

CAYMAN ISLANDS



PHARMACY ACT, 2024

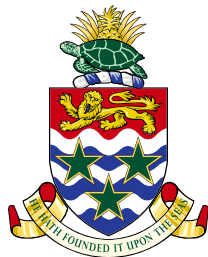
(Act 8 of 2024)

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CAYMAN ISLANDS



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CAYMAN ISLANDS

(Act 8 of 2024)

I Assent,

**Franz Manderson**
Acting Governor

Date: 15th August, 2024

PHARMACY ACT, 2024**(Act 8 of 2024)**

AN ACT TO REPEAL AND REPLACE THE PHARMACY ACT, 1979 AND THE PHARMACY ACT, 1991; TO REGULATE THE IMPORTATION, MANUFACTURE, WHOLESALE, DISPENSING AND PRESCRIBING OF MEDICINES; AND FOR INCIDENTAL AND CONNECTED PURPOSES

ENACTED by the Legislature of the Cayman Islands.

PART 1 - PRELIMINARY**Short title and commencement**

1. (1) This Act may be cited as the Pharmacy Act, 2024.
- (2) This Act comes into force on such date as may be appointed by Order made by the Cabinet and different dates may be appointed for different provisions of this Act and in relation to different matters.

Interpretation

2. In this Act —

“**administer**”, in relation to a substance, means to apply, inject, insert or otherwise introduce the substance to the body of a human being or animal, in its existing state or after it has been dissolved, dispensed in, diluted or mixed with another substance;

“**animal**” means a mammal, bird, reptile, amphibian or fish, but does not include a human being;

“**approved form**” means a form approved by the registrar under section 37;

“**approved medicine**” means a medicine —

- (a) approved by a corresponding authority of a country prescribed by the regulations;
- (b) approved by the Council under section 12; or
- (c) taken to be an approved medicine under section 13 or 14;

“**assemble**”, in relation to a medicine, means —

- (a) to enclose the medicine, with or without other medicines, in a container that is labelled before the medicine is supplied; or
- (b) if the medicine is already enclosed in the container in which it is to be supplied, to label the container before it is supplied;

“**authorized dispenser**” means a person who is permitted to dispense a pharmacy medicine or prescription-only medicine under section 6(1);

“**certified pharmacy**” means a pharmacy that is a certified health care facility under section 5 of the *Health Practice Act (2021 Revision)*;

“**Chief Medical Officer**” means the person appointed to the position of that title in the ministry responsible for health;

“**Chief Nursing Officer**” means the person appointed to the position of that title in the ministry responsible for health;

“**compound**”, in relation to a medicine, means to prepare, mix, alter or assemble multiple ingredients to make the medicine in order to dispense the medicine;

“**constable**” has the meaning assigned by section 2 of the *Police Act (2021 Revision)*;

“**corresponding authority**” means the authority of another country that performs corresponding functions to those of the Council under this Act;

“**Council**” means the Pharmacy Council established under section 21 of the *Health Practice Act (2021 Revision)*;

“**dispense**”, in relation to a medicine, means to supply the medicine, in the course of business or of providing professional services, to a person for the purpose of the medicine being self-administered by the person or administered to an animal;

“**dispensing licence**” means a licence referred to in section 6 and issued under section 9;

“**emergency use authorization**” means an authorization issued by the Cabinet under section 14;



“**exceptional use authorization**” means an authorization issued by the Council under section 13;

“**Government entity**” means any body of Government and includes a ministry, portfolio, statutory authority, company, board, department or office;

“**import**” means import into the Islands;

“**inspector**” means a person appointed to be an inspector under section 32;

“**internet or mail-order pharmacy**” means a pharmacy that —

- (a) operates by way of online or mail-order sales; and
- (b) does not have a physical premises where in-person pharmacy services are provided;

“**licence**” means a dispensing licence, a manufacturing licence, a product licence or a wholesale licence;

“**manufacturing licence**” means a licence referred to in section 4 and issued under section 9;

“**medicine**” means —

- (a) a substance or combination of substances —
 - (i) presented as having properties of preventing or treating disease in human beings or animals; or
 - (ii) that may be administered to human beings or animals with a view to restoring, correcting or modifying a physiological function by exerting a pharmacological, immunological or metabolic action, or making a medical diagnosis; or
- (b) a preparation developed from autologous or allogeneic biological materials;

“**narcotic**” means any of the following —

- (a) a drug included on the Yellow List of Narcotic Drugs Under International Control prepared by the International Narcotics Control Board; or
- (b) a controlled drug under the *Misuse of Drugs Act (2017 Revision)*;

“**non-approved medicine**” means a medicine other than an approved medicine;

“**pharmacist**” means a person who, under the *Health Practice Act (2021 Revision)* —

- (a) is registered to practice the profession of pharmacy; and
- (b) holds a practising licence to do so;

“**pharmacy medicine**” means a medicine prescribed as a pharmacy medicine by the regulations;

“**poison**” means a substance prescribed to be a poison by the regulations;

“**prescriber**” means —

- (a) a registered practitioner who is a medical doctor or dentist;
- (b) any other registered practitioner prescribed by the regulations; or
- (c) a veterinary surgeon;

“prescription” means a verbal or written instruction (including an instruction issued electronically) authorizing the dispensing of a medicine;

“prescription-only medicine” means medicine prescribed as a prescription-only medicine by the regulations;

“registered practitioner” has the meaning assigned by section 2 of the *Health Practice Act (2021 Revision)*;

“registrar” has the meaning assigned by section 2 of the *Health Practice Act (2021 Revision)*;

“scope of practice”, in relation to a registered practitioner, means the practitioner’s scope of practice under a code mentioned in section 35 of the *Health Practice Act (2021 Revision)*;

“ship’s physician” means a medical doctor employed to practise on board a commercial cruise ship or military vessel;

“supply” includes sell;

“veterinary surgeon” has the meaning assigned by section 2 of the *Veterinary Act (1997 Revision)*; and

“wholesale licence” means a licence referred to in section 5 and issued under section 9.

PART 2 - LICENCES AND AUTHORIZATIONS

Division 1 - Licence requirements

Importing medicines

3. (1) A person may import an approved medicine as follows —
- (a) a pharmacist may import an approved medicine for the purpose of dispensing it;
 - (b) a registered practitioner who is authorized to administer an approved medicine within the registered practitioner’s scope of practice may import the medicine for the purpose of administering the medicine to the practitioner’s patients in quantities that are appropriate for the size of the practitioner’s practice;
 - (c) under section 4(2)(a), the holder of a manufacturing licence for the manufacture of an approved medicine may import the ingredients (which



- may be medicines) specified in the licence for the purpose of manufacturing the medicine;
- (d) under section 5(2), the holder of a wholesale licence to sell an approved medicine by wholesale may import the medicine for the purpose of selling it;
 - (e) under section 6(2), the holder of a dispensing licence to dispense an approved medicine may import the medicine for the purpose of dispensing it;
 - (f) a person may import an approved medicine for self-administration if the following conditions are met —
 - (i) the quantity of the medicine is not of commercial value;
 - (ii) the medicine is contained in the original manufacturer's packaging or the container in which it was dispensed by a pharmacy and is labelled with a full description of the contents; and
 - (g) the Chief Medical Officer or the Director of Agriculture may import an approved medicine.
- (2) A person who imports a medicine except as specified in this section commits an offence and is liable on conviction to a fine of ten thousand dollars or to imprisonment for a term of two years, or to both.

Manufacturing licence

4. (1) Subject to subsection (3), a person shall not manufacture a medicine except in accordance with a manufacturing licence.
- (2) A manufacturing licence authorizes the holder of the licence to do any of the following in addition to manufacturing each medicine specified in the licence without an additional licence —
- (a) import the ingredients specified in the licence for the purpose of manufacturing the medicine; and
 - (b) sell the medicine by wholesale.
- (3) A person specified in section 17(1) does not require a manufacturing licence to compound a medicine.
- (4) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of twenty thousand dollars or to imprisonment for a term of five years, or to both.

Wholesale licence

5. (1) Subject to subsection (3), a person shall not sell an approved medicine by wholesale except in accordance with a wholesale licence.

- (2) A wholesale licence authorizes the holder of the licence to import, for the purpose of selling by wholesale, each medicine or class of medicines permitted to be sold by wholesale under the licence without an additional licence.
- (3) The holder of a manufacturing licence may sell a medicine specified in the licence by wholesale without a wholesale licence.
- (4) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of twenty thousand dollars or to imprisonment for a term of five years, or to both.

Dispensing licence

6. (1) A person shall not dispense a pharmacy medicine or a prescription-only medicine unless —
- (a) the person is a pharmacist or acting under the supervision of a pharmacist; or
 - (b) the person does so in accordance with a dispensing licence.
- (2) A dispensing licence authorizes the holder of the licence to import, for the purpose of dispensing, each medicine or class of medicines permitted to be dispensed under the licence without an additional licence.
- (3) Section 26 contains additional requirements for dispensing prescription-only medicines.
- (4) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of twenty thousand dollars or to imprisonment for a term of five years, or to both.

Division 2 - Licence applications

Application for licence

7. (1) A person may apply to the Council for a licence.
- (2) The application shall be —
- (a) made in the approved form; and
 - (b) accompanied by the prescribed fee and any documents prescribed by the regulations.

Criteria for issuing licence

8. (1) The criteria for issuing a manufacturing licence are as follows —
- (a) that the premises and equipment to be used for manufacturing operations under the licence are suitable for purpose;



- (b) that the manufacturing operations are to be supervised by a person who holds the qualifications prescribed by the regulations;
 - (c) that adequate records are to be kept of the operations and output; and
 - (d) any other criteria prescribed by the regulations.
- (2) The criteria for issuing a wholesale licence are as follows —
 - (a) that the premises and equipment to be used for storing and distributing medicines under the licence are suitable for purpose;
 - (b) that a pharmacist is to supervise the storage and distribution of medicines;
 - (c) that adequate records are to be kept of the storage and distribution of medicines; and
 - (d) any other criteria prescribed by the regulations.
- (3) The criteria for issuing a dispensing licence are as follows —
 - (a) that the premises and equipment to be used for storing and dispensing medicines under the licence are suitable for purpose;
 - (b) that a registered practitioner acting within the registered practitioner's scope of practice is to supervise the storage and dispensing of medicines;
 - (c) that adequate records are to be kept of the storage and dispensing of medicines;
 - (d) that adequate arrangements are in place for providing appropriate information to patients to whom medicines are dispensed;
 - (e) that a pharmacist, acting as a consultant, is to be available to advise the licence holder on dispensing medicines and to consult with patients when required; and
 - (f) any other criteria prescribed by the regulations.

Decision on licence application

9. The Council shall assess an application under section 7 and —
- (a) issue the licence if satisfied that the applicant —
 - (i) meets the criteria for the issue of the licence specified in section 8; and
 - (ii) is otherwise a fit and proper person to hold the licence; or
 - (b) refuse the application and give the applicant written notice of the refusal, including the reasons for it.

Duration and conditions of licence

10. (1) A licence may be issued for the period, not exceeding three years, specified in it and may be renewed.

- (2) A licence is subject to any conditions prescribed by the regulations and any additional conditions and limitations specified in it.

Variation, suspension and revocation of licence

- 11.** (1) The Council may vary the conditions of a licence if —
- (a) the Council becomes aware of information that, had it been known when the licence was issued, would have resulted in different or additional conditions being imposed on the licence; or
 - (b) the holder of the licence has breached a condition of the licence and the Council believes the variation is necessary to reduce the risk of a further breach.
- (2) Before deciding to vary the conditions of a licence, the Council shall —
- (a) give written notice to the holder of the licence —
 - (i) of the reasons for, and particulars of, the proposed variation; and
 - (ii) that the holder may make written submissions to the Council about the proposed variation within a reasonable period of at least twenty-one days stated in the notice; and
 - (b) have regard to any written submissions made by the holder within the stated period.
- (3) The Council may revoke a licence if —
- (a) the Council becomes aware of information that, had it been known when the application for the licence was made, would have resulted in the application being refused; or
 - (b) the holder of the licence has breached a condition of the licence in such a serious manner that the Council believes the licence should be revoked.
- (4) Before deciding to revoke the licence, the Council —
- (a) shall give written notice to the holder of the licence —
 - (i) of the reasons for the proposed revocation; and
 - (ii) that the holder may make written submissions to the Council about the proposed revocation within a reasonable period of at least twenty-one days stated in the notice; and
 - (b) may suspend the licence during the period stated in the notice; and
 - (c) shall have regard to any written submissions made by the holder within the stated period.
- (5) If the Council decides to vary, suspend or revoke a licence, the Council must give the holder written reasons for the decision.



Division 3 - Medicine approval and authorization

Medicine approval

- 12.** The Council may approve a medicine for the purposes of this Act if the Council is satisfied that the medicine —
- (a) is safe and effective for the purpose for which it is intended to be administered;
 - (b) is of good quality, according to the specification and method or proposed method of manufacture, and will be of good quality when sold and dispensed; and
 - (c) meets any other criteria prescribed by the regulations.

Exceptional use authorization

- 13.** (1) The Council may, on application by a prescriber, issue an exceptional use authorization for a non-approved medicine if the Council is satisfied that —
- (a) exceptional circumstances exist requiring the medicine to be made available; and
 - (b) the medicine is safe and effective for the purpose for which it is intended to be administered.
- (2) The non-approved medicine is taken to be an approved medicine for the purposes of this Act for the duration of the exceptional use authorization.
- (3) The exceptional use authorization —
- (a) may be issued for the use of the medicine generally or be limited to the administration of the medicine to a specified patient or class of patients; and
 - (b) is subject to the conditions and limitations specified in it.

Emergency use authorization

- 14.** (1) The Cabinet may, by notice published in the *Gazette*, issue an emergency use authorization for a non-approved medicine.
- (2) The non-approved medicine is taken to be an approved medicine for the purposes of this Act for the duration of the emergency use authorization.
- (3) The emergency use authorization is subject to the conditions and limitations specified in it.

PART 3 - DEALING WITH MEDICINES

Division 1 - Medicines generally

Dealing with non-approved medicine

- 15.** (1) A person shall not —
- (a) import;
 - (b) sell by wholesale;
 - (c) dispense;
 - (d) prescribe;
 - (e) administer; or
 - (f) manufacture,
- a non-approved medicine.
- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of twenty thousand dollars or to imprisonment for a term of five years, or to both.

Supplying a medicine

- 16.** (1) Subject to subsection (2), a person shall not supply a medicine by retail unless —
- (a) the person —
 - (i) is a pharmacist, or is acting under the supervision of a pharmacist; and
 - (ii) supplies the medicine from a certified pharmacy; or
 - (b) the person does so in accordance with a dispensing licence.
- (2) Subsection (1) does not apply to the supply of a medicine in the following circumstances —
- (a) the supply is made from premises capable of being contained and locked so as to exclude the public; and
 - (b) the medicine —
 - (i) has been prepared for retail supply and placed in a container or package that has not been opened since the medicine was prepared;
 - (ii) is supplied under the authority of the person holding a manufacturing or wholesale licence for the medicine; and
 - (iii) is not a pharmacy medicine, prescription-only medicine or narcotic.



- (3) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of twenty thousand dollars or to imprisonment for a term of five years, or to both.

Compounding a medicine

- 17.** (1) A person shall not compound a medicine unless the person is —
- (a) a pharmacist;
 - (b) acting under the supervision of a pharmacist;
 - (c) the holder of a dispensing licence who is authorized to compound the medicine under the licence;
 - (d) a registered practitioner acting within the registered practitioner's scope of practice; or
 - (e) a veterinary surgeon acting within the veterinary surgeon's scope of practice.
- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of twenty thousand dollars or to imprisonment for a term of five years, or to both.

Altering a medicine

- 18.** (1) A person shall not —
- (a) alter the composition of a medicine with the intention of supplying the medicine to another person; or
 - (b) supply, or possess with the intention of supplying, a medicine with an altered composition.
- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of twenty thousand dollars or to imprisonment for a term of five years, or to both.
- (3) Subsection (1) is not contravened only because a medicine contains extraneous matter as a result of the process of manufacturing or compounding the medicine.

Labelling of medicine

- 19.** (1) A person shall not supply, or possess with the intention of supplying, a medicine in a container that is labelled in a way that —
- (a) falsely describes the medicine;
 - (b) is likely to be misleading in relation to the nature, efficacy or quality of the medicine; or
 - (c) does not meet the requirements prescribed by the regulations for the labelling of the medicine.

- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of ten thousand dollars or to imprisonment for a term of two years, or to both.

Information accompanying medicine

20. (1) A person shall not supply with a medicine, or possess with the intention of supplying with a medicine, written information (whether contained in a leaflet or otherwise) that —
- (a) falsely describes the medicine;
 - (b) is likely to be misleading in relation to the nature, efficacy or quality of the medicine; or
 - (c) does not meet the requirements prescribed by the regulations or specified in the *Misuse of Drugs Act (2017 Revision)* for information accompanying the medicine.
- (2) Subject to subsection (3), a person shall not supply a medicine unless the medicine is supplied with a patient information leaflet written in English.
- (3) The Chief Medical Officer may, on a case-by-case basis, issue an exemption from the requirement to supply a medicine with a patient information leaflet written in English if it is not reasonably practicable to comply with the requirement.
- (4) A person who contravenes subsection (1) or (2) commits an offence and is liable on conviction to a fine of ten thousand dollars or to imprisonment for a term of two years, or to both.

Vending machines

21. (1) A person (“person A”) shall not —
- (a) install a vending machine on premises occupied by person A for the purpose of supplying a pharmacy medicine, prescription-only medicine or narcotic;
 - (b) dispense a pharmacy medicine, prescription-only medicine or narcotic using a vending machine; or
 - (c) allow another person to install a vending machine on premises occupied by person A for the purpose of supplying a pharmacy medicine, prescription-only medicine or narcotic.
- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of ten thousand dollars or to imprisonment for a term of two years, or to both.
- (3) In this section, “**vending machine**” means a machine or device from which a product can be obtained, including by —
- (a) electronic funds transfer; or



- (b) inserting money, a token or something else.

Internet and mail order pharmacies

- 22.** (1) A person shall not dispense a pharmacy medicine or prescription-only medicine from an internet or mail-order pharmacy.
- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of twenty thousand dollars or to imprisonment for a term of five years, or to both.

Division 2 - Pharmacy medicines, prescription-only medicines and other restricted products

Wholesale of pharmacy or prescription-only medicine

- 23.** (1) The holder of a wholesale licence or a manufacturing licence shall not sell a pharmacy medicine or a prescription-only medicine by wholesale except to —
- (a) a pharmacist operating a certified pharmacy;
 - (b) the holder of a dispensing licence;
 - (c) in the case of a pharmacy medicine, a registered practitioner who is authorized to administer the medicine within the registered practitioner's scope of practice;
 - (d) in the case of a prescription-only medicine, a prescriber who is authorized to administer the medicine within the prescriber's scope of practice; or
 - (e) a Government entity that is authorized by law to use the medicine.
- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of twenty thousand dollars or to imprisonment for a term of five years, or to both.

Prescribing medicine

- 24.** (1) Subject to subsection (2), a person shall not issue a prescription unless —
- (a) the person is a prescriber acting within the prescriber's scope of practice; and
 - (b) if the prescription is for a narcotic, the prescriber complies with the requirements prescribed by the regulations for prescribing narcotics.
- (2) A ship's physician who is not a registered practitioner may issue a prescription for the supply (not exceeding a seven-day supply or, if the medicine is packaged in units that each exceed a seven-day supply, a single unit), of a medicine other than a narcotic, but may not authorize a refill of the prescription.

- (3) A person who contravenes subsection (1) or (2) commits an offence and is liable on conviction to a fine of twenty thousand dollars or to imprisonment for a term of five years, or to both.

Altering a prescription

- 25.** (1) A person shall not alter a prescription unless the person is —
- (a) the prescriber who issued the prescription;
 - (b) a pharmacist acting under the instruction of the prescriber who issued the prescription; or
 - (c) a pharmacist acting within the pharmacist's scope of practice.
- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of ten thousand dollars or to imprisonment for a term of two years, or to both.

Dispensing a prescription-only medicine

- 26.** (1) Subject to subsection (2), an authorized dispenser shall not dispense a prescription-only medicine unless —
- (a) the person does so in accordance with a prescription issued within —
 - (i) twelve months before the date of dispensing the medicine; or
 - (ii) if the regulations prescribe a different period, that period; and
 - (b) if the medicine is a narcotic, the person complies with the requirements prescribed by the regulations for dispensing narcotics.
- (2) Notwithstanding subsection (1) —
- (a) a prescriber may dispense a prescription-only medicine to the prescriber's patients in accordance with a dispensing licence;
 - (b) a veterinary surgeon may dispense a prescription-only medicine within the scope of the veterinary surgeon's practice; or
 - (c) a pharmacist may dispense a supply of a prescription-only medicine without a prescription ("emergency supply") if the following conditions are satisfied —
 - (i) the medicine is not a narcotic;
 - (ii) the medicine is essential to maintaining the patient's life or continuing therapy and, in the professional opinion of the pharmacist, interrupting the therapy would be detrimental to the patient's well-being;
 - (iii) the request for the emergency supply has not been made immediately after a previous emergency supply of the medicine was given and used;



- (iv) the pharmacist makes a complete record of the particulars of the supply; and
 - (v) the amount dispensed does not exceed a fourteen-day supply or, if the medicine is packaged in units that each exceed a fourteen-day supply, a single unit.
- (3) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of ten thousand dollars or to imprisonment for a term of two years, or to both.

Interchangeable and bio-equivalent generic medicines

- 27.** (1) Subject to subsection (3), an authorized dispenser may substitute an interchangeable and bio-equivalent generic medicine for a product specified in a prescription if the authorized dispenser —
- (a) informs the patient; and
 - (b) labels the container of the medicine with the name of the medicine dispensed rather than the brand name of the medicine prescribed.
- (2) A medicine is an interchangeable and bio-equivalent generic medicine in relation to a specified product if it —
- (a) is an approved medicine; and
 - (b) has the same active ingredient, in the same strength and the same dosage, as the specified product.
- (3) An authorized dispenser shall not substitute an interchangeable and bio-equivalent medicine if the substitution is prohibited by the regulations.
- (4) A person who contravenes subsection (1) or (3) commits an offence and is liable on conviction to a fine of five thousand dollars or to imprisonment for a term of one year, or to both.

Dispensing samples

- 28.** (1) An authorized dispenser shall not dispense a prescription-only medicine by way of a sample except in accordance with the regulations.
- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of five thousand dollars or to imprisonment for a term of one year, or to both.

Administering a prescription-only medicine

- 29.** (1) A person shall not administer a prescription-only medicine to another person unless the person is —
- (a) the prescriber of the medicine;

- (b) acting in accordance with the written directions of the prescriber of the medicine; or
 - (c) a registered practitioner acting within the registered practitioner's scope of practice.
- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of ten thousand dollars or to imprisonment for a term of two years, or to both.

Possessing a prescription-only medicine

- 30.** (1) A person shall not possess a prescription-only medicine unless the person —
- (a) has a prescription for the medicine in the person's name;
 - (b) is acting under the directions of a person who holds a prescription for the medicine;
 - (c) is an authorized dispenser;
 - (d) is a registered practitioner who, acting within the registered practitioner's scope of practice, may dispense the medicine;
 - (e) is a veterinary surgeon who, acting within the veterinary surgeon's scope of practice, may dispense the medicine;
 - (f) is acting under the written directions of a prescriber; or
 - (g) holds a manufacturing or wholesale licence for the medicine.
- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of five thousand dollars or to imprisonment for a term of one year, or to both.

Supply and labelling of poison

- 31.** (1) The Council may, in accordance with the requirements of the regulations, certify a person to be a person who may possess a poison.
- (2) A person ("person A") shall not supply another person ("person B") with a poison unless —
- (a) person A is a pharmacist who supplies the poison from a certified pharmacy; and
 - (b) person B is certified by the Council under subsection (1).
- (3) A person who contravenes subsection (2) commits an offence and is liable on conviction to a fine of ten thousand dollars or to imprisonment for a term of two years, or to both.
- (4) A person shall not supply a poison, or possess a poison with the intention of supplying it, unless the container of the poison is labelled as follows —
- (a) with the name of the poison;



- (b) with the word “poison”;
 - (c) with the name and address of the certified pharmacy from which the poison is supplied;
 - (d) with an indication of the general or specific risk taken in its use, together with any safety precautions to be taken; and
 - (e) in accordance with any other requirements prescribed by the regulations.
- (5) A person who contravenes subsection (4) commits an offence and is liable on conviction to a fine of five thousand dollars or to imprisonment for a term of one year, or to both.

PART 4 - ENFORCEMENT

Appointment of inspectors

- 32.** (1) The registrar may, in writing, appoint a person to be an inspector.
- (2) An inspector shall assist the Council and has the functions and powers conferred on an inspector by this Act, subject to any restrictions specified in the inspector’s instrument of appointment.
- (3) In exercising powers or performing functions, an inspector shall comply with the directions of the registrar.
- (4) The registrar shall issue each inspector with an identity card —
- (a) stating the inspector’s name and the expiry date of the card;
 - (b) showing a recent photograph of the inspector; and
 - (c) signed by the Chief Medical Officer or a person designated by the Chief Medical Officer.
- (5) An inspector exercising a power or performing a function in relation to a person shall, if asked by the person, produce the inspector’s identity card.

Powers of inspectors

- 33.** (1) An inspector may exercise any of the powers specified in subsection (2) if the inspector reasonably believes it is necessary to do so for the administration or enforcement of this Act.
- (2) Subject to section 34, the inspector may do any of the following in relation to a place —
- (a) enter, inspect and search the place;
 - (b) require a person at the place to give the inspector specified information (including documents) or assistance to access anything at the place;
 - (c) inspect and take samples of anything found in the place that appears to the inspector to be —

- (i) a medicine;
 - (ii) a container or package used or intended to be used to contain a medicine or a document used or intended to be used to provide information to accompany a medicine; or
 - (iii) anything used in connection with the manufacture or assembly of a medicine, including an ingredient of the medicine;
- (d) seize anything the inspector reasonably believes —
 - (i) is connected with the commission, or intended commission, of an offence under this Act; or
 - (ii) provides evidence of the commission of an offence under this Act; and
- (e) make a record relating to the exercise of a power under this subsection (including, for example, by making a copy of a document, taking a photograph or making a video recording).

Restrictions on powers of entry

- 34.** (1) This section applies if an inspector intends to enter a place under section 33(2).
- (2) If the place is not a residence, the inspector shall, as far as practicable, give notice to the owner of the place of the inspector's intention to enter the place before doing so.
- (3) If the place is a residence, the inspector shall not enter the place unless —
- (a) the inspector has given written notice to the owner of the place of the inspector's intention to enter the place; and
 - (b) the inspector —
 - (i) is permitted by the owner (whether orally or in writing) to enter the place; or
 - (ii) is authorized to enter the place under a warrant issued by a Magistrate.
- (4) A Magistrate may issue the warrant only if satisfied that the warrant is reasonably required in the circumstances for the enforcement of this Act.
- (5) The warrant may authorize the inspector, with the assistance of a constable, to use force to enter the place.

Dealing with seized items

- 35.** (1) An inspector shall give a receipt to a person for anything the inspector seizes from the person under this Act.
- (2) The registrar has control of anything seized under this Act.



- (3) The registrar may retain anything seized that is relevant to the prosecution of an offence under this Act until the proceedings for the offence have ended.

Obstructing an inspector

- 36.** (1) A person who obstructs an inspector who is exercising a power or performing a function under this Act, including by destroying any document or thing during the course of a search conducted under section 33(2), commits an offence and is liable on conviction to a fine of five thousand dollars or to imprisonment for a term of one year, or to both.
- (2) A person who fails to comply with a requirement given to the person by an inspector who is exercising a power or performing a function under this Act commits an offence and is liable on conviction to a fine of five thousand dollars or to imprisonment for a term of one year, or to both.

PART 5 - MISCELLANEOUS

Approved forms

- 37.** The registrar may approve forms for use under this Act.

Protection from liability and indemnity

- 38.** (1) Each of the following persons is not civilly or criminally liable for an act done or omitted to be done by the person in good faith in the exercise of a power or performance of a function under this Act —
- (a) the registrar;
 - (b) an inspector;
 - (c) a member of the Council; or
 - (d) the Chief Medical Officer.
- (2) The Government shall indemnify each of the following persons against all claims, damages, costs, charges or expenses incurred for an act done or omitted to be done in good faith in the exercise of a power or performance of a function under this Act —
- (a) the registrar;
 - (b) an inspector;
 - (c) a member of the Council; or
 - (d) the Chief Medical Officer.

Confidentiality of information

- 39.** (1) A person commits an offence if —

- (a) the person obtains information in the course of performing a function connected with the administration of this Act or exercising a power under this Act;
 - (b) the person recklessly or intentionally discloses the information to another person; and
 - (c) the disclosure is not —
 - (i) for a purpose connected with the administration of this Act, including a legal proceeding arising out of the operation of this Act; or
 - (ii) to a person who is otherwise entitled to the information.
- (2) A person who contravenes subsection (1) commits an offence and is liable on conviction to a fine of ten thousand dollars or to imprisonment for a term of one year, or to both.

Appeals

40. Where a person is aggrieved by a decision of the Council under this Act —

- (a) the person may appeal against the decision to the Health Appeals Tribunal established by section 4 of the *Health Practice Act (2021 Revision)*; and
- (b) Schedule 2 to the *Health Practice Act (2021 Revision)* applies to the conduct of the appeal.

Regulations

41. (1) The Cabinet, on the advice of the Council, the Chief Medical Officer or the Chief Nursing Officer, may make regulations prescribing all matters that are required or permitted by this Act to be prescribed, or are necessary to be prescribed to give effect to the purposes of this Act.

- (2) Without limiting subsection (1), the Cabinet may make regulations —
- (a) controlling, regulating or prohibiting the manufacture, sale, supply, possession, export or import of any medicine;
 - (b) regulating the issue of prescriptions;
 - (c) regulating the storage and disposal of medicines;
 - (d) prohibiting or regulating the advertising of medicines;
 - (e) regulating the keeping of records relating to the supply of medicines; and
 - (f) prescribing fees and providing for the waiver, refund and discount of fees.
- (3) Regulations made under this Act may —
- (a) make different provision in relation to different cases or circumstances;
 - (b) apply in respect of particular persons or particular cases or particular classes of persons or particular classes of cases, and define a class by reference to any circumstances whatsoever; or



- (c) contain such transitional, consequential, incidental or supplementary provisions as appear to the Cabinet to be necessary or expedient for the purposes of the regulations.
- (4) Regulations made under this Act may create an offence punishable by a penalty not exceeding the maximum penalty prescribed under this Act.

PART 6 - REPEALS

Repeals

42. The following are repealed —

- (a) the *Pharmacy Act, 1979*; and
- (b) the *Pharmacy Act, 1991*.

Passed by the Parliament the 23rd day of July, 2024.

Hon. Sir Alden McLaughlin
Speaker

Zena Merren-Chin
Clerk of the Parliament