeine, except when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of dihydrocodeine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of dihydrocodeine

Dihydromorphine
Dimenoxadol
Dimepheptanol
Dimethylthiambutene
Dioxyphethyl Butyrate

Diphenoxylate, other than preparations containing per dosage unit, not more than 2.5 mg of diphenoxylate (base) and a quantity of atropine sulphate equivalent to at least 1 per cent of the dose of diphenoxylate

Dipipanone
Drotebanol
Econamine
Ethylmethylthiambutene

Ethylmorphine, except when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of ethylmorphine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of ethylmorphine

Authorised by the ACT Parliamentary Counsel—also accessible at www.legislation.act.gov.au
Etonitazene
Etorphine
Etoxeridine
Fentanyl
Furethidine
Hydrocodone
Hydromorphinol
Hydromorphone
Hydroxypethidine
Isomethadone
Levorphan
Levorphan
Levophenacylmorphan
Levorphan
Metazocine
Methadone
Methadone intermediate
Methaqualone
Methylnalorphine
Methyldesorphine
Methyldihydromorphine
Methylphenidate
1-Methyl-4-phenylpiperidine-4-carboxylic Acid
Metopon
Moramide intermediate
Morpheridine
Morphine
Morphine Methobromide and other pentavalent nitrogen morphine derivatives
Morphine-N-oxide
Morphine
Nicocodine, except when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of nicocodine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of nicocodine
Nicodicodine, except when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of nicodicodine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of nicodicodine
Nicomorphine
Noracymethadol
Norcodeine, except when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of norcodeine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of norcodeine
Norlevorphan
Normethadone
Normorphine
Norpipanone
Opium
Oxycodone
Oxymorphone
Pentazocine
Pethidine
Pethidine intermediate A
Pethidine intermediate B
Pethidine intermediate C
Phenadoxone
Phenampromide
Phenazocine
Phencyclidine
Phenmetrazine
Phenomorphan
Phenoperidine
Pholcodine, except when compounded with one or more other medicaments—
(a) in divided preparations containing not more than 100 mg of pholcodine per dosage unit; or
(b) in undivided preparations with a concentration of not more than 2.5 per cent of pholcodine
Piminodine
Pirimethamine
Proheptazine
Properidine
Propiram
Racemethorphan
Racemoramide
Racemorphan
Thebacon
Thebaine
Trimeperidine
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### WARNING STATEMENTS TO BE INCLUDED IN LABELS ON SCHEDULED SUBSTANCES

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<tr>
<th>Item No.</th>
<th>Substance</th>
<th>Warning statement</th>
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<tbody>
<tr>
<td>1</td>
<td>Benzoyl Peroxide</td>
<td>Avoid contact with the skin and eyes</td>
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<tr>
<td></td>
<td>5-Chloro-3-methyl-4-nitropyrazole</td>
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<td>Cyclohexanone Peroxide</td>
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<td>Hydrofluoric acid, hydrosilicofluoric acid, their salts and other fluorine</td>
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<tr>
<td></td>
<td>compounds</td>
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<td>Oxalic acid and metallic oxalates</td>
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<tr>
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<td>Oxathioquinox</td>
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<td>Phenol and any homologue of phenol boiling below 220°C</td>
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<tr>
<td></td>
<td>Sodium chlorate</td>
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<td>Sulphuric acid</td>
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<td>Zinc chloride</td>
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<td>2</td>
<td>Acrolein</td>
<td>Avoid contact with skin and eyes and avoid breathing its</td>
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<td></td>
<td>Amidothion</td>
<td>dust (or vapour)</td>
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<td>Aminocarb</td>
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<td></td>
<td>Aniline</td>
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<td>Arsenic, organic compounds when prepared for use as herbicides and</td>
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<td></td>
<td>defoliants</td>
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<td>Azoobenzene</td>
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<td>Chlorinating compounds and bleaches</td>
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<td>Chlorpyrifos</td>
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<td>Chromates and dichromates of alkali metals and ammonium</td>
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<td>Dicophane, except for human therapeutic use</td>
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Authorised by the ACT Parliamentary Counsel—also accessible at www.legislation.act.gov.au
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<td>Diethylene dioxide</td>
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<td>Arsenic, organic compounds, when prepared</td>
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<td>for use as herbicides or defoliants</td>
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<td></td>
<td>DSMA</td>
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<td>Endothal</td>
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<td>Insecticidal preparations</td>
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<td>Liquid epoxy resins and all amines and</td>
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<td>organic anhydrides used as curing agents for</td>
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<td>epoxy resins</td>
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<td>7</td>
<td>Asthma sprays containing adrenaline,</td>
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<tr>
<td></td>
<td>natural or synthetic, its salts, noradrenaline</td>
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<tr>
<td></td>
<td>and substances structurally derived from</td>
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<tr>
<td></td>
<td>substitution in the amine group, and the salts</td>
</tr>
<tr>
<td></td>
<td>of such substances and of noradrenaline</td>
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<tr>
<td>8</td>
<td>8-Hydroxyquinoline, its derivatives and their</td>
</tr>
<tr>
<td></td>
<td>salts when prepared for internal use</td>
</tr>
</tbody>
</table>

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<table>
<thead>
<tr>
<th>Item No.</th>
<th>Substance</th>
<th>Warning statement</th>
</tr>
</thead>
<tbody>
<tr>
<td>9</td>
<td>Amines, aromatic, including phenylene diamine, tolune diamine and other aromatic amines when used in hair dyes</td>
<td>Warning—this product contains ingredients which may cause skin irritation of certain individuals and a preliminary test according to accompanying directions should first be made. This product must not be used for dyeing the eyelashes or eyebrows; to do so may be injurious to the eye.</td>
</tr>
<tr>
<td>10</td>
<td>Antibiotic preparations for intramammary treatment of animals</td>
<td>Warning—milk from animals treated with this preparation is unfit for human consumption and must be discarded for (here state number) hours following the cessation of treatment to ensure that the milk is free from residues.</td>
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<td>11</td>
<td>Chlortetracycline in preparations for topical application to animals for ocular use</td>
<td>Warning—should not be used for human beings. For animal treatment only.</td>
</tr>
<tr>
<td></td>
<td>Neomycin in preparations for topical application to animals for ocular use</td>
<td></td>
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<tr>
<td></td>
<td>Oxytetracycline in preparations for topical application to animals for ocular use</td>
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<td></td>
<td>Tetracycline in preparations for topical application to animals for ocular use</td>
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<tr>
<td></td>
<td>Sulphanilamide in preparations for the treatment of animals, except in animal feed-stuffs containing 200 parts per million or less of sulphaquinoxaline</td>
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<tr>
<td></td>
<td>Testosterone propionate and testosterone dipropionate in preparations for the treatment of animals</td>
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<tr>
<td>12</td>
<td>Antibiotic premixes for growth promotion purposes</td>
<td>The concentration of antibiotics in the feed as given to stock should not exceed 100 parts per million of the active antibiotic principle.</td>
</tr>
<tr>
<td>13</td>
<td>Dichlorvos, when impregnated in plastic resin strip material containing 20 per cent or less of dichlorvos</td>
<td>Do not use in food cupboards. Do not use in nurseries and sick rooms where people may be continuously exposed.</td>
</tr>
<tr>
<td>14</td>
<td>Hexachlorophane, in preparations for skin cleansing purposes containing 3 per cent or less of hexachlorophane</td>
<td>For external washing only. Rinse skin thoroughly after use.</td>
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<tr>
<td>15</td>
<td>Benzene</td>
<td>If you can smell the vapour it is harmful to health on prolonged exposure.</td>
</tr>
<tr>
<td></td>
<td>Carbon tetrachloride</td>
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<tr>
<td></td>
<td>Tetrachloroethane</td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>Glazing preparations containing lead compounds</td>
<td>Unless adequately fired, utensils glazed with this preparation must not be used as containers for food or beverages; to do so may cause lead poisoning.</td>
</tr>
<tr>
<td>17</td>
<td>Antihistamine substances</td>
<td>This medication may cause drowsiness. If affected do not drive a vehicle or operate machinery. Avoid alcohol.</td>
</tr>
<tr>
<td>18</td>
<td>Trichloroisocyanuric acid and its salts</td>
<td>Highly reactive oxidizing chlorine compound may cause fire or explosion or produce severe burns. Do not allow to get damp. Store under cover in a dry, clean, well ventilated place. Do not allow to come into contact with acids, reducing agents, ammonium compounds, wood shavings, saw dust, papers, fabric, petrol, kerosene or other combustible material.</td>
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<tr>
<td>Item No.</td>
<td>Substance</td>
<td>Warning statement</td>
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<tr>
<td>19</td>
<td>Captafol</td>
<td>Warning—this product contains ingredients which may cause skin irritation in certain individuals. Avoid contact with skin and eyes and avoid breathing its dust</td>
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</tbody>
</table>
**FIRST AID DIRECTIONS**

**PART I**

<table>
<thead>
<tr>
<th>Paragraph</th>
<th>Description</th>
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<tbody>
<tr>
<td>a</td>
<td>If poisoning occurs, contact a doctor or Poisons Information Centre.</td>
</tr>
<tr>
<td>b</td>
<td>If swallowed, induce vomiting. Use Ipecac Syrup (APF) if available.</td>
</tr>
<tr>
<td>c</td>
<td>If swallowed, do NOT induce vomiting. Give plenty of water or milk.</td>
</tr>
<tr>
<td>d</td>
<td>Avoid giving milk or oils.</td>
</tr>
<tr>
<td>e</td>
<td>Avoid giving alcohol.</td>
</tr>
<tr>
<td>f</td>
<td>If skin contact occurs, remove contaminated clothing and wash skin thoroughly.</td>
</tr>
<tr>
<td>g</td>
<td>Remove from contaminated area. Apply artificial respiration if not breathing.</td>
</tr>
<tr>
<td>h</td>
<td>If swallowed, induce vomiting. Use Ipecac Syrup (APF) if available. Give one Atropine tablet every quarter hour until dryness of the mouth occurs. If poisoned by skin absorption, remove contaminated clothing and wash skin thoroughly.</td>
</tr>
<tr>
<td>i</td>
<td>If poisoning occurs get to a doctor or hospital quickly. If swallowed induce vomiting. Use Ipecac Syrup (APF) if available.</td>
</tr>
<tr>
<td>j</td>
<td>If swallowed, give vegetable oil or milk and induce vomiting. Use Ipecac Syrup (APF) if available. If spill on skin, flood area with vegetable oil or water.</td>
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<tr>
<td>k</td>
<td>If breathing, crush one amyl nitrite ampoule in handkerchief and hold under patient's nose for 1 or 2 seconds. Repeat up to 5 times at intervals of one minute. If not breathing wipe patient's lips and apply artificial respiration.</td>
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<td>l</td>
<td>Give activated charcoal and keep patient quiet, in a dark place if possible.</td>
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**PART II**

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<td>Tetrachloroethane</td>
<td>a, b, e, f, g</td>
</tr>
<tr>
<td>Tetrachloroethylene</td>
<td>a, b, d, e, f, g</td>
</tr>
<tr>
<td>Tetrachlorvinphos</td>
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<tr>
<td>Tetradifon</td>
<td></td>
</tr>
<tr>
<td>Thallium</td>
<td>a, b</td>
</tr>
<tr>
<td>Thiazalluron</td>
<td></td>
</tr>
<tr>
<td>Thiomethon</td>
<td>a, b, h</td>
</tr>
<tr>
<td>Thiourea</td>
<td></td>
</tr>
<tr>
<td>Thiram</td>
<td>a, b</td>
</tr>
<tr>
<td>Tin, organic compounds</td>
<td>a, b, f</td>
</tr>
<tr>
<td>o-Tolidine</td>
<td>a, b</td>
</tr>
<tr>
<td>Toluene</td>
<td></td>
</tr>
<tr>
<td>Toxaphene</td>
<td>a, b, f</td>
</tr>
<tr>
<td>Triadimefon</td>
<td></td>
</tr>
<tr>
<td>Tricloxane</td>
<td>a, b, e</td>
</tr>
<tr>
<td>S,S,S-Tributylphosphorothiolate</td>
<td></td>
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<tr>
<td>1,1,1-Trichloroethane</td>
<td>a, h</td>
</tr>
<tr>
<td>Trichloroethylene</td>
<td>a, d, e, f, g</td>
</tr>
<tr>
<td>Trichloroisocyanuric acid</td>
<td>a, c, f</td>
</tr>
<tr>
<td>Trichlorophenol</td>
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<tr>
<td>Trichlorophene</td>
<td></td>
</tr>
<tr>
<td>Trietazine</td>
<td>a, h</td>
</tr>
<tr>
<td>Triethyl phosphate</td>
<td>a, b</td>
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<tr>
<td>Turpentine oil</td>
<td>a, b</td>
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<tr>
<td>Vamidothion</td>
<td>a, h</td>
</tr>
<tr>
<td>Vernolate</td>
<td>a, b</td>
</tr>
<tr>
<td>Warfarin</td>
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</tr>
<tr>
<td>White spirit</td>
<td>a, c</td>
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<tr>
<td>Xylene</td>
<td></td>
</tr>
<tr>
<td>Xylenols</td>
<td>a, c, f</td>
</tr>
<tr>
<td>Zinc chloride</td>
<td></td>
</tr>
<tr>
<td>Zinc p-phenolsulphonate</td>
<td>a</td>
</tr>
<tr>
<td>Zinc pyrithione</td>
<td>a, b</td>
</tr>
<tr>
<td>Zinc sulphate</td>
<td></td>
</tr>
<tr>
<td>Zineb</td>
<td>a, c</td>
</tr>
<tr>
<td>Ziram</td>
<td>a, b, f</td>
</tr>
</tbody>
</table>

Columns 1 and 2 represent the Substance of Part I and the Paragraphs of Part I, respectively.
Prohibited Substances

4-Bromo-2,5-dimethoxyamphetamine
4-Bromo-3,5-dimethoxyamphetamine
3-Bromo-4-methoxyamphetamine
4-Bromo-3-methoxyamphetamine
Bufotenine
Cannabis
Cannabis resin
Desomorphine
Diethyltryptamine
2,4-Dimethoxyamphetamine
2,5-Dimethoxyamphetamine
3,4-Dimethoxyamphetamine
3,4-Dimethoxy-5-ethoxyamphetamine
2,5-Dimethoxy-4-ethoxyamphetamine
4,5-Dimethoxy-2-ethoxyamphetamine
2,5-Dimethoxy-4-methylamphetamine
2,3-Dimethoxy-4,5-methylenedioxyamphetamine
2,5-Dimethoxy-3,4-methylenedioxyamphetamine
3,4-Methoxyphenylethylamine
Dimethyltryptamine
4,5-Ethylidenedioxy-3-methoxyamphetamine
Heroin
Ketobemidone
Lysergic Acid
Lysergide
Mesaline, and any substance structurally derived from methoxyphenylethylamine, except methoxyphenamine
4-Methoxyamphetamine
2-Methoxy-3,4-methylenedioxyamphetamine
2-Methoxy-4,5-methylenedioxyamphetamine
3-Methoxy-4,5-methylenedioxyamphetamine
4-Methoxy-2,3-methylenedioxyamphetamine
2-Methoxy-3,4-methylenedioxyphenylethylamine
3-Methoxy-4,5-methylenedioxyphenylethylamine
4-Methoxyphenylethylamine
3,4-Methylenedioxyamphetamine
Psilocin
Psilocybin
Tetrahydrocannabinol, being—

(a) a substance having the structural designation 1-hydroxy-3-pentyl-6a, 7, 8, 10a-tetrahydro-6,6,9-trimethyl-6H-dibenzo (b,d) pyran or 2'-hydroxy-4'-pentyl-3,4,5,6-tetrahydro-1,8,8-trimethyl-8H dibenzo (b,d) pyran; or

(b) a 3- or 4'-alkyl homologue within one of those structural designations.

2,3,4,5-Tetramethoxyamphetamine
2,3,4-Trimethoxyamphetamine
2,3,5-Trimethoxyamphetamine
2,3,6-Trimethoxyamphetamine
2,4,5-Trimethoxyamphetamine
2,4,6-Trimethoxyamphetamine
3,4,5-Trimethoxyamphetamine
2,4,5-Trimethoxyphenylethylamine

Authorised by the ACT Parliamentary Counsel—also accessible at www.legislation.act.gov.au
SCHEDULE 13
Section 48
EXEMPT GOODS

Blankets moth-proofed with Dieldrin during manufacture in accordance with the directions of the Commonwealth Scientific and Industrial Research Organization

Ceramics
Electrical components, including electric lamps
Explosives
Glazed pottery
Inorganic pigments not specified in Schedule 6
Matches
Paints
Paper
Photographic paper and film
Timber
Vitreous enamels
Wallboard