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#### Definitions

1 In these bylaws:

"Act" means the Pharmacy Operations and Drug Scheduling Act,

"attestation" means the attestation referred to in section 2(2)(d)(ii) of the Act,

"BC Annual Report" means an annual report filed with the BC Registry Services;

"British Columbia Company Summary" means a summary issued by the BC Registry Services;

"central pharmacy" means a community pharmacy that holds one or more telepharmacy licences;

"Central Securities Register" means the register maintained under section 111(1) of the *Business Corporations Act* [SBC 2002] C.57 as amended;

"community pharmacy" means a pharmacy licensed to sell or dispense drugs to the public, but does not include a telepharmacy;

"**Community Pharmacy Standards of Practice**" means the standards, limits and conditions for practice established under section 19(1)(k) of the *Health Professions Act* respecting community pharmacies;

"controlled drug substances" means a drug which includes a substance listed in the Schedules in the regulations made pursuant to the *Controlled Drugs and Substances Act* (Canada), and Part G of the *Food and Drug Regulations* (Canada);

"controlled prescription program" means a program approved by the board, to prevent prescription forgery and reduce inappropriate prescribing of drugs;

"**criminal record history**" means the results of a criminal record search of Royal Canadian Mounted Police and local police databases, in the form approved by the board;

"direct owner" has the same meaning as in section 1 of the Act,

"direct supervision" means real time audio and visual observation by a full pharmacist of pharmacy services performed at a telepharmacy consistent with a pharmacy manager's responsibilities as set out in section 18(2);

"dispensary" means the area of a community pharmacy or a telepharmacy that contains Schedule I and II drugs;

"drug" has the same meaning as in section 1 of the Act,

#### "electronic signature" means

(a) information in electronic form that a person has created or adopted in order to sign a record, other than with respect to a prescription signed by a full

pharmacist for the purpose of prescribing, that is in, attached to or associated with a record, is secure and is only reproducible and used by that person, and,

(b) with respect to a prescription signed by a full pharmacist for the purpose of prescribing, the electronic signature must meet the requirements of paragraph
(a) and must be a unique mark personally applied by that pharmacist;

"full pharmacist" means a member of the College who is registered in the class of registrants established in section 41(a) of the bylaws under the *Health Professions Act*,

#### "health authority" includes

- (a) a regional health board designated under the Health Authorities Act,
- (b) the Provincial Health Services Authority,
- (c) First Nations Health Authority, and
- (d) Providence Health Care Society;

"hospital" has the same meaning as in section 1 of the Hospital Act;

"hospital pharmacy" means a pharmacy licensed to operate in or for a hospital;

"hospital pharmacy satellite" means a physically separate area on or outside the hospital premises used for the provision of pharmacy services which is dependent upon support and administrative services from the hospital pharmacy;

"Hospital Pharmacy Standards of Practice" means the standards, limits and conditions for practice established under section 19(1)(k) of the Health Professions Act respecting hospital pharmacies;

**"incentive"** has the same meaning as in Part 1 of Schedule "F" of the bylaws of the College under the *Health Professions Act;* 

"indirect owner" has the same meaning as in section 1 of the Act;

"manager" has the same meaning as in section 1 of the Act;

"outsource prescription processing" means to request another community pharmacy to prepare or process a prescription drug order;

"patient's representative" means a person who is authorized to act on a patient's behalf;

"personal health information" has the same meaning as in section 25.8 of the *Health Professions Act*;

"pharmacy" has the same meaning as in section 1 of the Act;

"pharmacy education site" means a pharmacy

- (a) that has Schedule I, II and III drugs, but no controlled drug substances,
- (b) that is licensed solely for the purpose of pharmacy education, and
- (c) from which pharmacy services are not provided to any person;

#### "pharmacy security" means

- (a) measures to prevent unauthorized access and loss of Schedule I, IA, II and III drugs, and controlled drug substances,
- (b) measures providing for periodic and post-incident review of pharmacy security,
- (c) measures to protect against unauthorized access, collection, use, disclosure or disposal of personal health information;

"pharmacy services" has the same meaning as in section 1 of the bylaws of the College under the *Health Professions Act*;

"**pharmacy technician**" has the same meaning as in section 1 of the bylaws of the College under the *Health Professions Act*;

"prescription drug" means a drug referred to in a prescription;

"professional products area" means the area of a community pharmacy that contains Schedule III drugs;

"professional service area" means the area of a community pharmacy that contains Schedule II drugs;

"**record**" has the same meaning as the definition of record in Schedule 1 of the *Freedom of Information and Protection of Privacy Act*,

"Residential Care Facilities and Homes Standards of Practice" means the standards, limits and conditions for practice established under section 19(1)(k) of the Health Professions Act respecting residential care facilities and homes;

"rural and remote community" means a community set out in Schedule "H";

"Schedule I, Schedule IA, Schedule II, or Schedule III", as the case may be, refers to the drugs listed in Schedule I, IA, II or III of the *Drug Schedules Regulation*;

"**signature**" on a record means either a handwritten signature in ink or an electronic signature;

"**support person**" has the same meaning as in the *Act* except that it does not include a pharmacy technician;

"**telepharmacy**" means a pharmacy located in a rural and remote community that is licensed to provide pharmacy services;

"**Telepharmacy Standards of Practice**" means the standards, limits and conditions for practice established under section 19(1)(k) of the *Health Professions Act* respecting the operation of telepharmacies.

# PART I – Pharmacy Licences

### Licence Types

2 (1) The registrar may issue a licence for any of the following:

- (a) a community pharmacy;
- (b) a hospital pharmacy;
- (c) a pharmacy education site; or
- (d) a telepharmacy.

### **New Community Pharmacy Licence**

- 3 (1) Applicants for a new community pharmacy licence must submit an application consistent with the type of ownership under section 5(2) of the *Act*.
  - (2) A direct owner may apply for a new community pharmacy licence by submitting:
    - (a) an application in Form 1A;
    - (b) the fee(s) specified in Schedule "A";
    - (c) a diagram professionally drawn to scale, including the measurements and entrances of the pharmacy, demonstrating compliance with the physical requirements in the bylaws and applicable policies;
    - (d) Form 10A;
    - (e) photographs or video demonstrating compliance with the physical requirements in the bylaws and applicable policies; and
    - (f) a copy of the pharmacy's valid business licence issued by the jurisdiction to the direct owner, if applicable.
  - (3) In addition to the requirements in subsection (2), a direct owner described in section 5(2)(b) or (c) of the *Act* must submit:
    - (a) an email contact of each indirect owner;
    - (b) a copy of the power(s) of attorney, if applicable;
    - (c) a copy of the current British Columbia Company Summary; and
    - (d) a certified true copy of the Central Securities Register if a direct owner is or includes a corporation that is not traded publicly.
  - (4) If an indirect owner is a company incorporated under the *Company Act* or the *Business Corporations Act* that is not traded publicly, the following must be submitted for that company:
    - (a) an email contact of each indirect owner;

- (b) a copy of the power(s) of attorney, if applicable;
- (c) a copy of the current British Columbia Company Summary; and
- (d) a certified true copy of the Central Securities Register.
- (5) Proof of eligibility in Form 5 and a criminal record history in accordance with section 14 must be submitted by the following:
  - (a) any pharmacist who is a direct owner described in section 5(2)(a) of the *Act*;
  - (b) indirect owner(s); and
  - (c) the manager.

#### **Community Pharmacy Licence Renewal**

- 4 (1) A direct owner may apply to renew a community pharmacy licence no later than 30 days prior to the expiry of the existing pharmacy licence by submitting:
  - (a) an application in Form 2A;
  - (b) the fee(s) specified in Schedule "A";
  - (c) a copy of the pharmacy's valid business licence issued by the jurisdiction to the direct owner, if applicable; and
  - (d) a copy of the current British Columbia Company Summary or the most recently filed BC Annual Report, if a direct owner is or includes a corporation.
  - (2) At the time of the renewal application, an attestation in Form 5 must be submitted by:
    - (a) any pharmacist who is a direct owner described in section 5(2)(a) of the *Act*;
    - (b) indirect owner(s); and
    - (c) the manager.
  - (3) An application submitted later than 30 days prior to the expiry of the pharmacy licence is subject to the fee(s) specified in Schedule "A".

#### **Community Pharmacy Licence Reinstatement**

- 5 (1) A direct owner may apply to reinstate a community pharmacy licence that has been expired for 90 days or less by submitting:
  - (a) an application in Form 3A;
  - (b) the fee(s) specified in Schedule "A";

- (c) a copy of the pharmacy's valid business licence issued by the jurisdiction to the direct owner, if applicable; and
- (d) a copy of the current British Columbia Company Summary, if the direct owner is or includes a corporation.
- (2) At the time of the reinstatement application, an attestation in Form 5 must be submitted by:
  - (a) any pharmacist who is a direct owner described in section 5(2)(a) of the *Act*;
  - (b) indirect owner(s); and
  - (c) the manager.

### **New Hospital Pharmacy Licence**

- 6 (1) Applicants for a new hospital pharmacy licence must submit an application consistent with the type of ownership under section 5(2) of the *Act*.
  - (2) A direct owner may apply for a new hospital pharmacy licence by submitting:
    - (a) an application in Form 1C;
    - (b) the fee(s) specified in Schedule "A"; and
    - (c) a diagram professionally drawn to scale, including the measurements and entrances of the pharmacy, demonstrating compliance with the physical requirements in the bylaws and applicable policies.
  - (3) The manager must submit an attestation in Form 5 and a criminal record history in accordance with section 14.
  - (4) A pharmacy located in a hospital which dispenses drugs to staff, out-patients or the public and which is not owned or operated by a health authority, must be licensed as a community pharmacy or telepharmacy.

#### Hospital Pharmacy Licence Renewal

- 7 (1) A direct owner may apply to renew a hospital pharmacy licence no later than 30 days prior to the expiry of the existing pharmacy licence by submitting:
  - (a) an application in Form 2C; and
  - (b) the fee(s) specified in Schedule "A".
  - (2) At the time of the renewal application, the manager must submit an attestation in Form 5.
  - (3) An application submitted later than 30 days prior to the expiry of the pharmacy licence is subject to the fee(s) specified in Schedule "A".

# Hospital Pharmacy Licence Reinstatement

- 8 (1) A direct owner may apply to reinstate a pharmacy licence that has been expired for 90 days or less by submitting:
  - (a) an application in Form 3C; and
  - (b) the fee(s) specified in Schedule "A".
  - (2) At the time of the reinstatement application, the manager must submit an attestation in Form 5.

# **New Pharmacy Education Site Licence**

- 9 (1) Applicants for a new pharmacy education site licence must submit an application consistent with the type of ownership under section 5(2) of the *Ac*t.
  - (2) A direct owner may apply for a new pharmacy education site licence by submitting:
    - (a) an application in Form 1F; and
    - (b) the fee(s) specified in Schedule "A".
  - (3) The manager must submit an attestation in Form 5 and a criminal record history in accordance with section 14.

#### **Pharmacy Education Site Licence Renewal**

- 10 (1) A direct owner may apply to renew a pharmacy education licence no later than 30 days prior to the expiry of the existing pharmacy licence by submitting:
  - (a) an application in Form 2F; and
  - (b) the fee(s) specified in Schedule "A".
  - (2) At the time of the renewal application, the manager must submit an attestation in Form 5.
  - (3) An application submitted later than 30 days prior to the expiry of the pharmacy licence is subject to the fee(s) specified in Schedule "A".

#### **Pharmacy Education Site Licence Reinstatement**

- 11 (1) A direct owner may apply to reinstate a pharmacy education site licence that has been expired for 90 days or less by submitting:
  - (a) an application in Form 3F; and
  - (b) the fee(s) specified in Schedule "A".
  - (2) At the time of the reinstatement application, the manager must submit an attestation in Form 5.

#### New Telepharmacy Licence

- 12 A direct owner of a community pharmacy may apply for a new telepharmacy licence by submitting:
  - (a) an application in Form 1B;
  - (b) the fee(s) specified in Schedule "A";
  - (c) a diagram professionally drawn to scale, including the measurements and entrances of the telepharmacy, demonstrating compliance with the physical requirements in the bylaws and applicable policies;
  - (d) Form 10B;
  - (e) photographs or video demonstrating compliance with the physical requirements in the bylaws and applicable policies; and
  - (f) if applicable, a copy of the telepharmacy's valid business licence issued to the direct owner by the jurisdiction in which the telepharmacy is located.

### **Conditions for Telepharmacy Licence**

- 12.1 (1) The registrar must not issue a telepharmacy licence to a central pharmacy unless
  - (a) the proposed telepharmacy will be the only telepharmacy or community pharmacy located in the rural and remote community,
  - (b) the proposed telepharmacy is located at least 25 kilometers away from any other telepharmacy or community pharmacy,
  - (c) the proposed name on the external signage of the telepharmacy described in section 18(2)(r) includes the word "telepharmacy",
  - (d) except for a pharmacy located at an address listed in Schedule "F", the proposed telepharmacy does not have a licence as a community pharmacy,
  - (e) the central pharmacy applicant and the telepharmacy will have the same direct owner, and
  - (f) the central pharmacy is in compliance, and the telepharmacy will be in compliance, with the *Telepharmacy Standards of Practice*.
  - (2) A telepharmacy licence issued under subsection (1) is valid only for the location stated on the telepharmacy licence.

#### **Telepharmacy Licence Renewal**

- 13 (1) A direct owner may apply to renew a telepharmacy licence no later than 30 days prior to the expiry of the existing telepharmacy licence by submitting:
  - (a) an application in Form 2B;

- (b) the fee(s) specified in Schedule "A"; and
- (c) if applicable, a copy of the telepharmacy's business licence issued by the jurisdiction in which the telepharmacy is located.
- (2) An application submitted later than 30 days prior to the expiry of the telepharmacy licence is subject to the fee(s) specified in Schedule "A".

### **Telepharmacy Licence Reinstatement**

- 13.1 A direct owner may apply to reinstate a telepharmacy licence that has been expired for 90 days or less by submitting:
  - (a) an application in Form 3B;
  - (b) the fee(s) specified in Schedule "A"; and
  - (c) if applicable, a copy of the telepharmacy's valid business licence issued to the direct owner by the jurisdiction in which the telepharmacy is located.

### Criminal Record History of Direct Owner, Indirect Owner(s) and Manager

14 A direct owner, indirect owner(s) and a manager must submit a criminal record history pursuant to section 5.1 of the *Act*, in the form approved by the board.

### **Unlawful Operation**

- 15 (1) Pursuant to section 7(1) of the *Act*, persons listed in Schedule "B" are authorized under this bylaw to store, dispense or sell drugs or devices to the public.
  - (2) Pursuant to section 7(3) of the *Act*, the registrar may authorize the direct owner, indirect owner(s) or manager of an unlicensed pharmacy, or a full pharmacist to continue the operation of the pharmacy for a period not exceeding 90 days, for the limited purpose of transferring drugs and personal health information on the premises to another licensed pharmacy.
  - On receiving a referral under section 16(6), the application committee may consider whether to authorize the operation of the pharmacy pursuant to section 7(3) of the *Act* pending a determination under section 4(4)(b) of the *Act* as to relevance or risk to the public.

#### **PART II - All Pharmacies**

#### Change in Direct Owner, Indirect Owner(s) or Manager

- 16 (1) If a direct owner changes, the registrar may issue a new pharmacy licence upon receipt of the following from the new direct owner:
  - (a) Form 8A;
  - (b) the fee(s) specified in Schedule "A";
  - (c) a copy of the pharmacy's valid business licence issued by the jurisdiction to the new direct owner, if applicable; and

- (d) the documents listed in sections 3(3), 3(4) and 3(5) as applicable.
- (2) If there is a change of indirect owner(s) the following must be submitted by the direct owner:
  - (a) Form 8B;
  - (b) the fee(s) specified in Schedule "A";
  - (c) a Notice of Change of Directors, if applicable;
  - (d) a certified true copy of the Central Securities Register, if there is a change of shareholder(s) of a non-publicly traded corporation; and
  - (e) the documents listed in sections 3(3), 3(4) and 3(5), as applicable.
- (3) If the change in subsection (2) includes a new indirect owner(s), proof of eligibility in Form 5 and a criminal record history in accordance with section 14 must be submitted by the new indirect owner(s).
- (4) If there is a change of manager, the registrar may issue a new pharmacy licence and telepharmacy licence if applicable, upon receipt of:
  - (a) Form 8C submitted by the direct owner;
  - (b) the fee(s) specified in Schedule "A"; and
  - (c) proof of eligibility in Form 5 and a criminal record history in accordance with section 14 submitted by the new manager.
- (5) In the event that a direct owner, indirect owner(s) or manager is no longer eligible under section 3 of the *Act*, the direct owner, indirect owner(s) or manager must submit a notice in Form 6.
- (6) On receipt of a Form 6 under subsection (5), the registrar must refer the matter to the application committee who may act under sections 4(3), 4(4), and 4(5) of the *Act*.

# Changes to the Pharmacy Premises and Name

- 17 (1) If there is a change in the name of a corporation that is a direct owner, the registrar may amend the pharmacy licence, and telepharmacy licence if applicable, upon receipt of the following from the direct owner:
  - (a) Form 8D;
  - (b) the fee(s) specified in Schedule "A";
  - (c) a copy of the pharmacy's valid business licence issued by the jurisdiction to the direct owner with the new corporation name, if applicable; and
  - (d) a copy of the Alteration to the Notice of Articles.

- (2) If there is a change in the name of a corporation that is an indirect owner, the following must be submitted by the direct owner:
  - (a) Form 8D;
  - (b) the fee(s) specified in Schedule "A"; and
  - (c) a copy of the Alteration to the Notice of Articles.
- (3) If there is a change in the name on the external signage described in section 18(2)(q) or section 18(2)(r), or in the operating name of the pharmacy, the registrar may amend the pharmacy or telepharmacy licence upon receipt of the following from the direct owner:
  - (a) Form 8E;
  - (b) the fee(s) specified in Schedule "A";
  - (c) for a change of operating name, a copy of the pharmacy's valid business licence with the new operating name issued by the jurisdiction to the direct owner, if applicable; and
  - (d) for a change of the name on the external signage, photographs or video demonstrating compliance with section 18(2)(q) or 18(2)(r).
- (4) If there is a change in location of the pharmacy, the registrar may issue a new pharmacy licence upon receipt of the following from the direct owner:
  - (a) Form 8F;
  - (b) the fee(s) specified in Schedule "A";
  - (c) the requirements in sections 3(2)(c), (d) and (e) for a community pharmacy, or
  - (d) the requirements in section 6(2)(c) for a hospital pharmacy;
  - (e) a copy of the pharmacy's valid business licence with the address of the new location issued by the jurisdiction to the direct owner, if applicable; and
  - (f) photographs or video demonstrating compliance with section 18(2)(ee)(v).
- (5) If there is a change in layout of the pharmacy, the direct owner must submit the following:
  - (a) Form 8G;
  - (b) the fee(s) specified in Schedule "A"; and
  - (c) a diagram, photographs or video to demonstrate the changes in layout in accordance with sections 3(2)(c), (d) and (e) for a community pharmacy;

- (d) a diagram to demonstrate the changes in layout in accordance with section 6(2)(c) for a hospital pharmacy; or
- (e) a diagram, photographs or video to demonstrate the changes in layout in accordance with sections 12(c), (d) and (e) for a telepharmacy.
- 17.1 (1) A direct owner of a pharmacy that is permanently closing must notify the registrar by submitting the following at least 30 days before closure:
  - (a) an application in Form 4A;
  - (b) the fee(s) specified in Schedule "A";
  - (c) documents demonstrating compliance with sections 18(2)(ee)(i), (ii), (iii) and (iv); and
  - (d) photographs or video demonstrating compliance with section 18(2)(ee)(v).
  - (2) The manager of the pharmacy receiving drugs, medical devices, and/or patient and prescription records from the closing pharmacy must submit Part 2 of Form 4A within 14 days of receiving date the drugs, medical devices, and/or patient and prescription records.

### Responsibilities of Manager, Direct Owners, Directors, Officers and Shareholders

- 18 (1) A full pharmacist may not act as manager of more than one pharmacy location, unless the pharmacy of which the full pharmacist is manager includes
  - (a) a telepharmacy,
  - (b) a hospital pharmacy,
  - (c) a hospital pharmacy satellite, or
  - (d) a pharmacy education site.
  - (2) A manager must do all of the following:
    - (a) personally manage and be responsible for the daily operation of the pharmacy;
    - (b) ensure compliance with all legislation, bylaws, policies, procedures applicable to the operation of a pharmacy;
    - (c) establish policies and procedures
      - (i) to specify the duties to be performed by registrants and support persons,
      - (ii) for inventory management, product selection, and proper destruction of non-usable drugs and devices,
      - (iii) for pharmacy security,

- (iv) for emergency preparedness, and
- (v) for drug recall of pharmacy inventory;
- (d) ensure all policies and procedures are in writing and regularly maintained;
- (e) ensure that pharmacy staff are trained in policies and procedures;
- (f) ensure that all steps in the drug recall procedure are documented, if the procedure is initiated;
- (g) ensure that all individuals working in the pharmacy who present themselves as registrants have been granted and maintain registration with the College, in accordance with the policies approved by the board;
- (h) notify the registrar of any appointments, resignations or terminations of registrants employed at the pharmacy as those changes occur;
- (i) cooperate with inspectors acting under section 17 of the *Act* or section 28 or 29 of the *Health Professions Act*,
- (j) ensure that
  - registrant and support persons staff levels are commensurate with workload volumes and patient care requirements are met at all times in accordance with the bylaws, Code of Ethics and standards of practice, and
  - (ii) adequate and appropriate resources are in place to enable pharmacy staff to devote the time needed to conduct the continuous quality improvement and reporting activities required under section 24 or 29, as applicable, and
  - (iii) meeting quotas, targets or similar measures do not compromise patient safety or compliance with the bylaws, Code of Ethics or standards of practice;
- (k) ensure that all records related to the purchase and receipt of controlled drug substances are signed by a full pharmacist;
- ensure safe and secure storage of all Schedule I, II, and III drugs and controlled drug substances for all aspects of pharmacy practice, in accordance with the policies approved by the board;
- (m) ensure that pharmacy records containing personal information about patients are secure from unauthorized access, use, disclosure, modification and destruction;
- (n) ensure that each individual working in the pharmacy presents themselves to the public in a manner that clearly identifies their registration class;

- (o) ensure that registrants identify themselves in a manner that clearly differentiates them from other individuals working in the pharmacy who are not registrants;
- (p) immediately notify the registrar in writing of ceasing to be the pharmacy's manager;
- (q) ensure that at a minimum, the name on the external signage of a community pharmacy must be correctly and consistently used on labels and directory listings;
- (r) if the pharmacy is a central pharmacy, ensure that at a minimum, the name on the external signage of a telepharmacy must be correctly and consistently used on labels and directory listings;
- (s) ensure that narcotic reconciliation is performed in accordance with the policies approved by the board;
- notify the registrar of any incident of loss of narcotic and controlled drug substances within 24 hours;
- advise the registrar if the pharmacy is providing pharmacy services over the internet, and provide to the registrar the internet address of every website operated or used by the pharmacy;
- (v) ensure the pharmacy contains the reference material and equipment in accordance with the policies approved by the board;
- (w) require anyone who will access the in-pharmacy computer system to sign an undertaking in a form approved by the registrar to maintain the confidentiality of patient personal health information;
- (x) retain the undertakings referred to in subsection (w) in the pharmacy for 3 years after employment or any contract for services has ended;
- (y) provide the registrar with access to the pharmacy and premises as defined in section 20(1) in cases where a pharmacy licence has been cancelled or suspended due to loss of eligibility under section 3 of the *Act*,
- (z) ensure that no incentive is provided to a patient or patient's representative for the purpose of inducing the patient or patient's representative to
  - (i) deliver a prescription to a particular registrant or pharmacy for dispensing of a drug or device specified in the prescription, or
  - (ii) obtain any other pharmacy service from a particular registrant or pharmacy;

- (aa) notify the registrar of persistent non-compliance by a direct owner and indirect owner(s) with their obligations under the bylaws to the *Act*,
- (bb) notify the registrar of any change of telephone number, fax number, electronic mail address or any other information previously provided to the registrar;
- (cc) in the event of an anticipated temporary closure, which is permitted for no more than 14 consecutive days,
  - (i) notify patients and the public of the anticipated temporary closure at least 30 days prior to the start of the closure in accordance with the policies approved by the board,
  - (ii) document steps taken to comply with the bylaws and applicable policies on anticipated temporary closures,
  - (iii) contact all patients whose prepared prescriptions are ready for pick-up to advise of the closure and provide them with the opportunity to obtain their prepared prescriptions prior to the closure start date,
  - (iv) make alternate arrangements with local prescribers, as appropriate, and
  - (v) return any prepared prescriptions in the pharmacy to inventory and reverse those prescriptions in PharmaNet;
- (dd) in the event of an unanticipated temporary closure due to unforeseen circumstances, which is permitted for no more than 90 days,
  - (i) notify the registrar of closures of 15 to 90 days in accordance with the policies approved by the board,
  - (ii) where possible, contact all patients whose prescriptions are ready for pick-up to advise of the closure and provide them with the opportunity to obtain their prepared prescriptions,
  - (iii) where possible, notify patients, the public, and local prescribers of the closure and alternate means of obtaining essential pharmacy services during the closure in accordance with the policies approved by the board,
  - (iv) apply for a new pharmacy licence if the closure will exceed 90 days, and
  - (v) return any prepared prescriptions in the pharmacy to inventory and reverse those prescriptions in PharmaNet;
- (ee) in the event of a permanent pharmacy closure, cancellation, or expiry of the pharmacy licence

- (i) provide for the safe and secure transfer and appropriate storage of all Schedule I, II, and III drugs and controlled drug substances,
- advise the registrar in writing of the disposition of all drugs and prescription records at the time of a closure, in accordance with policies approved by the board,
- (iii) provide the registrar with a copy of the return invoice and any other documentation sent to Health Canada in respect of the destruction of all controlled drug substances,
- (iv) arrange for the secure transfer and continuing availability of the prescription records at another pharmacy, or at storage facility that is monitored and secured from unauthorized access, and
- (v) remove all signs and advertisements from the closed pharmacy premises;
- (3) In the event of a suspension of the pharmacy licence for a period of more than 14 days,
  - (a) the manager and the direct owner must complete and submit Form 4C, and
  - (b) the registrar may direct a manager to do any of sections 18(2)(ee)(i), (iii) or (iv).
- Subsection (2)(z) does not prevent a manager, direct owner or indirect owner(s) from
  - (a) providing free or discounted parking to patients or patient's representatives,
  - (b) providing free or discounted delivery services to patients or patient's representatives, or
  - (c) accepting payment for a drug or device by a credit or debit card that is linked to an incentive.
- (5) Subsection (2)(z) does not apply in respect of a Schedule III drug or an unscheduled drug, unless the drug has been prescribed by a practitioner.
- (6) A pharmacy education site's manager must ensure that only registrants and instructors are present in the pharmacy education site and must also comply with subsections (2)(a), (b), (c)(ii), (d), (e), (i), (p), (ee)(i) and (ee)(ii).
- (7) A direct owner, directors and officers must do all of the following:
  - (a) ensure compliance with subsections (2)(c)(i), (c)(iii), (c)(iv), (c)(v), (i), (j), (l), (q), (r), (y) and (z) and, as applicable, section 24 or 29;

- (b) ensure that the requirements to hold a pharmacy licence under the *Act* are met at all times; and
- notify the registrar of any change of name, address, telephone number, electronic mail address or any other information previously provided to the registrar;
- (8) Shareholders must comply with subsections (2)(i) and (7)(c).
- (9) A direct owner, manager, directors, and officers must ensure compliance with the National Association of Pharmacy Regulatory Authorities standards as approved by the board from time to time, applicable to the operation of a pharmacy.

# Sale and Disposal of Drugs

- 19 (1) Schedule I, II, and III drugs and controlled drug substances must only be sold or dispensed from a pharmacy.
  - (2) A registrant must not sell or dispense a quantity of drug that will not be used completely prior to the manufacturer's expiry date, if used according to the directions on the label.
  - (3) If the manufacturer's expiry date states the month and year but not the date, the expiry date is the last day of the month indicated.
  - (4) Every registrant practising in a pharmacy is responsible for the protection from loss, theft or unlawful sale or dispensing of all Schedule I, II, and III drugs and controlled drug substances in or from the pharmacy.
  - (5) A registrant must not sell, dispense, dispose of or transfer a Schedule I drug except
    - (a) on the prescription or order of a practitioner,
    - (b) for an inventory transfer to a pharmacy by order of a registrant in accordance with the policies approved by the board,
    - (c) by return to the manufacturer or wholesaler of the drug, or
    - (d) by destruction, in accordance with the policies approved by the board.
  - (6) Drugs included in the controlled prescription program must not be sold or dispensed unless
    - (a) the registrant has received the prescription on the prescription form approved by both the board and the College of Physicians and Surgeons of British Columbia, and
    - (b) the prescription form is signed by the patient or the patient's representative upon receipt of the dispensed drug.
  - (6.1) Despite subsection (6), a registrant may dispense drugs included in the controlled prescription program upon receipt of a verbal prescription from a practitioner if

doing so is permitted under a section 56 exemption to the *Controlled Drugs and Substances Act.* The pharmacy must receive the original prescription form, or a copy of the completed form transmitted by facsimile, from the practitioner as soon as reasonably possible.

- (7) A new prescription from a practitioner is required each time a drug is dispensed, except for
  - (a) a part-fill,
  - (b) a prescription authorizing repeats,
  - (c) a full pharmacist-initiated renewal or adaptation, or
  - (d) an emergency supply for continuity of care.
- (8) Subsection (6) does not apply to prescriptions written for
  - (a) residents of a facility or home subject to the requirements of the *Residential Care Facilities and Homes Standards of Practice*, or
  - (b) patients admitted to a hospital.

# **Drug Procurement/Inventory Management**

20 (1) In this section:

#### "premises" means:

- (a) a hospital as defined in the *Hospital Act*, or
- (b) the building or part of the building, within which the pharmacy is located, and includes loading spaces and excludes other businesses in the building.
- (2) A full pharmacist may authorize the purchase of Schedule I, II, or III drugs or controlled drug substances only from
  - (a) a wholesaler or manufacturer licensed to operate in Canada, or
  - (b) another pharmacy in accordance with the policies approved by the board.
- (3) A registrant must record a transfer of drugs that occurs for any reason other than for the purpose of dispensing in accordance with a practitioner's prescription.
- (4) All drug shipments must be delivered unopened to
  - (a) the pharmacy, or
  - (b) an area of the premises other than the pharmacy if the storage of the drug shipment is temporary, safe and secure.
- (5) Non-usable and expired drugs must be stored in the pharmacy in an area separate from other pharmacy stock or drug products until final disposal.

(6) A full pharmacist must not purchase Schedule I, II and III drugs and controlled drug substances unless they are for sale or dispensing in or from a pharmacy.

### Interchangeable Drugs

21 When acting under section 25.91 of the *Health Professions Act*, a full pharmacist must determine interchangeability of drugs by reference to Health Canada's Declaration of Equivalence, indicated by the identification of a Canadian Reference Product in a Notice of Compliance for a generic drug.

### **Returned Drugs**

22 No registrant may accept for return to stock or reuse any drug previously dispensed except in accordance with section 11(3) of the *Residential Care Facilities and Homes Standards of Practice,* section 5(2) of the *Hospital Pharmacy Standards of Practice,* or section 5 of the *Dispensing Drugs for the Purpose of Medical Assistance in Dying Standards, Limits and Conditions.* 

### Records

- 23 (1) All prescriptions, patient records, invoices and documentation in respect of the purchase, receipt or transfer of Schedule I, II and III drugs and controlled drug substances must be retained for a period of not less than three years from the date
  - (a) a drug referred to in a prescription was last dispensed, or
  - (b) an invoice was received for pharmacy stock.
  - (2) Despite subsection (1), a registrant must not destroy prescriptions, patient records, invoices and documentation as described in subsection (1) until the completion of any audit or investigation for which the registrant has received notice.
  - (3) Registrants, support persons, managers, direct owners, and indirect owners must not, for commercial purposes, disclose or permit the disclosure of information or an abstract of information obtained from a prescription or patient record which would permit the identity of the patient or practitioner to be determined.
- 23.1 (1) All records required to be kept under bylaws of the College or other legislation that regulates the practice of pharmacy shall be readable, complete, filed systematically and maintained in a manner that is secure, auditable and allows for easy retrieval.
  - (2) Notwithstanding subsection (1), a prescription record that is valid must be retrievable immediately.
  - (3) For the purpose of subsection (2):
    - (a) a prescription is valid for a period of up to two years from the prescribing date, unless the prescription is for a benzodiazepine or other targeted substance, in which case the prescription is valid for a period of up to one year from the prescribing date, and

- (b) despite paragraph (a), a prescription for a benzodiazepine or other targeted substance is valid for a period of up to two years from the prescribing date, if permitted by a section 56 exemption to the *Controlled Drugs and Substances Act.*
- (4) With respect to prescriptions for drugs included in the controlled prescription program, the original prescription form or a paper copy of the completed form transmitted by facsimile must be retained, regardless of whether or not such prescription form has also been stored electronically.
- (5) Prescriptions stored electronically must accurately reflect the original prescription, including the original colour composition of that prescription.
- 23.2 (1) A pharmacy manager must ensure that a policy is in place that:
  - (a) describes the pharmacy's records filing system, the records format and the method and system for storing records;
  - (b) is compliant with the sections 23.1, 23.2 and 23.3 requirements; and
  - (c) is readily accessible to and understood by pharmacy staff.
  - (2) With respect to electronic records, the policy must include a description of the process for the preservation, storage and backing up of records that is compliant with section 23.3 requirements.
- 23.3 (1) A pharmacy may maintain electronic records containing personal health information if the pharmacy has the equipment, software and systems necessary for the input, storage, use, protection and retrieval of records that are required to be kept under bylaws of the College or other legislation that regulates the practice of pharmacy.
  - (2) For purposes of subsection (1), the equipment, software and systems must:
    - (a) be capable of storing the electronic records for the periods required by applicable law;
    - (b) keep the records secure from unauthorized access, use, disclosure, modification and destruction;
    - (c) for audit purposes, be capable of uniquely identifying each time an electronic record is accessed and modified;
    - (d) be capable of restricting the functions that may be used by an authorized person;
    - (e) be capable of tracing alterations to records by identifying the original entry, the identity of the individual who made the alteration and the date of the alteration;

- (f) be capable of searching and sorting electronic prescription records chronologically, and by drug name, drug strength, patient, prescriber, prescription number and transaction number;
- (g) ensure that electronic records can be stored, backed up and recovered in accordance with subsection (3); and
- (h) provide for a deliberate and auditable procedure to be carried out by the pharmacy manager or by an authorized person prior to the destruction of any electronic record that includes information identifying the pharmacy manager or authorized person who destroyed the record and the date, time and reason for its destruction.
- (3) A pharmacy manager must ensure that electronic records are preserved and backed up at least once daily and that such electronically preserved and backed up records are stored:
  - (a) in a location resistant to environment perils including but not limited to fires and floods;
  - (b) so that they are secure from unauthorized access, use, modification, destruction and disclosure; and
  - (c) in a manner that would enable the backed up records, once restored, to be compliant with section 23.1(1) requirements.
- Notwithstanding subsections (1), (2) and (3), a pharmacy that presently stores electronic records has six months from the date this section comes into effect to bring itself into full compliance with the requirements of subsections (1), (2) and (3).

# Signage within the Pharmacy

- 23.4 (1) A pharmacy must clearly display, within the pharmacy, the signage provided by the College for the purpose of this section.
  - (2) The signage referred to in subsection (1) may be placed within the dispensary area but in any case must, at all times when patients or the public may be present in the pharmacy.
    - (a) be placed in a manner and location that makes it
      - (i) clearly visible and readable to members of the public from every consultation area or counter where a member of the public can obtain a full pharmacist's advice, and
      - (ii) visually distinctive from other signage.
    - (b) be in good condition or be displayed on an electronic sign that is in good working condition, and

- (c) comply with any applicable colour or minimum size requirements or other technical specifications, or additional criteria respecting manner or location of placement, approved by the board from time to time for the purpose of this section.
- (3) The following pharmacies are exempt from the requirements in subsections (1) and (2):
  - (a) a community pharmacy or telepharmacy, if it is never open to the public and has no external signage identifying it as a pharmacy;
  - (b) a hospital pharmacy, if pharmacy services are never provided in or from it on an outpatient basis;
  - (c) a pharmacy education site.

# **PART III – Community Pharmacies**

#### **Community Pharmacy's Manager – Quality Management**

- 24 (1) A community pharmacy's manager must establish and maintain written quality management policies and procedures for the community pharmacy that
  - (a) ensure pharmacy staff, equipment, and facilities comply with all legislation, bylaws and policies applicable to the operation of a community pharmacy,
  - (b) include a continuous quality improvement program for the pharmacy,
  - (c) include a process for reporting, documenting, analyzing, following up on, and learning from medication incidents and near misses in the pharmacy, and
  - (bd) include a process to monitor compliance with the quality management policies and procedures<del>, and</del>.
  - (c) include a process for reporting, documenting and following up on known, alleged and suspected errors, incidents and discrepancies.
  - (2) The policies and procedures for the program referred to in subsection (1)(b) must include adequate and appropriate processes for all of the following:
    - (a) identifying root causes and contributing factors for medication incidents and near misses and performing a root-cause analysis as appropriate;
    - (b) reviewing and assessing summary reports and analyses of pharmacyspecific data;
    - (c) reviewing and assessing objective analyses from available regional-, provincial-, and national-level data;
    - (d) holding team meetings on a routine basis;

- (e) completing safety self-assessments on a routine basis;
- (f) following up with team members involved in medication incidents and near misses and encouraging them to seek peer support when appropriate;
- (g) ensuring that pharmacy policies and procedures are reviewed and updated based on the pharmacy's root-cause analyses, safety selfassessments, summary reports and analyses, and objective analyses from pharmacy-specific data and available regional-, provincial-, and national-level data;
- (h) implementing improvements to the pharmacy's procedures in accordance with the pharmacy's continuous quality improvement plan;
- (i) developing a monitoring process to determine the efficacy of implemented improvements to the pharmacy's procedures;
- (j) implementing further updates to the pharmacy's procedures if previous improvements are not effective.
- (3) After June 1, 2026, a community pharmacy's manager must ensure that a team meeting is held for the pharmacy
  - (a) at least once within the one-year period immediately following that date if, on that date, the pharmacy is operating under a valid pharmacy license issued, renewed or reinstated on or before that date, and
  - (b) at least once within the one-year period immediately following each subsequent anniversary of that date if, on such anniversary date, the pharmacy is operating under a valid pharmacy license issued, renewed or reinstated on or before such anniversary date,

but in any case, not later than one year after the date of the last preceding team meeting, if any, held for the pharmacy as required under paragraph (a) or (b), as applicable.

- (4) A community pharmacy's manager must ensure that that all team meetings held as required under subsection (3) are adequately documented in the pharmacy's records including documentation of all of the following for each team meeting:
  - (a) date of the meeting;
  - (b) names of the pharmacy staff members in attendance;
  - (c) topics of discussion;
  - (d) any resulting improvement plans.

- (5) After June 1, 2026, and subject to subsection (6), a community pharmacy's manager must ensure that a safety self-assessment is completed for the pharmacy
  - (a) at least once within the one-year period immediately following that date if, on that date, the pharmacy was operating under a valid pharmacy license issued, renewed or reinstated on or before that date, and
  - (b) at least once within the one-year period immediately following each subsequent anniversary of that date if, on such anniversary date, the pharmacy is operating under a valid pharmacy license issued, renewed or reinstated on or before such anniversary date.
- (6) After the initial safety self-assessment has been completed for a pharmacy as required under subsection (5)(a) or (b), as applicable, further safety self-assessments are required to be completed for the pharmacy only once within each successive three-year period following the date such first safety self-assessment is completed, but in any case, not later than three years after the date of the last preceding safety self-assessment completed for the pharmacy as required under subsection (5).
- (7) A community pharmacy's manager must ensure that all safety self-assessments required under subsection (5) are completed using a software tool or platform that is appropriate for the pharmacy's licence type and that has the necessary technical features and capabilities to consistently and reliably generate adequate safety self-assessments for the pharmacy, such as the Medication Safety Self-Assessment solutions offered by Institute for Safe Medication Practices Canada, the Pharmacy Safety Self-Assessment tool within the Pharmapod software solution offered by Think Research Corporation, or an equivalent software tool or platform.
- (8) The policies and procedures for the process referred to in subsection (1)(c) must clearly set out
  - (a) the steps that pharmacy staff must take when a medication incident or near miss occurs, including the steps for disclosure, and
  - (b) the pharmacy's criteria for determining whether a near miss must be reported to a national database, which at a minimum must include the following criteria:
    - (i) if the event had reached the patient, it is likely that actual harm would have been caused;
    - (ii) the event has been a recurring problem for the pharmacy;
    - (iii) the event provides a learning opportunity for the pharmacy specifically or for pharmacy practice in general.

- (9) A community pharmacy's manager must select, implement, and maintain a reporting platform that
  - (a) has processes in place to de-identify data, ensuring there are no patient or pharmacy staff identifiers once data leaves the reporting platform,
  - (b) is able to integrate with a national database to share de-identified medication incident and near miss reports, and
  - (c) at a minimum, enables and requires pharmacy staff to enter information for all of the following data fields, and includes all these data fields in all reports submitted to a national database:
    - (i) date incident occurred;
    - (ii) type of medication incident;
    - (iii) incident discovered by (position/job title only);
    - (iv) medication system stages involved in this incident;
    - (v) medication(s) involved;
    - (vi) degree of harm to patient due to incident;
    - (vii) incident description/how the incident was discovered;
    - (viii) contributing factors of the incident.
- (210) If a community pharmacy is a central pharmacy, the quality management policies and procedures in required by subsection (1) must include and apply to all telepharmacies associated with the central pharmacy and must comply with the Telepharmacy Standards of Practice.
- (11) In this section:

"**contributing factor**" means a circumstance, action or influence that is thought to have played a part in the origin or development of a medication incident or near miss, or to increase the risk of a medication incident or near miss;

"culture of patient safety" means the component of organizational culture involving the shared beliefs, attitudes, values, norms and behavioural characteristics of employees, and that influences staff member attitudes and behaviours in relation to their organization's ongoing patient safety performance, resulting in an enabling patient safety culture characterized by leadership that leads by example, transparent communication, psychological safety facilitating reporting of errors, patient and family engagement, and a commitment to ongoing improvement;

"**de-identify**" and "**de-identified**", in relation to data and reports referred to in this section, means the data or report does not include any information that could be

used to identify the individual who completed or submitted the report, any pharmacy staff involved in the medication incident or near miss, any patient or patient's representative, or any other individual;

"just culture" means the environment of a workplace in which consideration is given to wider systemic issues when things go wrong, enabling professionals and those operating the system to learn without fear of retribution, and where, to encourage reporting of safety issues, inadvertent human error, freely admitted, is generally not subject to sanction, but people are held to account where there is evidence of unprofessional conduct or deliberate acts;

"medication incident" means any preventable event that may cause or lead to inappropriate medication use or patient harm that has reached the patient and that may be related to professional practice, drug products, procedures, and systems, and include prescribing, order communication, product labelling/packaging/nomenclature, compounding, dispensing, distribution, administration, education, monitoring, and use;

"national database" means a data repository that

- (a) contains de-identified medication incident and near miss reporting data submitted from across Canada,
- (b) is appropriate for the pharmacy's licence type, and
- is an established component of the collaborative pan-Canadian program for reducing and preventing harmful medication incidents known as the Canadian Medication Incident Reporting and Prevention System (CMIRPS);

"**near miss**" means an event that could have resulted in unwanted consequences but did not because, either by chance or through timely intervention, the event did not reach the patient;

"**peer support**" means emotional and practical support between two people who share a common experience, such as a mental health challenge or illness;

"**reporting platform**" means the computer software used by pharmacy staff for recording medication incidents and near misses at the pharmacy level and reporting them to a national database;

"**root cause**" means the most fundamental reason, or one of several fundamental reasons, a suspected failure, a medication incident, a near miss, or a situation in which performance does not meet expectations has occurred.

"**root cause analysis**" means an objective analytical process that can be used to perform a comprehensive, system-based review of critical incidents including without limitation the identification of the root and contributory factors, the determination of risk reduction strategies, and the development of action plans along with measurement strategies to evaluate the effectiveness of the plans; "**safety self-assessment**" means a process used regularly by pharmacy staff to proactively identify potential safety concerns, which may help decrease the number of medication incidents and near misses and identify opportunities for improvement at a pharmacy in order to mitigate risks to patients;

"**team meeting**" means a regular meeting of pharmacy staff to proactively review and assess summary reports and analyses of pharmacy-specific data and available summary reports and analyses of available regional-, provincial-, and national-level data and determine how to address them.

### **Community Pharmacy and Telepharmacy Premises**

- 25 (1) In locations where a community pharmacy or telepharmacy does not comprise 100 per cent of the total area of the premises, the community pharmacy manager or the central pharmacy manager in the case of a telepharmacy, must ensure that
  - (a) the professional products area extends not more than 25 feet from the perimeter of the dispensary and is visually distinctive from the remaining areas of the premises by signage, and
  - (b) a sign reading "Medication Information" is clearly displayed to identify a consultation area or counter at which a member of the public can obtain a full pharmacist's advice.
  - (2) Subject to subsection (3), the dispensary area of a community pharmacy or a telepharmacy must
    - (a) be at least 160 square feet,
    - (b) be inaccessible to the public by means of gates or doors across all entrances,
    - (c) include a dispensing counter with at least 30 square feet of clear working space, in addition to service counters,
    - (d) contain adequate shelf and storage space that is clean and organized,
    - (e) contain a double stainless steel sink with hot and cold running water,
    - (f) contain an adequate stock of drugs to provide full dispensing services, and
    - (g) contain a refrigerator.
  - (3) A telepharmacy that was authorized by the registrar to provide pharmacy services as a telepharmacy remote site as of January 1, 2017 is exempt from the requirements in subsections (2)(a) and (c) until such time as it commences a renovation of all or part of the premises.
  - (4) In all new and renovated community pharmacies or telepharmacies, an appropriate area must be provided for patient consultation that

- (a) ensures privacy and is conducive to confidential communication, and
- (b) includes, but is not limited to, one of the following:
  - (i) a private consultation room, or
  - (ii) a semiprivate area with suitable barriers.

### **Community Pharmacy and Telepharmacy Security**

- 26 (1) A community pharmacy or telepharmacy must:
  - (a) keep Schedule IA drugs in a locked metal safe inside the dispensary that is secured in place and equipped with a time delay lock set at a minimum of five minutes;
  - (b) install and maintain a security camera system that:
    - (i) has date/time stamp images that are archived and available for no less than 30 days; and
    - (ii) is checked daily for proper operation; and
  - (c) install and maintain motion sensors in the dispensary.
  - (2) When no full pharmacist is present and the premises in which the pharmacy is located are accessible to non-registrants, the pharmacy must be secured as follows:
    - (a) if the premises in which the pharmacy is located are closed and accessible to non-registrant staff:
      - (i) the dispensary area must be secured by a monitored alarm; and
      - (ii) subject to subsection (2.1), Schedule I and II drugs, controlled drug substances and personal health information, are secured by physical barriers; or
    - (b) if the pharmacy is closed but other areas of the premises in which the pharmacy is located are open:
      - (i) the dispensary area must be secured by a monitored alarm;
      - (ii) subject to subsection (2.1), Schedule I, and II drugs, controlled drug substances and personal health information, are secured by physical barriers; and
      - (iii) Schedule III drugs are inaccessible to anyone other than full pharmacists, temporary pharmacists and pharmacy technicians.
  - (2.1) A community pharmacy or telepharmacy that exists on the date this provision comes into force and is not renovated during the period must comply with

sections 26(2)(a)(ii) and (b)(ii) no later than three years after the date that provision comes into force.

- (2.2) For the purposes of subsection (2), a full pharmacist is deemed to be present at a telepharmacy when he or she is engaged in direct supervision of the telepharmacy.
- (3) Subject to subsection (5), a community pharmacy or a telepharmacy must clearly display at all external entrances that identify the premises as a pharmacy, and at the dispensary counter signage provided by the College.
- (4) The manager, direct owner or indirect owner(s) of a community pharmacy or telepharmacy that does not stock IA drugs must complete a declaration attesting that Schedule IA drugs are never stocked on the premises.
- (5) A pharmacy that is never open to the public and has no external signage identifying it as a pharmacy is exempt from the requirements in subsection (3).

### Permitted Activities of a Community Pharmacy without a Full Pharmacist Present

- 27 (1) Except as provided in subsection (2), a community pharmacy must not operate unless a full pharmacist is present.
  - (2) A community pharmacy may carry on the activities set out in subsection (3) without a full pharmacist present only if:
    - (a) the registrar is notified of the hours during which a full pharmacist is not present;
    - (b) the pharmacy is secured in accordance with section 26(2); and
    - (c) the hours when a full pharmacist is on duty are posted.
  - (3) Subject to subsection (2) if a full pharmacist is not present, only the following activities may be carried out:
    - (a) pharmacy technicians may access the dispensary to perform activities outlined in section 4 of the *Community Pharmacy Standards of Practice*, that do not require pharmacist supervision, except if any such activity involves patient interaction; and
    - (b) receive drug shipments under section 20(4).
  - (4) Nothing contained in this section relieves a pharmacy manager of their responsibilities under section 18(2)(a).

#### **Outsource Prescription Processing**

- 28 (1) A community pharmacy may outsource prescription processing if
  - (a) all locations involved in the outsourcing are community pharmacies,

- (b) all prescriptions dispensed are labeled and include an identifiable code that provides a complete audit trail for the dispensed drug, and
- (c) a notice is posted informing patients that the preparation of their prescriptions may be outsourced to another pharmacy.
- (2) The manager of an outsourcing community pharmacy must ensure that all applicable standards of practice are met in processing prescriptions at all locations involved in the outsourcing.
- (3) In this section, "community pharmacy" includes a hospital pharmacy.

### **PART IV – Hospital Pharmacies**

#### Hospital Pharmacy's Manager – Quality Management

- 29 (1) A hospital pharmacy's manager must establish and maintain written quality management policies and procedures for the hospital pharmacy that
  - (a) ensure pharmacy staff, equipment, and facilities comply with all legislation, bylaws and policies applicable to the operation of a hospital pharmacy,
  - (b) include a continuous quality improvement program for the pharmacy,
  - (c) include a process for reporting, documenting, analyzing, following up on, and learning from medication incidents and near misses in the pharmacy,
  - (bd) include a process to monitor compliance with the quality management policies and procedures,
  - (c) include a process for reporting, documenting and following up on known, alleged and suspected errors, incidents and discrepancies,
  - (de) document periodic audits of the drug distribution process,
  - (ef) include a process to review patient-oriented recommendations,
  - (fg) include a process that reviews a full pharmacist's documentation notes in the hospital's medical records,
  - (gh) include a process to evaluate drug use, and
  - (hi) regularly update policies and procedures for drug use control and patientoriented pharmacy services in collaboration with the medical and nursing staff and appropriate committees.
  - (2) Section 24(2) to (9) and (11) applies to a hospital pharmacy and the hospital pharmacy's manager as if they were, respectively, a community pharmacy and the community pharmacy's manager.

(23) If sample drugs are used within a hospital, the hospital pharmacy's manager must ensure that the pharmacy oversees the procurement, storage and distribution of all sample drugs.

### **After Hours Service**

- 30 (1) If continuous pharmacy services are not provided in a hospital, the hospital pharmacy's manager must ensure that urgently needed drugs and patient-oriented pharmacy services are available at all times by
  - (a) providing a cabinet which must
    - be a locked cabinet or other secure enclosure located outside of the hospital pharmacy, to which only authorized persons may obtain access,
    - (ii) be stocked with a minimum supply of drugs most commonly required for urgent use,
    - (iii) not contain controlled drug substances unless they are provided by an automated dispensing system,
    - (iv) contain drugs that are packaged to ensure integrity of the drug and labeled with the drug name, strength, quantity, expiry date and lot number, and
    - (v) include a log in which drug withdrawals are documented, and
  - (b) arranging for a full pharmacist to be available for consultation on an oncall basis.
  - (2) When a hospital pharmacy or hospital pharmacy satellite is closed, the premises must be equipped with a security system that will detect unauthorized entry.

# **PART V – Telepharmacies**

#### **Telepharmacy Operation**

- 31 (1) A telepharmacy must not remain open and prescriptions must not be dispensed without a full pharmacist physically present on duty at the telepharmacy, unless
  - (a) a full pharmacist at the central pharmacy is engaged in direct supervision of the telepharmacy in accordance with the *Telepharmacy Standards of Practice*, and
  - (b) subject to subsection (2), a pharmacy technician is physically present on duty at the telepharmacy.
  - (2) A telepharmacy located at an address listed in Schedule "G" is exempt from the requirements in subsection (1)(b).

- (3) A telepharmacy must have a security system that prevents the public and nonpharmacy staff from accessing the professional services area and the dispensary area, including any area where personal health information is stored.
- (4) Prescriptions and labels relating to prescriptions dispensed at a telepharmacy must identify the prescription as having been dispensed at that telepharmacy.
- (4.1) Prescriptions and labels relating to prescriptions dispensed at a pharmacy listed in Schedule "F" must distinguish between those dispensed when it is operating as a telepharmacy from when it is operating as a community pharmacy.
- (5) The manager of a central pharmacy, or a full pharmacist designated by the manager, must
  - (a) inspect and audit its telepharmacy at least 4 times each year, at intervals of not less than 2 months,
  - (b) record each inspection and audit in the prescribed form, and
  - (c) provide the inspection and audit records to the registrar immediately upon request.
- (6) A telepharmacy located at an address listed in Schedule "G" must perform a monthly count of narcotics at the telepharmacy and retain a record of each monthly count signed by the supervising pharmacist for three years at both the central pharmacy and the telepharmacy location, and provide the signed record to the registrar immediately upon request.
- (7) A telepharmacy must not continue to provide pharmacy services for more than 30 days after
  - (a) its location ceases to be a rural and remote community,
  - (b) a community pharmacy is established within the community, or
  - (c) a community pharmacy is established within 25 kilometers of the location of the telepharmacy.
- (8) In accordance with sections 18(2)(c) and (d), a telepharmacy must have policies and procedures on site that outline the methods for ensuring the safe and effective distribution of pharmacy products and delivery of pharmaceutical care by the telepharmacy.
- (9) All transactions in PharmaNet must be distinguishable between the central pharmacy and telepharmacy.

# PART VI – PharmaNet

### **Application of Part**

32 This Part applies to every pharmacy that connects to PharmaNet.

#### Definitions

33 In this Part:

"**patient record**" means the patient record described in section 11(2) of the *Community Pharmacy Standards of Practice* and in the *British Columbia Professional and Software Conformance Standards, Electronic Health Information Exchange* as the "patient record (pharmacy)".

"**PharmaNet**" means "PharmaNet" as defined in section 1 of the *Information Management Regulation*, B.C. Reg. 74/2015;

#### **Operation of PharmaNet**

34 A pharmacy must connect to PharmaNet.

#### Data Collection, Transmission of and Access to PharmaNet Data

- 35 (1) A registrant must enter the prescription information and record it in PharmaNet at the time of dispensing and keep the patient record current.
  - (2) A registrant may collect and record patient information in PharmaNet, or access, use and disclose a patient's PharmaNet record only for the purposes of:
    - (a) dispensing a drug;
    - (b) providing patient consultation;
    - (c) evaluating a patient's drug usage;
    - (d) claims adjudication and payment by an insurer; or
    - (e) providing pharmacy services to, or facilitating the care of, the individual whose personal information is being collected, accessed, used or disclosed.
  - (3) A registrant must revise information in PharmaNet pertaining to corrected billings for prescriptions billed to the patient or a payment agency other than PharmaCare and record the reason for the revision within 120 days of the original entry in PharmaNet.
  - (4) A registrant must reverse information in PharmaNet, for any drug that is not released to the patient or the patient's representative, and record the reason for the reversal no later than 30 days from the date of the original entry of the prescription information in PharmaNet.
  - (5) If a registrant is unable to comply with the deadlines in subsection (3) or (4), he or she must provide the information required to make the correction to the Ministry of Health as soon as possible thereafter.

# PART VII – Confidentiality

### Confidentiality

36 A registrant must take reasonable steps to confirm the identity of a patient, patient's representative, registrant or practitioner before providing any pharmacy service that requires accessing, using or disclosing of patient personal health information.

# PART VIII – College

### Forms

37 The registrar may establish forms for the purposes of the Act.

### Use, Disclosure and Retention of Criminal Record History Information

- 38 (1) The College may disclose criminal record history information only for the purpose of licensing pharmacies or for the purpose of regulating registrants (including for the discipline of registrants).
  - (2) The College must retain criminal record history information only for so long as is permitted by the applicable College records retention and disposal provisions established by the College.