



**Model State Pharmacy Act  
and Model Rules of the  
National Association of Boards of Pharmacy**

**August 2021**

# Mission Statement of the National Association of Boards of Pharmacy

## ***NABP Mission Statement***

The National Association of Boards of Pharmacy (NABP) is the independent, international, and impartial Association that assists its member boards in protecting the public health.

## ***Vision Statement***

Innovating and collaborating today for a safer public health tomorrow.

## ***NABP Member Boards of Pharmacy***

Alabama State Board of Pharmacy  
Alaska Board of Pharmacy  
Arizona State Board of Pharmacy  
Arkansas State Board of Pharmacy  
California State Board of Pharmacy  
Colorado State Board of Pharmacy  
Connecticut Commission of Pharmacy  
Delaware State Board of Pharmacy  
District of Columbia Board of Pharmacy  
Florida Board of Pharmacy  
Georgia State Board of Pharmacy  
Guam Board of Examiners for Pharmacy  
Hawaii State Board of Pharmacy  
Idaho State Board of Pharmacy  
Illinois Department of Financial and Professional Regulation, Division of Professional Regulation – State Board of Pharmacy  
Indiana Board of Pharmacy  
Iowa Board of Pharmacy  
Kansas State Board of Pharmacy  
Kentucky Board of Pharmacy  
Louisiana Board of Pharmacy  
Maine Department of Professional and Financial Regulation, Office of Professional and Occupational Regulation – Board of Pharmacy  
Maryland Board of Pharmacy

Massachusetts Board of Registration in Pharmacy  
Michigan Board of Pharmacy  
Minnesota Board of Pharmacy  
Mississippi Board of Pharmacy  
Missouri Board of Pharmacy  
Montana Board of Pharmacy  
Nebraska Department of Health and Human Services, Division of Public Health, Licensure Unit  
Nevada State Board of Pharmacy  
New Hampshire Board of Pharmacy  
New Jersey State Board of Pharmacy  
New Mexico Board of Pharmacy  
New York State Board of Pharmacy  
North Carolina Board of Pharmacy  
North Dakota State Board of Pharmacy  
State of Ohio Board of Pharmacy  
Oklahoma State Board of Pharmacy  
Oregon State Board of Pharmacy  
Pennsylvania State Board of Pharmacy  
Puerto Rico Board of Pharmacy  
Rhode Island Board of Pharmacy  
South Carolina Department of Labor, Licensing, and Regulation – Board of Pharmacy  
South Dakota State Board of Pharmacy  
Tennessee Board of Pharmacy

Texas State Board of Pharmacy  
Utah Board of Pharmacy  
Vermont Board of Pharmacy  
Virgin Islands Board of Pharmacy  
Virginia Board of Pharmacy  
Washington State Pharmacy Quality Assurance Commission  
West Virginia Board of Pharmacy  
Wisconsin Pharmacy Examining Board  
Wyoming State Board of Pharmacy

### ***Bahamas:***

Bahamas Pharmacy Council\*

### ***Canada:***

Alberta College of Pharmacy\*  
College of Pharmacists of British Columbia\*  
College of Pharmacists of Manitoba\*  
New Brunswick College of Pharmacists\*  
Newfoundland and Labrador Pharmacy Board\*  
Nova Scotia College of Pharmacists\*  
Ontario College of Pharmacists\*  
Prince Edward Island College of Pharmacists\*  
Quebec Order of Pharmacists\*  
Saskatchewan College of Pharmacy Professionals\*

\* Associate Member

# **Model State Pharmacy Act and Model Rules of the National Association of Boards of Pharmacy**

**August 2021**

**Published by:**

National Association of Boards of Pharmacy  
1600 Feehanville Drive  
Mount Prospect, IL 60056  
847/391-4406

Lemrey “Al” Carter, PharmD, MS, RPh  
Executive Director/Secretary

© 2021, National Association of Boards of Pharmacy. All rights reserved.

No part of this publication may be reproduced in any manner without the written permission of the executive director/secretary of the National Association of Boards of Pharmacy. Violation of the copyright will lead to prosecution under federal copyright laws.

ARTICLE I TITLE, PURPOSE, AND DEFINITIONS .....	88
<i>Introductory Comment to Article I</i> .....	88
Section 101. Title of Act .....	88
Section 102. Legislative Declaration .....	88
Section 103. Statement of Purpose.....	88
Section 104. Practice of Pharmacy.....	99
Section 105. Definitions.....	99
Section 105(l2). Comment.....	3434
Section 105(z5). Comment.....	3434
ARTICLE II BOARD OF PHARMACY .....	3939
<i>Introductory Comment to Article II</i> .....	3939
Section 201. Designation.....	3939
Section 202. Membership.....	4040
Section 203. Qualifications.....	4040
Section 204. Appointment.....	4040
Section 205. Terms of Office.....	4141
Section 206. Vacancies.....	4141
Section 207. Removal.....	4141
Section 208. Organization.....	4141
Section 209. Compensation of Board Members.....	4242
Section 210. Meetings.....	4242
Section 211. Employees.....	4242
Section 212. Rules.....	4242
Section 213. Powers and Responsibilities.....	4343
ARTICLE III LICENSING .....	4747
<i>Introductory Comment to Article III</i> .....	4747
Section 301. Unlawful Practice.....	4747
Section 302. Qualifications for Licensure by Examination.....	4848
Section 303. Qualifications for Licensure Transfer.....	4949
Section 304. Renewal of Licenses.....	5050
Section 305. Continuing Pharmacy Education.....	5050
Section 306. Pharmacy Practice Experience Program Standards; Pharmacy Intern Licensure.....	5050
Section 307. Licensure of Certified Pharmacy Technicians.....	5151
Section 308. Licensure of Certified Pharmacy Technician Candidates.....	5151
ARTICLE IV DISCIPLINE .....	5353
<i>Introductory Comment to Article IV</i> .....	5353
Section 401. Disciplinary Action Terms.....	5353
Section 402. Grounds, Penalties, and Reinstatement.....	5353
Section 403. Procedure.....	5656
ARTICLE V LICENSING OF FACILITIES .....	5858
<i>Introductory Comment to Article V</i> .....	5858
Section 501. Licensing.....	5858
Section 502. Application.....	6060
Section 503. Notifications.....	6161
Section 504. Grounds, Penalties, and Reinstatement.....	6161
Section 505. Criminal Offense; Forfeiture of Property.....	6363
ARTICLE VI OTHER .....	6464
Section 601. Severability.....	6464
Section 602. Effective Date.....	6464
MODEL RULES FOR PHARMACY INTERNS .....	6565
Section 1. Licensure.....	6565
Section 2. Identification.....	6565
Section 3. Supervision.....	6565
Section 4. Change of Address.....	6666
Section 5. Evidence of Completion.....	6666
MODEL STANDARDS FOR PHARMACY PRACTICE EXPERIENCE PROGRAMS.....	6767

Section 1. Preceptor.....	6767
Section 2. Pharmacy Practice Experience Programs.....	6767
Section 3. Global Exchange Pharmacy Students.....	6767
MODEL RULES FOR INSTITUTIONAL PHARMACY.....	6868
Section 1. Applicability.....	6868
Section 2. Absence of Pharmacist at a Pharmacy Located Within an Institutional Facility.....	6868
Section 3. Emergency Kit Use by Institutional Facilities.....	6868
Section 4. Drug Distribution and Pharmacist Care Services.....	6969
Section 5. Shared Pharmacy Services Utilization for Immediate Need.....	6969
Section 6. Packaging of Previously Dispensed Medication.....	7070
Section 7. Institutional Pharmacy Delivery Room.....	7070
MODEL RULES FOR THE PRACTICE OF PHARMACY.....	7171
Introductory Comment.....	7171
Section 1. Facility.....	7171
Section 2. Security.....	7272
Section 3. Personnel.....	7272
Section 4. Prescription Drug Order Processing.....	7676
Section 5. Record Keeping.....	8181
Section 6. Pharmacist Care Services.....	8585
Section 7. Continuous Quality Improvement Program.....	8787
Section 8. Shared Pharmacy Services.....	8888
Section 9. Automated Pharmacy Systems.....	9191
Section 10. Return and Reuse of Prescription Drugs.....	9393
Section 11. Prescription Drug Repository Programs.....	9393
Section 12. Disposal of Controlled Substances.....	9494
Section 13. Prepackaging.....	9494
Section 14. Provision of Pharmacist Care Services Outside of a Licensed Pharmacy.....	9595
Section 15. Approval of Pharmacy Practice Initiatives.....	9595
Section 16. Unprofessional Conduct.....	9696
MODEL RULES FOR PUBLIC HEALTH EMERGENCIES.....	9797
Section 1. Purpose and Scope.....	9797
Section 2. Definitions.....	9797
Section 3. Emergency Prescription Drug Order.....	9898
Section 4. Public Health Emergency Refill Dispensing.....	9898
Section 5. Temporary Recognition of Nonresident Licensure.....	9999
Section 6. Temporary Pharmacy Facilities or Mobile Pharmacies.....	100+00
MODEL RULES FOR NUCLEAR/RADIOLOGIC PHARMACY.....	101+01
Section 1. Purpose and Scope.....	101+01
Section 2. Definitions.....	101+01
Section 3. General Requirements for Pharmacies Providing Radiopharmaceutical Services.....	102+02
Section 4. Other Requirements.....	103+03
MODEL RULES FOR COMPOUNDED OR REPACKAGED PHARMACEUTICALS.....	104+04
Section 1. Purpose and Scope.....	104+04
Section 2. Notification.....	104+04
Section 3. Policy and Procedure Manual.....	104+04
Section 4. Physical Requirements.....	104+04
Section 5. Records and Reports.....	105+05
Section 6. Delivery Service.....	105+05
Section 7. Disposal of Hazardous and/or Infectious Wastes.....	105+05
Section 8. Quality Assurance.....	105+05
Section 9. Compounded Drug Preparations for Veterinary Use.....	106+06
MODEL RULES FOR OUTSOURCING FACILITIES.....	107+07
Section 1. Purpose and Scope.....	107+07
Section 2. Registration.....	107+07
Section 3. Notification.....	107+07
Section 4. Requirements.....	107+07

MODEL RULES FOR THE LICENSURE OF MANUFACTURERS, REPACKAGERS, THIRD-PARTY LOGISTICS PROVIDERS, AND WHOLESALE DISTRIBUTORS .....		108108
Section 1. Requirements for Licensure.....		108108
Section 2. Minimum Qualifications.....		110110
Section 3. Personnel.....		111111
Section 4. Minimum Requirements for the Storage and Handling of Prescription Drugs and for Establishment and Maintenance of Prescription Drug Records.....		113113
Section 5. Security.....		113113
Section 6. Storage.....		114114
Section 7. Operations/Reporting Requirements.....		114114
Section 8. Due Diligence.....		116116
Section 9. Record Keeping.....		116116
Section 10. Policies and Procedures.....		117117
Section 11. Prohibited Acts.....		118118
Section 12. Criminal Acts.....		119119
Section 13. Salvaging and Reprocessing.....		120120
Section 14. Inspection and Accreditation by a Third Party.....		120120
MODEL RULES FOR THE LICENSURE OF MEDICAL GAS AND MEDICAL GAS RELATED EQUIPMENT WHOLESALE DISTRIBUTORS.....		121121
Section 1. Definitions.....		121121
Section 2. Requirements for Licensure.....		124124
Section 3. Minimum Qualifications.....		126126
Section 4. Personnel.....		127127
Section 5. Minimum Requirements for the Storage and Handling of Medical Gases or Medical Gas Related Equipment and for Establishment and Maintenance of Medical Gas or Medical Gas Related Equipment Records.....		127127
Section 6. Security.....		128128
Section 7. Storage.....		129129
Section 8. Examination of Materials.....		129129
Section 9. Returned, Damaged, and Outdated Medical Gases or Medical Gas Related Equipment.....		129129
Section 10. Due Diligence.....		130130
Section 11. Record Keeping.....		130130
Section 12. Policies and Procedures.....		131131
Section 13. Prohibited Acts.....		132132
Section 14. Criminal Acts.....		132132
Section 15. Salvaging and Reprocessing.....		133133
Section 16. Inspection.....		133133
APPENDIX A MULTISTATE PHARMACY INSPECTION BLUEPRINT.....		134134
APPENDIX B SAMPLE PHARMACY AUTOMATION POLICY AND PROCEDURE OUTLINE.....		147147
APPENDIX C GUIDELINES FOR DISCIPLINARY SANCTIONS.....		149149
APPENDIX D COMMUNITY PHARMACY QUALITY-RELATED EVENT (QRE) DATA COLLECTION FORM .....		152152
COMMUNITY PHARMACY CONTINUOUS QUALITY IMPROVEMENT PROGRAM INSPECTION FORM.....		155155
COMMUNITY PHARMACY QUALITY SELF-AUDIT .....		157157
APPENDIX E MODEL PRESCRIPTION MONITORING PROGRAM ACT.....		163163
Section 1. Short Title.....		163163
Section 2. Legislative Findings.....		163163
Section 3. Purpose.....		163163
Section 4. Definitions.....		163163
Section 5. Establishment of a Prescription Monitoring Program.....		164164
Section 6. Reporting of Prescription Monitoring Program Information.....		164164
Section 7. Access to Prescription Monitoring Program Information/Confidentiality.....		165165
Section 8. Interoperability.....		166166
Section 9. Unlawful Acts and Penalties.....		166166
Section 10. Evaluation, Data Analysis, and Reporting.....		167167

<i>Section 11. Rules and Regulations.</i> .....	<del>167</del> 167
<i>Section 12. Severability.</i> .....	<del>167</del> 167
<i>Section 13. Effective Date.</i> .....	<del>167</del> 167
APPENDIX F MODEL RULES FOR THE PRACTICE OF TELEPHARMACY .....	<del>168</del> 168
(a) <i>General Requirements</i> .....	<del>168</del> 168
(b) <i>Remote Dispensing Site Requirements</i> .....	<del>168</del> 168

# National Association of Boards of Pharmacy

## Model State Pharmacy Act

### Article I

#### Title, Purpose, and Definitions

##### Introductory Comment to Article I

*Article I of the Model State Pharmacy Act and Model Rules of the National Association of Boards of Pharmacy (Model Act) sets forth the foundation upon which the Act is constructed. It clearly declares and acknowledges that safeguarding the public interest is the foremost compelling reason for regulating the Practice of Pharmacy and the Distribution of Drugs and related Devices. It also circumscribes the activities included within the Practice of Pharmacy, as well as the definitions of several other terms used throughout the Act.*

NABP created the Model Act to provide State Boards of Pharmacy with model language that may be used when developing state laws or board rules for the respective States. *NABP believes that it is both desirable and necessary to recognize that the public interest must be the central precept in the Model Act and its administration, and that State Boards of Pharmacy must constantly strive to achieve the principles enunciated in Article I of the Act.*

*An ACT concerning the regulation of the Practice of Pharmacy in this State and related matters.*

*Be it enacted. . . .*

##### Section 101. Title of Act.

This Act shall be known as the “\_\_\_\_\_ Pharmacy Practice Act.”

##### Section 102. Legislative Declaration.

The Practice of Pharmacy in the State of \_\_\_\_\_ is declared a professional practice affecting the public health, safety, and welfare and is subject to regulation and control in the public interest.<sup>1</sup> It is further declared to be a matter of public interest and concern that the Practice of Pharmacy, as defined in this Act, merit and receive the confidence of the public and that only qualified persons be permitted to engage in the Practice of Pharmacy, and to ensure the quality of Drugs and related Devices Distributed in the State of \_\_\_\_\_. This Act shall be liberally construed to carry out these objectives and purposes.

##### Section 103. Statement of Purpose.

It is the purpose of this Act to promote, preserve, and protect the public health, safety, and welfare by and through the effective control and regulation of the Practice of Pharmacy; the licensure of Pharmacists; the registration of Certified Pharmacy Technicians and Certified Pharmacy Technician Candidates; the licensure, control, and regulation of all sites or Persons, in or out of this State, that Distribute, Manufacture, or sell Drugs (or Devices used in the Dispensing and Administration of Drugs), within this State, and the regulation and control of such other materials as

---

<sup>1</sup> Pharmacy is a learned profession affecting public health and welfare and should be declared as such by the State Legislature. The Practice of Pharmacy, from time to time, has been erroneously viewed, even by government agencies, as a commercial business rather than a profession. The status of Pharmacy as a profession has been, and will continue to be, of particular importance in litigation.



may be used in the diagnosis, treatment, and prevention of injury, illness, and disease of a patient or other individual.<sup>2</sup>

#### **Section 104. Practice of Pharmacy.**

The “Practice of Pharmacy” means, but is not limited to, the interpretation, evaluation, Dispensing, and/or implementation of Medical Orders, and the initiation and provision of Pharmacist Care Services. The Practice of Pharmacy also includes continually optimizing patient safety and quality of services through effective use of emerging technologies and competency-based training.<sup>3</sup>

#### **Section 105. Definitions.**

- (a) “Active Ingredients” refer to chemicals, substances, or other Components of articles intended for use in the diagnosis, cure, mitigation, treatment, or prevention of diseases in humans or animals or for use as nutritional supplements.
- (b) “Added Substances” mean the ingredients necessary to prepare the Drug Product but are not intended or expected to cause a human pharmacologic response if administered alone in the amount or concentration contained in a single dose of the Compounded Drug Product or alter the composition and effectiveness of the Compounded Drug Product. The term “added substances” is used synonymously with the terms “inactive ingredients,” “excipients,” “flavoring agents,” and “pharmaceutic ingredients.”
- (c) “Administer” means the direct application of a Drug to the body of a patient or research subject by injection, inhalation, ingestion, or any other means.
- (d) “Adulterated”: A Drug or Device shall be deemed to be Adulterated:
  - (1) if:
    - (i) it consists in whole or in part of any filthy, putrid, or decomposed substance; or
    - (ii) it has been produced, prepared, packed, or held under insanitary conditions whereby it may have been contaminated with filth, or whereby it may have been rendered injurious to health; or if the methods used in, or the facilities or controls used for, its manufacture, processing, packing, or holding do not conform to or are not operated or administered in conformity with current good manufacturing practices to ensure that such Drug or Device meets the requirements of this part as to safety and has the identity and strength, and meets the quality and purity characteristics that it purports or is represented to possess; or
    - (iii) its container is composed, in whole or in part, of any poisonous or deleterious substance that may render the contents injurious to health; or

---

<sup>2</sup> The Statement of Purpose is designed to define the general scope of the Pharmacy Act. It provides for the control and regulation of the Practice of Pharmacy and the licensure of facilities engaged in the Distribution of Drugs and related Devices. A Board will have full knowledge of the whereabouts of Drugs in the legitimate stream of intrastate and interstate commerce, providing it with the ability to better prevent diversion, effectuate recalls, ensure the quality of Drugs Dispensed or Administered to patients, and effectively protect the public.

<sup>3</sup> The definition of the “Practice of Pharmacy” is one of the most important, and perhaps one of the most discussed, clauses in the NABP *Model Act*. The definition is purposely expressed in broad terms to provide substantial latitude to the Board of Pharmacy in the adoption of implementing rules. Inclusion of or authorization for specific activities, such as the Administration of medications, is purposely not delineated in the definition in order to be empowering and not proscriptive. To assist states in the interpretation of the revised definition of the “Practice of Pharmacy,” the *Model Act* includes the definition of “Pharmacist Care Services” and Model Rules for the Provision of Pharmacist Care Services for those states that need to include specific lists of activities or authorizations due to legislative and/or administrative policies and procedures.

The definition also acknowledges that pharmacy is a dynamic profession and a broad definition of the practice will permit the Board to make necessary changes from time to time to meet the changing practice. Such changes may be affected by new or amended rules, which would be promulgated pursuant to the requirements of the State Administrative Procedures Act, affording all interested parties an opportunity to review and comment on any proposed rules.

- (iv) it bears or contains, for purposes of coloring only, a color additive that is unsafe within the meaning of the Federal Food, Drug, and Cosmetic Act (Federal Act); or it is a color additive, the intended use of which, in or on such Drugs or Devices, is for purposes of coloring only, and is unsafe within the meaning of the Federal Act;
- (2) if it purports to be or is represented as a Drug, the name of which is recognized in an official compendium, and its strength differs from, or its quality or purity falls below, the standard set forth in the compendium. Such a determination as to strength, quality, or purity shall be made in accordance with the tests or methods of assay set forth in the compendium, or in the absence of or inadequacy of these tests or methods of assay, those prescribed under authority of the Federal Act. No Drug defined in an official compendium shall be deemed to be Adulterated under this paragraph because it differs from the standard of strength, quality, or purity therefore set forth in the compendium, if its difference in strength, quality, or purity from that standard is plainly stated on its label. Whenever a Drug is recognized in both the United States Pharmacopeia (USP) and the Homeopathic Pharmacopoeia of the United States it shall be subject to the requirements of the USP unless it is labeled and offered for sale as a homeopathic Drug, in which case it shall be subject to the Homeopathic Pharmacopoeia of the United States and not those of the USP;
- (3) if it is not subject to paragraph (2) and its strength differs from, or its purity or quality falls below, that which it purports or is represented to possess; or
- (4) if it is a Drug and any substance has been mixed or packed therewith so as to reduce its quality or strength; or substituted wholly or in part therefore.
- (e) “Affiliated Entity” means legally separate covered entities that are affiliated and that designate themselves as a single covered entity for the purposes of this section.
- (f) “Automated Pharmacy Systems” include, but are not limited to, mechanical systems that perform operations or activities, Compounding or Administration, relative to the storage, packaging, Dispensing, or Distribution of medications, and which collect, control, and maintain all Transaction Information.
- (g) “Beyond-Use Date” means a date placed on a prescription label at the time of Dispensing that is intended to indicate to the patient or caregiver a time beyond which the contents of the prescription are not recommended to be used.<sup>4</sup>
- (h) “Bioburden” means the total number of microorganisms associated with a specific item prior to sterilization.
- (i) “Biological Product” is
  - (1) regulated by Food and Drug Administration (FDA);
  - (2) used to diagnose, prevent, treat, and cure diseases and medical conditions;
  - (3) a diverse category of Products and generally large, complex molecules;
  - (4) produced through biotechnology in a living system, such as a microorganism, plant cell, or animal cell, and often more difficult to characterize than small molecule Drugs.
- (j) “Biosimilar Product” is a Biological Product that is highly similar to and has no clinically meaningful differences from an existing FDA-approved Reference Product.
- (k) “Board of Pharmacy” or “Board” means the governmental regulatory body empowered to regulate pharmaceutical practices including granting and disciplining licenses of individuals and companies.
- (l) “Cease and Desist” is an order of the Board prohibiting a licensee or other Person or entity from continuing a particular course of conduct that violates the Pharmacy Practice Act or its rules and regulations.<sup>5</sup>
- (m) “Censure” is a severe formal reproof of a licensee for violation of the Pharmacy Practice Act or rules and regulations, and may require specific redress; for example, restitution of fees.

<sup>4</sup> In determining a Beyond-Use Date for a specific Drug Product, the Pharmacist may use the recommendations provided in the most recent edition of the United States Pharmacopeia-National Formulary (USP-NF).

<sup>5</sup> No proof of actual damage is required for issuance of a Cease and Desist order.

- (n) “Centralized Performance Database” means aggregate data from a large number of pharmacies concerning Quality-Related Events and patients for whom pharmaceutical Products and services have been provided at the pharmacies, and from which patient identifiers have been removed.
- (o) “Centralized Prescription Processing” means the processing by a Pharmacy of a request from another Pharmacy to fill or refill a Prescription Drug Order or to perform processing functions such as Dispensing, Drug Utilization Review (DUR), claims adjudication, refill authorizations, and therapeutic interventions.
- (p) “Certified Pharmacy Technician”<sup>6</sup> means personnel licensed by the Board who have completed a certification program approved by the Board and may, under the supervision of a Pharmacist, perform certain activities involved in the Practice of Pharmacy that are within their scope of certification and as assigned by the Pharmacist, but excluding clinical patient care activities such as, but not limited to:
  - (1) Drug Utilization Review (DUR);
  - (2) clinical conflict resolution; and
  - (3) Patient Counseling.
- (q) “Certified Pharmacy Technician Candidate” means personnel licensed by the Board who intend to complete a certification program approved by the Board and may, under the supervision of a Pharmacist, perform certain activities involved in the Practice of Pharmacy that are within their scope of education and training and as assigned by the Pharmacist, but excluding clinical patient care activities such as, but not limited to:
  - (1) Drug Utilization Review (DUR);
  - (2) clinical conflict resolution; and
  - (3) Patient Counseling.
- (r) “Chain Pharmacy Warehouse” means a permanent physical location for Drugs and/or Devices that acts as a central warehouse and performs intracompany sales and transfers of Prescription Drugs or Devices to chain Pharmacies, which are members of the same affiliated group, under common ownership and control. Chain Pharmacy Warehouses must be licensed as Wholesale Distributors.
- (s) “Chart Order” means a lawful order entered on the chart or a medical record of an inpatient or resident of an Institutional Facility by a Practitioner or his or her licensed health care designee for a Drug or Device and shall be considered a Prescription Drug Order provided that it contains:
  - (1) the full name of the patient;
  - (2) date of issuance;
  - (3) name, strength, and dosage form of the Drug prescribed;
  - (4) directions for use; and
  - (5) if written or electronic, the prescribing Practitioner’s signature<sup>7</sup> or the signature of the Practitioner’s licensed health care designee (including the name of the prescribing Practitioner).

Bidirectional transmission of Chart Orders between the Institutional Pharmacy and the Institutional Facility is allowed. The Pharmacist-in-Charge shall ensure that the Institutional Pharmacy has policies and procedures for a Practitioner to delegate the transmittal of a Chart Order to a licensed nurse employed by, or contracted by, the Institutional Facility and acting within the scope of his or her practice. Renewal of ongoing Chart Orders shall be signed by the prescriber at the appropriate time interval based on facility type and federal regulation, state law, or rule. Chart Orders shall be ongoing until such time as the Practitioner discontinues the order and such discontinuation is communicated to the Pharmacy, including but not limited to, by automatic stop order, unless otherwise indicated.

---

<sup>6</sup> The *Model Act* defines Certified Pharmacy Technician and Certified Pharmacy Technician Candidate separately to distinguish between the activities that can be performed. A Certified Pharmacy Technician is recognized, because of the completion of a Board-approved certification program, as having knowledge and skills that qualify them to assist the Pharmacist in the Practice of Pharmacy with limited patient care tasks that exceed routine Dispensing or Drug storage activities. Certified Pharmacy Technician Candidates are limited to routine Dispensing activities, Drug storage, medical coverage claims processing, and cashing.

<sup>7</sup> A Practitioner’s signature for Chart Orders is only required to be maintained at the Institutional Facility unless otherwise required for controlled substances by state and federal law.

- (t) “Closed Pharmacy” means a Pharmacy that purchases Drugs or Devices for a limited patient population and is not open for dispensing to the general patient population and cannot operate or be licensed as a Wholesale Distributor.
- (u) “Co-licensee” means a pharmaceutical manufacturer that has entered into an agreement with another pharmaceutical manufacturer to engage in a business activity or occupation related to the manufacture or distribution of a Prescription Drug.
- (v) “Collaborative Pharmacy Practice” is that Practice of Pharmacy whereby one or more Pharmacists have jointly agreed, on a voluntary basis, to work in conjunction with one or more Practitioners under protocol and in collaboration with Practitioner(s) to provide patient care services to achieve optimal medication use and desired patient outcomes.
- (w) “Collaborative Pharmacy Practice Agreement” is a written and signed agreement between one or more Pharmacists and one or more Practitioners that provides for Collaborative Pharmacy Practice as defined by law and the Rules of the Board.
- (x) “Common Carrier” means any person or entity who undertakes, whether directly or by any other arrangement, to transport property including Prescription Drugs for compensation.<sup>8</sup>
- (y) “Component” means any Active Ingredient or Added Substance intended for use in the Compounding of a Drug, including those that may not appear in such Drug.
- (z) “Compounding” means the preparation, mixing, assembling, altering, packaging, or Labeling of a Drug, Drug-Delivery Device, or Device, unless performed in a Food and Drug Administration (FDA)-registered Outsourcing Facility in conformance with Federal law, in accordance with a licensed Practitioner’s prescription, medication order, or initiative based on the Practitioner/patient/Pharmacist/compounder relationship in the course of professional practice. Compounding includes the following:
  - (1) preparation of Drug dosage forms for both human and animal patients;
  - (2) preparation of Drugs or Devices in anticipation of Prescription Drug Orders based on routine, regularly observed prescribing patterns<sup>9</sup>; and
  - (3) manipulation of commercial Products for patient-specific needs beyond FDA-approved Labeling.<sup>10</sup>
- (a2) “Consumer Survey” means a systematic record of consumer perceptions of the quality of pharmaceutical Products and services provided at a pharmacy.
- (b2) “Continuous Quality Improvement Program” means a system of standards and procedures to identify and evaluate quality-related events, and to constantly enhance the efficiency and effectiveness of the structures and processes of a pharmacy system that determine the outcomes of medication use. All information, communications, or data maintained as a component of such a system shall be privileged and confidential and not subject to discovery in civil litigation.<sup>11</sup>
- (c2) “Contraband Device” means a Device that is Counterfeit, stolen, Misbranded, obtained by fraud, purchased by a nonprofit institution for its own use and placed in commerce in violation of the own use agreement for that Device, or for which the documentation in existence has been forged, Counterfeited, falsely created, or contains any altered, false, or misrepresented information.
- (d2) “Contraband Drug” means a Drug which is Counterfeit, stolen, Misbranded, obtained by fraud, purchased by a nonprofit institution for its own use and placed in commerce in violation of the own use agreement for that Drug, that has inappropriately entered the Drug supply chain Distribution.

---

<sup>8</sup> The definition of “Common Carrier” specifically excludes Wholesale Distributors, which are defined separately.

<sup>9</sup> Anticipatorily Compounded Drugs may not be dispensed until receipt of a patient-specific Prescription Drug Order.

<sup>10</sup> Reconstitution of an FDA-approved Drug according to FDA-approved Labeling is not Compounding.

<sup>11</sup> States should continue efforts to develop and implement requirements for Continuous Quality Improvement (CQI) Programs in pharmacies, recognizing that CQI Programs enhance patient safety and operate most effectively when privilege of discovery laws and/or regulations protecting CQI data and information are enacted and included as a component of the CQI process.

- (e2) “Costs/Administrative Costs” is a monetary amount assessed a licensee to cover the cost of investigation and prosecution of a disciplinary action.
- (f2) “Counterfeit Device” means a Device which, or the container, shipping container, seal, or Product Labeling of which, without authorization, bears the trademark, trade name, or other identifying mark, imprint, or any likeness thereof, of a Manufacturer, processor, packer, or Distributor other than the Person or Persons who in fact Manufactured, processed, packed, Distributed, or Wholesale Distributed such Device and which thereby falsely purports or is represented to be the Product of, or to have been packed, Distributed, or Wholesale Distributed by, such other Manufacturer, processor, packer, or Distributor.
- (g2) “Counterfeit Drug” means a Drug which, or the container, shipping container, seal, or Product Labeling of which, without authorization, bears the trademark, trade name, or other identifying mark, imprint, or Device, or any likeness thereof, of a Manufacturer, processor, packer, or Distributor other than the Person or Persons who in fact Manufactured, processed, packed, Distributed, or Wholesale Distributed such Drug and which thereby falsely purports or is represented to be the Product of, or to have been packed, Distributed, or Wholesale Distributed by, such other Manufacturer, processor, packer, or Distributor.
- (h2) “Critical Areas” means areas designed to maintain sterility of sterile materials. Sterilized Product, container/closures, and equipment may be exposed in critical areas.
- (i2) “Critical Surfaces” are surfaces that may come into contact with or directly impact sterilized Product or containers/closures.
- (j2) “Cytotoxic” means a pharmaceutical that has the capability of killing living cells.
- (k2) “Declared Disaster Areas” are areas designated by state or federal authorities as those that have been adversely affected by a natural or man-made disaster and require extraordinary measures to provide adequate, safe, and effective health care for the affected population.
- (l2) “De-identified Health Information” means Health Information that does not identify an individual and with respect to which there is no reasonable basis to believe that the information can be used to identify an individual. De-identified health information must meet the specifications of the de-identified health information described in the HIPAA privacy rules (45 CFR §164.514(b)).  
(See comment list.)
- (m2) “Deliver” or “Delivery” means the actual, constructive, or attempted transfer of a Drug or Device from one Person to another, whether or not for a consideration.
- (n2) “Designated Record Set” means:
  - (1) A group of records maintained by or for a Pharmacy, Pharmacist, Pharmacy Benefits Manager, or Person licensed or registered by the Board that is:
    - (i) the medical records and billing records about patients maintained by or for a Pharmacy, Pharmacist, Pharmacy Benefits Manager, or Person licensed or registered by the Board;
    - (ii) the enrollment, payment, claims adjudication, and case or medical management record systems maintained by or for a health plan; or
    - (iii) used, in whole or in part, by or for the Pharmacy, Pharmacist, Pharmacy Benefits Manager, or Person licensed or registered by the Board to make decisions about patients.
  - (2) For purposes of this paragraph, the term “record” means any item, collection, or grouping of information that includes Protected Health Information and is maintained, collected, used, or disseminated by or for a Pharmacy, Pharmacist, Pharmacy Benefits Manager, or Person licensed or registered by the Board.

- (o2) “Designated Representative” means an individual designated by the Wholesale Distributor who will serve as the responsible individual of the Wholesale Distributor with the Board who is actively involved in and aware of the actual daily operation of the Wholesale Distributor.<sup>12</sup>
- (p2) “Device” means an instrument, apparatus, implement, machine, contrivance, implant, or other similar or related article, including any component part or accessory, which is required under Federal law to bear the label, “Caution: Federal or State law requires Dispensing by or on the order of a physician.”<sup>13</sup>
- (q2) “Digital Signature” means an electronic signature based upon cryptographic methods of originator authentication, and computed by using a set of rules and a set of parameters so that the identity of the signer and the integrity of the data can be verified.
- (r2) “Disinfection” means the process by which surface Bioburden is reduced to a safe level or eliminated.
- (s2) “Dispense” or “Dispensing” means the interpretation, evaluation, and implementation of a Prescription Drug Order, including the preparation, final Verification, and Delivery of a Drug or Device to a patient or patient’s agent in a suitable container appropriately labeled for subsequent Administration to, or use by, a patient.<sup>14</sup>
- (t2) “Dispenser” means a retail Pharmacy, hospital Pharmacy, a group of chain Pharmacies under common ownership and control that do not act as a Wholesale Distributor, or any other Person authorized by law to Dispense or Administer Prescription Drugs, and the affiliated warehouses or Distribution centers of such entities under common ownership and control that do not act as a Wholesale Distributor.
- (u2) “Distribute” or “Distribution” means to sell, offer to sell, deliver, offer to deliver, broker, give away, or transfer a Drug, whether by passage of title, physical movement, or both. The term does not include:
- (1) To Dispense or Administer;
  - (2) Delivering or offering to deliver a Drug by a common carrier in the usual course of business as a common carrier; or
  - (3) Providing a Drug sample to a patient by a Practitioner licensed to prescribe such Drug; a health care professional acting at the direction and under the supervision of a Practitioner; or the Pharmacy of a hospital or of another health care entity that is acting at the direction of such a Practitioner and that received such sample in accordance with the Act and regulations to administer or dispense.
- (v2) “Diversion Activity” means activity where evidence exists that controlled substances or Drugs of Concern are being diverted from legitimate channels.
- (w2) “Drug” means:
- (1) articles recognized as Drugs in any official compendium, or supplement thereto, designated from time to time by the Board for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in humans or other animals;<sup>15</sup>
  - (2) articles intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in humans or other animals;
  - (3) articles (other than food) intended to affect the structure or any function of the body of humans or other animals; and
  - (4) articles intended for use as a Component of any articles specified in clause (1), (2), or (3) of this definition.

---

<sup>12</sup> The “Designated Representative” should serve as a liaison to the Board who is extremely knowledgeable about and involved in the daily operations of the Wholesale Distributor. If a Wholesale Distributor is licensed by multiple states, it is not necessary for the Wholesale Distributor to have multiple Designated Representatives. One Designated Representative per Wholesale Distributor facility is sufficient.

<sup>13</sup> States at their option may want to consider limiting the definition of “Devices” to those Devices associated with the Dispensing, Administration, or use of Drugs.

<sup>14</sup> “Dispensing” includes the Delivery of a Drug or Device to the patient or the patient’s agent by the Pharmacist or the Pharmacist’s agent. Drugs and/or Devices mailed or shipped to a patient are not Dispensed until the Drugs and/or Devices are actually received by the patient or the patient’s agent.

<sup>15</sup> The official compendium recognized by Food and Drug Administration (FDA) and many State Boards of Pharmacy is the USP-NF.

- (x2) “Drug of Concern” means any prescription or over-the-counter medication that demonstrates a potential for abuse, particularly those identified by Boards of Pharmacy, law enforcement, and addiction treatment professionals.
- (y2) “Drug Utilization Review (DUR)”<sup>16</sup> includes but is not limited to the following activities:
- (1) Evaluation of the Prescription Drug Order(s) and patient record(s) for:
    - (i) known allergies;
    - (ii) rational therapy contraindications;
    - (iii) reasonable dose, duration of use, and route of Administration, considering age, gender, and other patient factors;<sup>17</sup>
    - (iv) reasonable directions for use;
    - (v) potential or actual adverse Drug reactions;
    - (vi) Drug-Drug interactions;
    - (vii) Drug-food interactions;
    - (viii) Drug-disease contraindications;
    - (ix) therapeutic duplication;
    - (x) proper utilization (including over- or under-utilization), and optimum therapeutic outcomes; and
    - (xi) abuse/misuse.
- (z2) “Electronic Signature” means an electronic sound, symbol, or process attached to or logically associated with a record and executed or adopted by a Person with the intent to sign the record.<sup>18</sup>
- (a3) “Emergency Medical Reasons” include, but are not limited to, transfers of a prescription Drug between a Wholesale Distributor or Pharmacy to alleviate a temporary shortage of a prescription Drug arising from delays in or interruption of regular distribution schedules; sales to nearby emergency medical services, ie, ambulance companies and firefighting organizations in the same state or same marketing or service area, or nearby licensed Practitioners of Prescription Drugs for use in the treatment of acutely ill or injured Persons; provision of minimal emergency supplies of Prescription Drugs to nearby nursing homes for use in emergencies or during hours of the day when necessary Prescription Drugs cannot be obtained; and transfers of Prescription Drugs by a retail Pharmacy to another retail Pharmacy to alleviate a temporary shortage.
- (b3) “Emergency Prescription Drug Order” means a standing Prescription Drug Order issued by the State Health Officer for Pharmacists to Dispense designated Prescription Drugs during a Public Health Emergency requiring mass Dispensing to expeditiously treat or provide prophylaxis to large numbers of Patients.
- (c3) “Emergency Situations,” for the purposes of authorizing an oral Prescription Drug Order of a Schedule II controlled substance, means those situations in which the prescribing Practitioner determines (1) that immediate Administration of the controlled substance is necessary for proper treatment of the patient, (2) that no appropriate alternative treatment is available, including Administration of a Drug that is not a Schedule II controlled substance, and (3) that it is not reasonably possible for the prescribing Practitioner to provide a written Prescription Drug Order to be presented to the Person Dispensing the substance, prior to the Dispensing.
- (d3) “Enteral” means within or by way of the gastrointestinal tract or intestine.

---

<sup>16</sup> DUR is also known to mean “Drug Use Review”; however, “Drug Utilization Review” is the preferred term.

<sup>17</sup> A “reasonable” dose, duration of use, and route of administration under “Drug Utilization Review” would be determined by taking into consideration patient-specific factors, including but not limited to, age, gender, and other patient factors, but dependent upon the information about the patient known to the pharmacist.

<sup>18</sup> The term “Electronic Signature” may have different meanings in various State laws and regulations. It is important to distinguish between “Electronic Signatures” and “Digital Signatures,” which provide a much higher level of security for electronically transmitted information.

- (e3) “Equivalent Drug Product” means a Drug Product that has the same established name, active ingredient(s), strength or concentration, dosage form, and route of Administration and which is formulated to contain the same amount of active ingredient(s) in the same dosage form and to meet the same compendial or other applicable standards (eg, strength, quality, purity, and identity), but which may differ in characteristics, such as shape, scoring, configuration, packaging, excipients (including colors, flavors, and preservatives), and expiration time.
- (f3) “Exclusive Distributor” means the Wholesale Distributor who directly purchased the Product from the Manufacturer and is the sole Distributor of that Manufacturer’s Product to a subsequent Repackager, Wholesale Distributor, or Dispenser.
- (g3) “External Entities” means those organizations that exist outside of the pharmacist-patient relationship and that participate in the implementation of Patient Compliance and Patient Intervention Programs. External Entities include, but are not limited to, health insurance carriers, health benefit management companies, and health care marketing enterprises. [Note: Depending on the activities conducted by External Entities, they may be construed as “business associates” as defined under HIPAA and its related privacy rules (45 CFR Part 160). If so, HIPAA and its privacy rules that apply to those External Entities acting as business associates shall take precedence over contrary state law. In addition, “business associate agreements,” as defined under HIPAA and its privacy rules, shall be required between a Pharmacist or Pharmacy and the External Entity acting as a business associate so as to prevent the unauthorized use or disclosure of Protected Health Information.]
- (h3) “FDA” means Food and Drug Administration, a federal agency within the United States Department of Health and Human Services, established to set safety and quality standards for Drugs, food, cosmetics, and other consumer Products.
- (i3) “Federal Act” means the Federal Food, Drug, and Cosmetic Act.
- (j3) “Fill date” means the actual date a new or refilled prescription is dispensed but not necessarily delivered to a patient from a pharmacy.
- (k3) “Fine/Civil Penalty” is a monetary penalty assessed a licensee for violation of the Pharmacy Practice Act or rules and regulations.
- (l3) “Global Exchange Pharmacy Student” means a current student of a non-US professional degree program of a school or college of Pharmacy who is participating in an exchange program administered by an ACPE-accredited or Board-approved US school or college of pharmacy for a limited duration in this state.
- (m3) “Health Care Entity” means any Person that provides diagnostic, medical, surgical, dental treatment, or rehabilitative care but does not include any retail Pharmacy or Wholesale Distributor.
- (n3) “Health Care Operations” means any of the following activities of the Pharmacy, Pharmacist, Pharmacy Benefits Manager, or Person licensed or registered by the Board to the extent that the activities are related to the provision of Pharmacist Care Services:
  - (1) conducting quality assessment and improvement activities, including outcomes evaluation and development of clinical guidelines, provided that the obtaining of generalizable knowledge is not the primary purpose of any studies resulting from such activities; population-based activities relating to improving health or reducing health care costs, protocol development, case management and care coordination, contacting of health care providers and patients with information about treatment alternatives; and related functions that do not include treatment;
  - (2) reviewing the competence or qualifications of health care professionals, evaluating practitioner and provider performance, health plan performance, conducting training programs in which students, trainees, or practitioners in areas of health care learn under supervision to practice or improve their skills as health care providers, training of non-health care professionals, accreditation, certification, licensing, or credentialing activities;
  - (3) underwriting, premium rating, and other activities relating to the creation, renewal, or replacement of a contract of health insurance or health benefits, and ceding, securing, or placing a contract for



reinsurance of risk relating to claims for health care (including stop-loss insurance and excess of loss insurance),<sup>19</sup> provided that the requirements of 45 CFR §164.514(g) are met, if applicable;

- (4) conducting or arranging for medical review, legal services, and auditing functions, including fraud and abuse detection and compliance programs;
- (5) business planning and development, such as conducting cost-management and planning-related analyses related to managing and operating the entity, including formulary development and Administration, development, or improvement of methods of payment or coverage policies; and
- (6) business management and general administrative activities, including, but not limited to:
  - (i) management activities relating to implementation of and compliance with the requirements of this Act;
  - (ii) customer service, including the provision of data analyses for policy holders, plan sponsors, or other customers, provided that Protected Health Information is not disclosed to such policy holder, plan sponsor, or customer;
  - (iii) resolution of internal grievances;
  - (iv) the sale, transfer, merger, or consolidation of all or part of the Pharmacy, Pharmacy Benefits Manager, or other entity that is or will be licensed or registered by the Board with another Pharmacy, Pharmacy Benefits Manager, or other entity licensed or registered by the Board and due diligence related to such activity; and
  - (v) creating de-identified health information or a limited data set, and fundraising for the benefit of the Pharmacy, Pharmacy Benefits Manager, or other entity licensed or registered by the Board.<sup>20</sup>

(o3) “Health Information” means any information, whether oral or recorded in any form or medium, that:

- (1) is created or received by a health care provider, health plan, public health authority, employer, life insurer, school or university, or health care clearinghouse, and
- (2) relates to the past, present, or future physical or mental health or condition of an individual; or the past, present, or future payment for the provision of health care to an individual.

(p3) “HIPAA” is the federal Health Insurance Portability and Accountability Act of 1996 (Public Law 104-191) and any amendments thereof.

(q3) “Home Infusion Pharmacy” means a Pharmacy that Compounds solutions for direct Administration to a patient in a private residence, Long-Term Care Facility, or hospice setting by means of parenteral, intravenous, intramuscular, subcutaneous, or intraspinal infusion.

(r3) “Illegitimate Product” means a Product for which credible evidence shows that the Product:

- (1) is Counterfeit, diverted, or stolen;
- (2) is intentionally Adulterated such that the Product would result in serious adverse health consequences or death to humans;
- (3) is the subject of a fraudulent Transaction; or
- (4) appears otherwise unfit for Distribution such that the Product would be reasonably likely to result in serious adverse health consequences or death to humans.

(s3) “Immediate Container” means a container and does not include package liners.

(t3) “Individually Identifiable Health Information” is information that is a subset of Health Information, including demographic information collected from an individual and

- (1) is created or received by a health care provider, health plan, employer, or health care clearinghouse; and

---

<sup>19</sup> 45 CFR §164.514(g) reads:

Standard: uses and disclosures for underwriting and related purposes. If a health plan receives protected health information for the purpose of underwriting, premium rating, or other activities relating to the creation, renewal, or replacement of a contract of health insurance or health benefits, and if such health insurance or health benefits are not placed with the health plan, such health plan may not use or disclose such protected health information for any other purpose, except as may be required by law.

<sup>20</sup> The word “fundraising” is contemplated to refer to generation of revenue through the sale of data, and is not intended to be used in the charitable sense.

- (2) relates to the past, present, or future physical or mental health or condition of an individual; the provision of health care to an individual; or the past, present, or future payment for the provision of health care to an individual; and
  - (i) that identifies the individual; or
  - (ii) with respect to which there is a reasonable basis to believe the information can be used to identify the individual.
- (u3) “Institutional Facility” means any organization whose primary purpose is to provide a physical environment for patients to obtain health care services, including but not limited to a(n):
  - (1) hospital;
  - (2) Long-Term Care Facility;
  - (3) convalescent home;
  - (4) nursing home;
  - (5) extended care facility;
  - (6) mental health facility;
  - (7) rehabilitation center;
  - (8) psychiatric center;
  - (9) developmental disability center;
  - (10) Drug abuse treatment center;
  - (11) family planning clinic;
  - (12) penal institution;
  - (13) hospice;
  - (14) public health facility;
  - (15) athletic facility;
  - (16) assisted living facility; and
  - (17) intermediate care facility for individuals with intellectual disabilities.
- (v3) “Institutional Pharmacy”<sup>21</sup> means any place that is registered with the State Board of Pharmacy pursuant to Article V of the Pharmacy Practice Act that provides Pharmacist Care Services to an Institutional Facility and where Drugs, Devices, and other materials used in the diagnosis and treatment of injury, illness, and disease (hereinafter referred to as “Drugs”) are Dispensed, Compounded, and Distributed.<sup>22</sup>
- (w3) “Interchangeable Product” is a Biosimilar Product that meets additional requirements.
- (x3) “Internal Test Assessment” means, but is not limited to, conducting those tests of quality assurance necessary to ensure the integrity of the test.
- (y3) “Intracompany Transaction” means any transaction between a division, subsidiary, parent, and/or affiliated or related company under the common ownership and control of a corporate entity.
- (z3) “ISO Class” means the description of an atmospheric environment characterized by the number of particles within a diameter per cubic foot of air.
- (a4) “Isolator” means a decontaminated unit, supplied with ISO Class 5 or higher air quality that provides uncompromised, continuous isolation of its interior from the external environment (eg, surrounding cleanroom air and Compounding Pharmacy personnel).
- (b4) “Label” means a display of written, printed, or graphic matter upon the immediate container of any Drug or Device.
- (c4) “Labeling” means the process of preparing and affixing a label to any Drug container exclusive, however, of the Labeling by a Manufacturer, packer, or Distributor of a Nonprescription Drug or commercially

---

<sup>21</sup> Although traditionally characterized as being physically part of an Institutional Facility, the Model Rules recognize that an Institutional Pharmacy may or may not be physically attached to an Institutional Facility.

<sup>22</sup> States should recognize that hospitals, in order to prepare for a disaster or emergency, may be stocking emergency supplies of medications in areas outside the licensed Pharmacy. Hospitals should be encouraged to expand the space allotted to the licensed Pharmacy area to accommodate the need to store emergency supplies.

packaged Legend Drug or Device. Any such label shall include all information required by Federal and State law or rule.

- (d4) (a4) “Long-Term Care Facility” means a nursing home, retirement care, mental care, or other facility or institution that provides extended health care to resident patients.
- (e4) (b4) “Manufacturer” means a Person, which may include a Virtual Manufacturer, engaged in the Manufacture of Drugs or Devices.
- (f4) (c4) “Manufacturing” means the production, preparation, propagation, conversion, or processing of a Drug or Device, either directly or indirectly, by extraction from substances of natural origin or independently by means of chemical or biological synthesis. Manufacturing includes the packaging or repackaging of a Drug or Device or the Labeling or relabeling of the container of a Drug or Device for resale by pharmacies, Practitioners, or other Persons.<sup>23</sup>
- (g4) “Marketing” means:
  - (1) To make a communication about a Product or service that encourages recipients of the communication to purchase or use the Product or service, unless the communication is made:
    - (i) to describe a health-related Product or service (or payment for such Product or service) that is provided by, or included in a plan of benefits of, the covered entity making the communication, including communications about the entities participating in a health care provider network or health plan network; replacement of, or enhancements to, a health plan; and health-related Products or services available only to a health plan enrollee that add value to, but are not part of, a plan of benefits;
    - (ii) for treatment of the patient; or
    - (iii) for case management or care coordination for the patient, or to direct or recommend alternative treatments, therapies, health care providers, or settings of care to the patient.
  - (2) An arrangement between a Pharmacy, Pharmacist, Pharmacy Benefits Manager, or Person licensed or registered by the Board, and any other entity whereby the Pharmacy, Pharmacist, Pharmacy Benefits Manager, or Person licensed or registered by the Board discloses Protected Health Information to the other entity in exchange for direct or indirect remuneration, for the other entity or its affiliate to make a communication about its own Product or service that encourages recipients of the communication to purchase or use that Product or service.
- (h4) “Medical Order” means a lawful order of a Practitioner that may or may not include a Prescription Drug Order.
- (i4) “Medication Adherence Monitoring Service” is defined as any structured activity that complements or supplements the existing responsibilities regarding the Dispensing of prescriptions and associated Patient Counseling, and that uses Protected Health Information to contact the patient or caregiver via phone, print, electronic media, or other means of communication, in order to improve patient compliance with and adherence to prescribed medication therapy and that involves the collection and analysis of data related to patient medication use.<sup>24</sup> Medication Adherence Monitoring Services may incorporate such efforts as refill reminder and patient education programs.
- (j4) “Medication-assisted Treatment (MAT)” is the use of medications, in combination with counseling and behavioral therapies, to provide a “whole-patient” approach to the treatment of substance use disorders. Medications used in MAT are approved by Food and Drug Administration (FDA).<sup>25</sup>
- (k4) “Medication Synchronization” refers to a component of Medication Therapy Management that recognizes the authority of the pharmacist, at the patient’s direction, to proactively adjust the

---

<sup>23</sup> Manufacturing also includes the Compounding of Drugs for office use of which can only be done by an FDA-registered Outsourcing Facility.

<sup>24</sup> Compliance refers to taking actions necessary to ensure patients receive prescribed medications initially, whereas adherence refers to taking actions necessary to ensure that medication therapy is continued.

<sup>25</sup> The Substance Abuse and Mental Health Services Administration also refers to MAT as “Medications for Opioid Use Disorder” (MOUD), which are FDA-approved medications for the treatment of opioid use disorders and currently include methadone, naltrexone, and buprenorphine.

medication quantity or refill schedule and to manage a patient's maintenance medications by coordinating the refill schedules to improve patient outcomes.<sup>26</sup>

- (l4) "Medication Therapy Management" is a distinct Pharmacist Care Service or group of services that optimize therapeutic outcomes for individual patients. Medication Therapy Management services are independent of, but can occur in conjunction with, the provision of a medication or a medical device. Medication Therapy Management encompasses a broad range of professional activities and responsibilities within the licensed Pharmacist's scope of practice. These services may include, but are not limited to, the following, according to the individual needs of the patient:
- (1) performing or obtaining necessary assessments of the patient's health status;
  - (2) formulating a medication treatment plan;
  - (3) selecting, initiating, modifying, or administering medication therapy;
  - (4) monitoring and evaluating the patient's response to therapy, including safety and effectiveness;
  - (5) performing a comprehensive medication review to identify, resolve, and prevent medication-related problems, including adverse drug events;
  - (6) documenting the care delivered and communicating essential information to the patient's other primary care providers;
  - (7) providing verbal education and training designed to enhance patient understanding and appropriate use of his or her medications;
  - (8) providing information, support services and resources designed to enhance patient adherence with his or her therapeutic regimens, such as Medication Synchronization;
  - (9) coordinating and integrating Medication Therapy Management services within the broader health care management services being provided to the patient; and
  - (10) such other patient care services as may be allowed by law.
- (m4) "Misbranded": A Drug or Device shall be deemed to be misbranded if the label is false or misleading in any particular; or the label does not bear the name and address of the Manufacturer, packer, or Distributor and does not have an accurate statement of the quantities of the active ingredients in the case of a Drug; or the label does not show an accurate monograph for Prescription Drugs.
- (n4) "Mobile Pharmacy" means a Pharmacy that is self-propelled or movable by another vehicle that is self-propelled.
- (o4) "National Association of Boards of Pharmacy (NABP)" means the association whose members are the Boards of Pharmacy, which association was established to assist Boards in developing, implementing, and enforcing uniform standards for the purpose of protecting the public health.
- (p4) "NABP e-Profile ID" means the unique identifier for permittees, licensees, and registrants that is provided by NABP. This unique, unduplicated identifier allows for, but is not limited to, the accurate identification and collection of licensure, disciplinary, and inspection information for permittees,

---

<sup>26</sup> Medication Synchronization can be effective in improving medication adherence and eliminating gaps in therapy by reducing the number of Pharmacy visits for patients on multiple-medication regimens. It is recommended that patients receive their synchronized refills by regular appointment with their Pharmacist, which allows for increased patient-Pharmacist interaction and the provision of comprehensive Medication Therapy Management services for chronic illnesses. In addition to facilitating medication adherence and improving patient outcomes, Medication Synchronization may also offer Pharmacies a mechanism to improve workload and inventory control. Other demonstrated advantages of medication synchronization include minimization of overall health costs and increased convenience for patients.

Medication Synchronization extends the pharmacist's authority to adjust medication use and quantities, not to exceed the total quantity prescribed or what is otherwise allowed by law.

"Medication Refill Consolidation," "Medication Schedule Synchronization," and "Medication Refill Synchronization" are other terms used for these types of services.

Medication Synchronization is used in the Dispensing of maintenance medications (excluding controlled substances (Schedules II-V) or those designated "as needed") for patients with chronic illnesses. Chronic illnesses are those diseases or conditions that are of long duration, require ongoing treatment, and can be controlled but not completely cured. The US National Center for Health Statistics defines a chronic disease as a condition lasting for three or more months. According to the Centers for Medicare and Medicaid Services, the most common chronic conditions among Medicare beneficiaries are hypertension, high cholesterol, heart disease, diabetes, and arthritis. Other common chronic illnesses include heart failure, depression, chronic kidney disease, osteoporosis, Alzheimer's disease, chronic obstructive pulmonary disease, atrial fibrillation, cancer, asthma, and stroke.

licensees, and registrants, both in-state as well as out-of-state, in a secure electronic profile that can be utilized for applicant submission, review, and/or Board action.

- (q4) “NABP Information Sharing Network”<sup>27</sup> means the information sharing network developed by NABP that collects, assesses, and allows for review and sharing of Compounding Pharmacy and physician information as described in the “MEMORANDUM OF UNDERSTANDING ADDRESSING CERTAIN DISTRIBUTIONS OF COMPOUNDED HUMAN DRUG PRODUCTS BETWEEN THE [insert STATE BOARD OF PHARMACY OR OTHER APPROPRIATE STATE AGENCY] AND THE U.S. FOOD AND DRUG ADMINISTRATION.”
- (r4) “Nonprescription Drug” means a Drug that may be sold without a prescription and that is labeled for use by the consumer in accordance with the requirements of the laws and rules of this State and the Federal government.
- (s4) “Nonresident Pharmacy” means a Pharmacy located outside this State.
- (t4) “Normal Distribution Channel” means a chain of custody for a Prescription Drug that goes from a Manufacturer of the Prescription Drug, the Manufacturer’s Co-Licensee, the Manufacturer’s Third-Party Logistics Provider, or the Manufacturer’s Exclusive Distributor to:
- (1) a Wholesale Distributor to a Pharmacy to a patient or other designated persons authorized by law to dispense or administer such Prescription Drug to a patient; or
  - (2) a Wholesale Distributor to a Chain Pharmacy Warehouse to that Chain Pharmacy Warehouse’s intracompany Pharmacy to a patient or other designated persons authorized by law to dispense or administer such Prescription Drug to a patient; or
  - (3) a Chain Pharmacy Warehouse to that Chain Pharmacy Warehouse’s intracompany Pharmacy to a patient or other designated persons authorized by law to dispense or administer such Prescription Drug to a patient; or
  - (4) as prescribed by the Board’s regulations.
- (u4) “Nuclear Pharmacy” means a Pharmacy providing radiopharmaceutical services or, as provided in the Model Rules for Nuclear/Radiologic Pharmacy, appropriate area of any Institutional Facility.
- (v4) “Outsourcing Facility”<sup>28</sup> means a facility at one geographic location or address that<sup>29</sup>:
- (1) is engaged in the Compounding of sterile drugs for human use;
  - (2) is registered as an Outsourcing Facility with FDA; and
  - (3) complies with all of the requirements of Section 503B of the Federal FD&C Act.
- (w4) “Parenteral” means by some other route than through the gastrointestinal tract such as, but not limited to, intravenous, subcutaneous, or intramuscular routes.
- (x4) “Patient Counseling” means the oral communication by the Pharmacist of information, as defined in the rules of the applicable Board, to the patient or caregiver, in order to ensure proper use of Drugs and Devices.
- (y4) “Patient Intervention Program” is defined as any structured activity that complements or supplements the existing responsibilities regarding the Dispensing of prescriptions and associated Patient Counseling, and that uses Protected Health Information to contact the patient or caregiver via phone, print, electronic media, or other means of communication, to discuss, inform, and/or affect patient therapy or choice of medications.
- (z4) “Peer Review” means a process that is part of an outcome-based, continuous quality improvement process that involves:

---

<sup>27</sup> The information sharing network was built by NABP pursuant to the NABP-FDA “Cooperative Agreement to Develop a System for the Collection, Management, and Sharing of Information on Compounding Pharmacies Distributing Interstate.”

<sup>28</sup> Outsourcing Facilities may engage in Compounding for animal use.

<sup>29</sup> Boards may choose to license an Outsourcing Facility as a Pharmacy; however, if a Pharmacy and an Outsourcing Facility are located at the same geographic location or address, or are located adjacent to said location or address, there must be a clear delineation between the two entities and both must comply with current Good Manufacturing Practices as defined by the Federal FD&C Act.

- (1) the setting and periodic re-evaluation of standards for quality by which a pharmacy operation will be evaluated;
  - (2) the collection of data necessary to identify when those standards are not being met and data necessary to evaluate the reason(s) the deficiency occurred;
  - (3) an objective review of the data by an appropriate peer review committee to make recommendations for quality improvement; and
  - (4) an appropriate feedback mechanism to ensure that the process is operating in a manner that continually improves the quality of care provided to patients.
    - (i) Peer review should not be a punitive activity or a performance evaluation.
- (a5) “Peer Review Committee”<sup>30</sup> means:
- (1) a committee that is authorized to evaluate the quality of pharmacy services or the competence of pharmacists and suggest improvements in pharmacy systems to enhance patient care; or
  - (2) a committee established by a person who owns a pharmacy or employs pharmacists that is authorized to evaluate the quality of pharmacy services or the competence of pharmacists and suggest improvements in pharmacy systems to enhance patient care.
- (b5) “Person” means an individual, corporation, subsidiary, partnership, association, organization, affiliate organization, or any other legal entity, including government.
- (c5) “Pharmacist” means an individual currently licensed by this State to engage in the Practice of Pharmacy. A Pharmacist is entitled to engage in the Practice of Pharmacy, as defined in this chapter, within or outside of a licensed Pharmacy, as defined in the Rules of the Board.
- (d5) “Pharmacist Care Services” is the provision by a Pharmacist of patient care activities within this State or into this State, as defined by the Rules of the Board, with or without the Dispensing of Drugs or Devices, intended to achieve outcomes related to the cure or prevention of a disease, elimination or reduction of a patient’s symptoms, or arresting or slowing of a disease process.<sup>31</sup>
- (e5) “Pharmacist-in-Charge” means a Pharmacist currently licensed in this State who accepts responsibility for the operation of a Pharmacy in conformance with all laws and rules pertinent to the Practice of Pharmacy and the Distribution of Drugs, and who is personally in full and actual charge of such Pharmacy and personnel.
- (f5) “Pharmacist’s Scope of Practice Pursuant to the Collaborative Pharmacy Practice Agreement” means those duties and limitations of duties placed upon one or more Pharmacists by the collaborating Practitioner or Practitioners, the Board, and applicable law, and includes the limitations implied by the scope of practice of the collaborating Practitioner or Practitioners.
- (g5) “Pharmacy” means any place within this State where Drugs are Dispensed and Pharmacist Care Services is provided and any place outside of this State where Drugs are Dispensed and Pharmacist Care Services is provided to residents of this State.

---

<sup>30</sup> A Peer Review Committee may be established to evaluate the quality of Pharmacy services or the competence of pharmacists and suggest improvements in Pharmacy systems to enhance patient care. Peer Review Committees may review documentation of quality-related activities in a pharmacy, assess system failures and personnel deficiencies, determine facts, and make recommendations or issue decisions in a written report that can be used for Continuous Quality Improvement purposes. A Peer Review Committee may include the members, employees, and agents of the committee, including assistants, investigators, attorneys, and any other agents that serve the committee in any capacity.

<sup>31</sup> Objectives of Pharmacist Care Services include cure of a disease, elimination or reduction of a patient’s symptomatology, arresting or slowing of a disease process, or prevention of a disease or symptomatology. Pharmacist Care Services should be provided by all Pharmacists within the Standard of Care to the extent of their abilities regardless of the practice setting.

- (h5) “Pharmacy Benefits Manager” means a Person that Administers the Prescription Drug/Device portion of health insurance plans on behalf of plan sponsors, such as self-insured employers, insurance companies, and health maintenance organizations, and that engages in or directs the Practice of Pharmacy.<sup>32</sup>
- (i5) “Pharmacy Intern”<sup>33</sup> means an individual who is:
- (1) currently licensed by this State to engage in the Practice of Pharmacy while under the supervision of a Pharmacist and is enrolled in a professional degree program of a school or college of pharmacy that has been approved by the Board and is satisfactorily progressing toward meeting the requirements for licensure as a Pharmacist; or
  - (2) a graduate of an approved professional degree program of a school or college of Pharmacy or a graduate who has established educational equivalency by obtaining a Foreign Pharmacy Graduate Examination Committee™ (FPGEC®) Certificate, who is currently licensed by the Board of Pharmacy for the purpose of obtaining practical experience as a requirement for licensure as a Pharmacist; or
  - (3) a qualified applicant awaiting examination for licensure or meeting Board requirements for re-licensure; or
  - (4) an individual participating in a residency or fellowship program.
- (j5) “Positive Patient Outcomes” include the cure or prevention of disease, elimination or reduction of a patient’s symptoms, or arresting or slowing of a disease process so as to improve the patient’s quality of life.
- (k5) “Practice Accountability Audit” means an evaluation of the Centralized Performance Database to determine which pharmacies are consistently in violation of Criteria and/or standards.
- (l5) “Practice of Telepharmacy” means the Practice of Pharmacy by registered Pharmacies and Pharmacists located within US jurisdictions through the use of Telepharmacy Technologies between a licensee and patients or their agents at distances that are located within US jurisdictions. The Practice of Telepharmacy is deemed to occur within the jurisdiction in which the patient is located and the jurisdiction(s) in which the pharmacist and, if applicable, pharmacy are located; therefore, such practice will be subject to the Pharmacy practice regulations of all jurisdictions’ Boards of Pharmacy.

---

<sup>32</sup> It is the performance of activities that encompass the Practice of Pharmacy that distinguishes Pharmacy Benefits Managers from Pharmacy Benefits Processors. The activities that may encompass the Practice of Pharmacy by Pharmacy Benefits Managers include, but are not limited to, the following:

- Disease state management
- Disease compliance management
- Drug adherence management
- Drug interaction management
- Drug utilization management
- Formulary management
- Generic alternative program management
- Generic incentive program management
- Medical and/or Drug data analysis
- Patient Drug Utilization Review (DUR) services
- Prior authorization services
- Provider profiling and outcomes assessment
- Refill reminder program management
- Therapy guidelines management
- Stop therapy protocol management
- Wellness management
- Maintenance of confidential patient information
- Direction or design of the clinical programs for a Pharmacy or a group of Pharmacies

<sup>33</sup> Most Pharmacy Interns are either enrolled in a professional degree program or postgraduate program (residency or fellowship), or have graduated from a Board-approved professional degree program and are awaiting examination. In some cases, however, Boards of Pharmacy also designate pharmacists whose licenses have lapsed or been inactive for a significant period of time as “Pharmacy Intern,” allowing these pharmacists to obtain practical experience so that their licenses can be reactivated. Additionally, Boards may grant the “Pharmacy Intern” designation to those Pharmacists seeking practical experience following a period of license suspension or revocation.

Boards of Pharmacy may consider limiting the Pharmacy Interns’ duration of registration especially if the boards find that Pharmacy Interns are not successfully progressing toward Pharmacist Licensure in an acceptable and reasonable time frame.

- (m5) “Practitioner” means an individual currently licensed, registered, or otherwise authorized by the appropriate jurisdiction to prescribe and Administer Drugs in the course of professional practice.<sup>34</sup>
- (n5) “Preceptor” means an individual who is currently licensed as a Pharmacist by the Board of Pharmacy, meets the qualifications as a Preceptor under the Rules of the Board, and participates in the instructional training of Pharmacy Interns.<sup>35</sup>
- (o5) “Prepackaging” means the act of transferring a Drug, manually or by use of an Automated Pharmacy System, from a Manufacturer’s or Distributor’s original container to another container in advance of receiving a Prescription Drug Order or for a Patient’s immediate need for Dispensing by a Pharmacy or Practitioner authorized to Dispense in the establishment in which the Prepackaging occurred.
- (p5) “Prescription Drug” or “Legend Drug” means a Drug that is required under Federal law to be labeled with either of the following statements prior to being Dispensed or Delivered: (1) “Rx Only”; or (2) “Caution: Federal law restricts this Drug to use by, or on the order of, a licensed veterinarian”; or (3) a Drug that is required by any applicable Federal or State law or rule to be Dispensed pursuant only to a Prescription Drug Order or is restricted to use by Practitioners only.
- (q5) “Prescription Drug Order” means a lawful order from a Practitioner for a Drug or Device for a specific patient, including orders derived from Collaborative Pharmacy Practice, where a valid Patient-Practitioner relationship exists, that is communicated to a Pharmacist in a licensed Pharmacy.
- (r5) “Primary Care” is the first level of contact of individuals, the family, and the community with the health care delivery system, bringing health care as close as possible to where people live and work, and constitutes the first element of a continuing health care process. (Areas of Primary Care where Pharmacists provide Pharmacist Care Services include, but are not limited to, the following: chronic disease management; smoking cessation; maternal and child health; immunizations; family planning; self-care consulting; Drug selection under protocol; treatment of common diseases and injuries; nutrition; and general health education and promotion.)
- (s5) “Probation” is a restriction of Pharmacy practice for a specified period of time.<sup>36</sup>
- (t5) “Product” means a Prescription Drug in a finished dosage form for Administration to a Patient without substantial further Manufacturing (such as capsules, tablets, and lyophilized Products before reconstitution), but does not include:
- (1) blood or blood components intended for transfusion;
  - (2) radioactive Drugs or radioactive Biological Products;
  - (3) imaging Drugs;
  - (4) intravenous Product that are intended to:
    - (i) replenish fluids and electrolytes;
    - (ii) maintain the equilibrium of water and minerals; or
    - (iii) irrigate.
  - (5) any medical gas
  - (6) homeopathic Drugs marketed in accordance with applicable Federal law; or
  - (7) a Drug Compounded in compliance with Federal law.
- (u5) “Product Identifier” means a standardized graphic that includes, in both human-readable form and on a machine-readable data carrier that conforms to the standards developed by a widely recognized international standards development organization, the Standardized Numerical Identifier, lot number, and expiration date of the Product.

---

<sup>34</sup> The definition of “Practitioner” anticipates that those persons other than Pharmacists who are permitted to prescribe and Administer Drugs will be specifically so authorized in other legislation.

<sup>35</sup> Preceptors should be appropriately qualified and possess ample experience for the proper instructional training of Pharmacy Interns. It is strongly encouraged that Preceptors pursue continuing professional development for their practitioner-educator role expectations.

<sup>36</sup> Licensee may be placed on Probation for a period of time subject to specific conditions determined by the Board. Probation may result from the Board’s decision to stay a license Revocation or Suspension judgment. The licensee may be permitted to continue practice only within conditions established by the Board, and violation of those conditions will end the stay and result in Revocation or Suspension.



- (v5) “Product Labeling” means all labels and other written, printed, or graphic matter upon any article or any of its containers or wrappers, or accompanying such article.
- (w5) “Product Quality and Characteristics” include sterility, potency, identity, strength, quality, and purity associated with environmental quality, preparation activities, and checks and tests.
- (x5) “Professional Performance Evaluation” means a peer review process in which a competency assessment is made of a pharmacist by another pharmacist for the purpose of improving the quality of the evaluated pharmacist’s performance.
- (y5) “Prospective Drug Utilization Review (DUR)” means a review of the patient’s Drug therapy and Prescription Drug Order as part of a Drug Utilization Review, as defined in the rules of the Board, prior to Dispensing the Drug.
- (z5) “Protected Health Information” means Individually Identifiable Health Information:  
Except as provided in paragraph (2) of this definition, that is:
- (i) transmitted by electronic media;
  - (ii) maintained in any medium described in the definition of electronic media at §162.103 of the HIPAA privacy rules (45 CFR Part 160);
  - (iii) transmitted or maintained in any other form or medium.
- Protected health information excludes individually identifiable health information in:
- (iv) education records covered by the Family Educational Right and Privacy Act, as amended 20 USC 1232(g);
  - (v) records described at 20 USC 1232(g)(4)(B)(iv); and
  - (vi) employment records held by a licensee in its role as an employer.
- (See comment list.)
- (a6) “Public Health Emergency” means an imminent threat or occurrence of an illness or health condition caused by terrorism, bioterrorism, epidemic or pandemic disease, novel and highly fatal infectious agent or biological toxin, or natural or man-made disaster, that poses a substantial risk of significant number of human fatalities or incidents of permanent or long-term disability that is beyond the capacity of local government or nongovernmental organizations to resolve.
- (b6) “Qualified Licensed Professional” means a non-Pharmacist individual (such as a physician, nurse, or technologist) who possesses a current state license, if applicable, and who has sufficient training and experience to safely handle and Dispense radiopharmaceuticals as defined by the respective requirements of [cite appropriate Nuclear Regulatory Commission (NRC) or Agreement State and State Board of Pharmacy law(s)].
- (c6) “Qualified Nuclear Pharmacist” means a currently licensed Pharmacist in the State of practice, who is certified as a Nuclear Pharmacist by the State Board of Pharmacy or by a certification Board recognized by the State Board of Pharmacy, or who meets the following standards:
- (1) Minimum standards of training for “authorized user status” of radioactive material [cite State Radiation Control Agency or NRC licensure guide].
  - (2) Completed a minimum of 200 contact hours of instruction in nuclear Pharmacy and the safe handling and the use of radioactive materials from a program approved by the State Board of Pharmacy, with emphasis in the following areas:
    - (i) radiation physics and instrumentation;
    - (ii) radiation protection;
    - (iii) mathematics of radioactivity;
    - (iv) radiation biology; and
    - (v) radiopharmaceutical chemistry.
  - (3) Attained a minimum of 500 hours of clinical nuclear Pharmacy training under the supervision of a qualified nuclear Pharmacist.

- (d6) “Quality-Related Event” means any departure from the appropriate Dispensing of a prescribed medication that is or is not corrected prior to the Delivery and/or Administration of the medication.<sup>37</sup> The term “Quality-Related Event” includes:
- (1) a variation from the prescriber’s prescription drug order, including, but not limited to:
    - (i) incorrect Drug;
    - (ii) incorrect Drug strength;
    - (iii) incorrect dosage form;
    - (iv) incorrect patient; or
    - (v) inadequate or incorrect packaging, labeling, or directions;
  - (2) a failure to identify and manage:
    - (i) over-utilization or under-utilization;
    - (ii) therapeutic duplication;
    - (iii) drug-disease contraindications;
    - (iv) drug-drug interactions;
    - (v) incorrect drug dosage or duration of drug treatment;
    - (vi) drug-allergy interactions; or
    - (vii) clinical abuse/misuse.
  - (3) The term also includes packaging or warnings that fail to meet recognized standards, the Delivery of a medication to the wrong patient, and the failure to detect and appropriately manage a significant actual or potential problem with a patient’s drug therapy.
- (e6) “Quality Self-Audit” means an internal evaluation at a pharmacy to assess the effectiveness of the Continuous Quality Improvement (CQI) Program.
- (f6) “Radiopharmaceutical Quality Assurance” means, but is not limited to, the performance of appropriate chemical, biological, and physical tests on potential radiopharmaceuticals and the interpretation of the resulting data to determine their suitability for use in humans and animals, including internal test assessment, authentication of Product history, and the keeping of proper records.
- (g6) “Radiopharmaceutical Service” means, but shall not be limited to, the procurement, storage, handling, preparation, Labeling, quality assurance testing, Dispensing, Delivery, record keeping, and disposal of radiopharmaceuticals and other Drugs.
- (h6) “Radiopharmaceuticals” are radioactive Drugs as defined by Food and Drug Administration (FDA) and the State Board of Pharmacy [cite appropriate law(s)].
- (i6) “Reference Product” is the single Biological Product, already approved by FDA, against which a proposed Biosimilar Product is compared.
- (j6) “Remote Dispensing Site” means a location, other than where a Pharmacist is located, where Drugs are maintained and prescriptions are filled by a Certified Pharmacy Technician and Dispensed under the direct, remote supervision of a Pharmacist.
- (k6) “Repackage” means the act of taking a Drug Product from the container in which it was Distributed by the Manufacturer and placing it into a different container without further manipulation of the Drug. Repackaging also includes the act of placing the contents of multiple containers, eg, vials, of the same finished Drug into one container, providing the container does not include other ingredients or is not further manipulated in any way.
- (l6) “Repackager” means a Person who owns or operates an establishment that Repackages and relabels a Product or package for:
- (1) further sale; or
  - (2) Distribution without a further Transaction.<sup>38</sup>

---

<sup>37</sup> Quality-Related Events may be recorded using the Community Pharmacy Quality-Related Event Data Collection Form found in Appendix D.

<sup>38</sup> Is not intended to include a Pharmacy, Pharmacist, or Outsourcing Facility that Dispenses or Distributes Repackaged Drugs.

- (m6) “Repository Program” means a program that is established to receive previously dispensed medications and redispense such to qualified individuals and/or to facilitate the proper disposal of unacceptable medications in compliance with state and environmental regulations.
- (n6) “Reprimand” is a formal reproof of a licensee for violation of the Pharmacy Practice Act or rules and regulations.
- (o6) “Return” means providing Product to the authorized immediate Trading Partner from which such Product was purchased or received, or to a Returns Processor or Reverse Logistics Provider for handling of such Product.
- (p6) “Returns Processor or Reverse Logistics Provider” means any Person who owns or operates an establishment that disposes or otherwise processes saleable or nonsaleable Product received from an authorized Trading Partner such that the Product may be processed for credit to the purchaser, Manufacturer, or seller or disposed of for no further Distribution.
- (q6) “Revocation” is the withdrawal of the license to practice Pharmacy. The Person no longer has the privilege to practice in the State.
- (r6) “Risk Level” of the Sterile Pharmaceutical means the level assigned to a Sterile Pharmaceutical by a Pharmacist that represents the probability that the Sterile Pharmaceutical will be contaminated with microbial organisms, spores, endotoxins, foreign chemicals, or other physical matter.
- (s6) “Sales Unit” means the unit of measure the manufacturer uses to invoice its customer for the particular Product.
- (t6) “Shared Pharmacy Services” means a system that allows a participating Pharmacist or Pharmacy pursuant to a request from another participating Pharmacist or Pharmacy to process or fill a Prescription Drug Order, which may include preparing, packaging, Labeling, Compounding for specific patients, Dispensing, performing Drug Utilization Reviews, conducting claims adjudication, obtaining refill authorizations, reviewing therapeutic interventions, and/or reviewing institutional facility orders.
- (u6) “Significant Adverse Drug Reaction” means any Drug-related incident that may result in serious harm, injury, or death to the patient.
- (v6) “Significant Quality-Related Event” means any Quality-Related Event that results in serious harm, injury, or death to the patient.
- (w6) “Significant Loss” means any loss of a Prescription Drug that exceeds a reasonable level established by like persons, which requires that loss to be reported to the Board or as required by Drug Enforcement Administration (DEA) or other state and/or federal agencies for Prescription Drugs and controlled substances.<sup>39</sup>
- (x6) “Specialty Drug” means a Drug used to treat a chronic or specific disease or condition that requires frequent communication with other health care providers, extensive patient monitoring and case management, and comprehensive counseling with the patient and/or caregiver.
- (y6) “Specialty Pharmacy” means a Pharmacy that is providing Specialty Pharmacy Practice services and where Drugs, Devices, and other materials used in the diagnosis and treatment of injury, illness, and disease are Dispensed and Compounded.

---

<sup>39</sup> Some factors to consider in determining a Significant Loss include:

- (a) the actual quantity of Prescription Drugs or controlled substances lost in relation to the type of business;
- (b) the specific Prescription Drugs or controlled substances lost;
- (c) whether the loss of the Prescription Drugs or controlled substances can be associated with access to those Prescription Drugs or controlled substances by specific individuals, or whether the loss can be attributed to unique activities that may take place involving the Prescription Drugs or controlled substances;
- (d) a pattern of losses over a specific time period, whether the losses appear to be random, and the results of efforts taken to resolve the losses;
- (e) whether the specific Prescription Drugs or controlled substances are likely candidates for diversion; and
- (f) local trends and other indicators of the diversion potential of the missing Prescription Drug or controlled substance.

If it is determined that the loss is not significant, a record of the occurrence should be kept for future reference. When a Significant Loss occurs in a Pharmacy that is registered in multiple states, all applicable Boards should be notified.

- (z6) “Specialty Pharmacy Practice” means the provision of Pharmacist Care Services, which involves Drugs used to treat chronic or specific diseases and conditions that require frequent communication with other health care providers, extensive patient monitoring and case management, and comprehensive counseling with the patient and/or caregiver.<sup>40</sup> Drugs Dispensed by a Specialty Pharmacy may also require instruction and training on complex administration processes and/or handling and storage considerations.
- (a7) “Specific Patient Need” means the transfer of a Product from one Pharmacy to another to fill a Prescription for an identified patient, but does not include the transfer of Product from one Pharmacy to another for the purpose of increasing or replenishing stock in anticipation of a potential need.
- (b7) “Standard of Care” means the degree of care a prudent and reasonable licensee or registrant with similar education, training, and experience will exercise under similar circumstances.
- (c7) “Standardized Numerical Identifier” means a set of numbers or characters used to uniquely identify each package or homogeneous case that is composed of the National Drug Code that corresponds to the specific Product, including the particular package configuration, combined with a unique alphanumeric serial number of up to 20 characters.
- (d7) “State of Emergency” means a governmental declaration, usually issued as a result of a Public Health Emergency, that may suspend certain normal functions of government, alert citizens to alter their normal behaviors, and/or direct government agencies to implement emergency preparedness plans.
- (e7) “Sterile Pharmaceutical” means any dosage form of a drug, including but not limited to, parenterals (eg, injectables, surgical irrigants, and ophthalmics) devoid of viable microorganisms.
- (f7) “Summary Suspension” is the Suspension of a license, which requires a licensee to cease Pharmacy practice immediately pending the results of a timely hearing.<sup>41</sup>
- (g7) “Suspect Product” means a Product for which there is reason to believe that such Product:
- (1) is potentially Counterfeit, diverted, or stolen;
  - (2) is potentially intentionally Adulterated such that the Product would result in serious adverse health consequences or death to humans;
  - (3) is potentially the subject of a fraudulent Transaction; or
  - (4) appears otherwise unfit for Distribution such that the Product would result in serious adverse health consequences or death to humans.
- (h7) “Suspension” is the withdrawal of the license to practice Pharmacy in the State for a specified period of time.
- (i7) “Suspicious Order” includes, but is not limited to, an unsubstantiated order with the following characteristic(s):
- (1) unusual size or frequency; or
  - (2) deviating substantially from a normal pattern.
- (j7) “Telepharmacy Technologies” means secure electronic communications, information exchange, or other methods that meet applicable state and federal requirements.
- (k7) “Temporary Pharmacy Facility” means a facility established as a result of a Public Health Emergency or State of Emergency to temporarily provide Pharmacy services within or adjacent to Declared Disaster Areas.
- (l7) “Therapeutic Interchange” means substitution by the Pharmacist of one medication for another medication with a similar therapeutic effect, at the time of dispensing.
- (m7) “Third-Party Logistics Provider” means an entity that:

---

<sup>40</sup> It should be noted that the rationale for Dispensing Drugs from a Specialty Pharmacy be based on the Drugs possessing certain characteristics such as a very narrow therapeutic range, life-endangering side effect profile, and/or requiring periodic laboratory or diagnostic testing, and that such characteristics require extensive Pharmacist interaction with both the patient and health care provider(s).

<sup>41</sup> If the Board believes it necessary to protect the public health and safety, it may summarily Suspend a license and order a prompt hearing on the matters in question.

- (1) Provides or coordinates warehousing, Distribution, or other services on behalf of a Manufacturer, but does not take title to the Prescription Drug or have general responsibility to direct the Prescription Drug's sale or disposition; and
  - (2) Is licensed as a Third-Party Logistics Provider.
- (n7) "Trading Partner" means:
- (1) a Manufacturer, Repackager, Wholesale Distributor, or Dispenser from whom a Manufacturer, Repackager, Wholesale Distributor, or Dispenser accepts direct ownership of a Product or to whom a Manufacturer, Repackager, Wholesale Distributor, or Dispenser transfers direct ownership of a Product; or
  - (2) a Third-Party Logistics Provider from whom a Manufacturer, Repackager, Wholesale Distributor, or Dispenser accepts direct possession of a Product or to whom a Manufacturer, Repackager, Wholesale Distributor, or Dispenser transfers direct possession of a Product.
- (o7) "Transaction" means the transfer of Product between Persons in which a change of ownership occurs. Transaction does not include:
- (1) intracompany Distribution of any Product between members of an affiliate or within a Manufacturer;
  - (2) the Distribution of a Product among hospitals or other Health Care Entities that are under common control;
  - (3) the Distribution of a Product for Emergency Medical Reasons, including a Public Health Emergency declaration pursuant to State or Federal law, except that a Drug shortage not caused by a Public Health Emergency shall not constitute an Emergency Medical Reason;
  - (4) the Dispensing of a Product pursuant to a Prescription;
  - (5) the Distribution of Product samples by a Manufacturer or a licensed Wholesale Distributor in accordance with State and Federal law;
  - (6) the Distribution of blood or blood components intended for transfusion;
  - (7) the Distribution of minimal quantities of Product by a licensed retail Pharmacy to a licensed Practitioner for office use;
  - (8) the sale, purchase, or trade of a Drug or an offer to sell, purchase, or trade a Drug by a charitable organization to a nonprofit affiliate of the organization to the extent otherwise permitted by State and Federal law;
  - (9) the Distribution of a Product pursuant to the sale or merger of a Pharmacy or Pharmacies or a Wholesale Distributor or Wholesale Distributors, except that any records required to be maintained for the Product shall be transferred to the new owner of the Pharmacy or Pharmacies or Wholesale Distributor or Wholesale Distributors;
  - (10) the Dispensing of a new animal Drug Product approved under Federal law;
  - (11) Products transferred to or from any facility that is licensed by the Nuclear Regulatory Commission or by a State pursuant to an agreement with such Commission;
  - (12) a combination Product that is:
    - (i) a Product composed of a Device and one or more other regulated Components (such as a Drug/Device, biologic/Device, or Drug/Device/biologic) that are physically, chemically, or otherwise combined or mixed and produced as a single entity;
    - (ii) two or more separate Products packaged together in a single package or as a unit and composed of a Drug and Device or a Device and Biological Product; or
    - (iii) two or more finished medical Devices plus one or more Drug or Biological Products that are packaged together in what is referred to as a "medical convenience kit" as described in (w6)(13);
  - (13) the Distribution of a collection of finished medical Devices, which may include a Product or Biological Product, assembled in kit form strictly for the convenience of the purchaser or user if:

- (i) the medical convenience kit is assembled in an establishment that is registered with FDA as a Device Manufacturer;
- (ii) the medical convenience kit does not contain a federally scheduled controlled substance;
- (iii) in the case of a medical convenience kit that includes a Product, the Person who Manufactured the kit:
  - (A) purchased such Product directly from the pharmaceutical Manufacturer or from a Wholesale Distributor that purchased the Product directly from the pharmaceutical Manufacturer;
  - (B) does not alter the primary container or label of the Product as purchased from the Manufacturer or Wholesale Distributor; and
- (iv) in the case of a medical convenience kit that includes a Product, the Product is:
  - (A) an intravenous solution intended for the replenishment of fluids and electrolytes;
  - (B) a Product intended to maintain the equilibrium of water and minerals in the body;
  - (C) a Product intended for irrigation or reconstitution;
  - (D) an anesthetic;
  - (E) an anticoagulant;
  - (F) a vasopressor; or
  - (G) a sympathomimetic;
- (14) the Distribution of an intravenous Product that, by its formulation, is intended for the replenishment of fluids and electrolytes (such as sodium, chloride, and potassium) or calories (such as dextrose and amino acids);
- (15) the Distribution of an intravenous Product used to maintain the equilibrium of water and minerals in the body, such as dialysis solutions;
- (16) the Distribution of a Product that is intended for irrigation, or sterile water, whether intended for such purposes or for injection;
- (17) the Distribution of a medical gas;
- (18) the Distribution or sale of any licensed biologic Product that meets the definition of Device under Federal law.

(p7) “Transaction History” means a statement in paper or electronic form that includes the Transaction Information of each prior Transaction going back to the Manufacturer of the Product.

(q7) “Transaction Information” means:

- (1) the proprietary or established name or names of the Product;
- (2) the strength and dosage form of the Product;
- (3) the National Drug Code number of the Product;
- (4) the container size;
- (5) the number of containers;
- (6) the lot number of the Product;
- (7) the Transaction date;
- (8) the shipment date, if more than 24 hours after the Transaction date;
- (9) the business name and address of the transferring Person; and
- (10) the business name and address of the transferee Person.

(r7) “Transaction Statement” is a statement, in paper or electronic form, that the entity transferring ownership in a Transaction:

- (1) is authorized under Federal law;
- (2) received the Product from a Person who is authorized as required under Federal law;
- (3) received Transaction Information and Transaction Statement from the prior owner of the Product, as required by Federal law;
- (4) did not knowingly ship a Suspect or Illegitimate Product;

- (5) had systems and processes in place to comply with Verification requirements outlined in Federal law;
- (6) did not knowingly provide false Transaction Information; and
- (7) did not knowingly alter the Transaction History.
- (s7) “USP Standards” means standards published in the current official United States Pharmacopeia or National Formulary.
- (t7) “Valid Patient-Practitioner Relationship”<sup>42</sup> means the following have been established:
  - (1) a Patient has a medical complaint;
  - (2) a medical history has been taken;
  - (3) a face-to-face physical examination adequate to establish the medical complaint has been performed by the prescribing practitioner or, in the instances of telemedicine, through telemedicine practice approved by the appropriate Practitioner Board; and
  - (4) some logical connection exists between the medical complaint, the medical history, and the physical examination and the Drug prescribed.
- (u7) “Verification” means determining whether the Product Identifier of a package or homogeneous case corresponds to the Standardized Numerical Identifier or lot number and expiration date assigned to the Product by the Manufacturer or Repackager in accordance with Federal law.
- (v7) “Veterinary Dispensing” means the interpretation, evaluation, and implementation of a veterinary Prescription Drug Order, including the preparation, final Verification, and Delivery of a Drug for a veterinary patient in a suitable container appropriately labeled for the client for subsequent Administration.
- (w7) “Virtual Manufacturer” means a Manufacturer that sells its own Prescription Drugs and/or Devices but never physically possesses the Product.
- (x7) “Virtual Wholesale Distributor” means a Wholesale Distributor that sells a Prescription Drug or Device but never physically possess the Product.
- (y7) “Warning” is a written notice issued to a licensee addressing possible errant conduct.<sup>43</sup>
- (z7) “Wholesale Distribution” means the Distribution of a Drug or Device to a Person other than a consumer or patient, or receipt of a Drug or Device by a Person other than the consumer or patient, but does not include:<sup>44</sup>

---

<sup>42</sup> A Valid Patient-Practitioner Relationship includes a relationship with a consulting Practitioner or a Practitioner to which a patient has been referred, or a covering Practitioner, or an appropriate Practitioner-Board-approved telemedicine Practitioner providing that a physical examination had been previously performed by the patient’s Practitioner.

To best protect the public, the issue of a Valid Patient-Practitioner Relationship should be addressed in each jurisdiction’s Medical Practice Act and the Consumer Fraud Protection Act or their equivalent.

A face-to-face physical examination is not required to establish a Valid Patient-Practitioner Relationship if:

- (a) the prescribing Practitioner is issuing a prescription or Dispensing a non-controlled substance legend drug in accordance with the Expedited Partner Therapy in the Management of Sexually Transmitted Diseases Guidance document issued by the United States Centers for Disease Control and Prevention;
- (b) the prescription, Administration, or Dispensing is through a public health clinic or other distribution mechanism approved by the state health authority in order to prevent, mitigate, or treat a pandemic illness, infectious disease outbreak, or intentional or accidental release of a biological, chemical, or radiological agent; or
- (c) the prescribing Practitioner is issuing a prescription through a telemedicine practice approved by the appropriate state agency that provides health care delivery, diagnosis, consultation, or treatment by means of audio, video, or data communications. Standard telephone, facsimile transmission, or both, in the absence of other integrated information or data, do not constitute telemedicine practices.
- (d) the State allows third-party prescribing of opioid reversal agents, such as naloxone, or other drugs as allowed by State law to a person other than the patient.

<sup>43</sup> A Warning may require that the licensee provide the Board with clarifying information. (May also be known as a Letter of Concern or Letter of Admonition.)

<sup>44</sup> Although “Devices” is included in both definitions of “Wholesale Distribution” and “Wholesale Distributor,” Federal law and some State laws do not define “Wholesale Distribution” as such. Wherever appropriate under the Model Rules, the term is included and recognized that Wholesale Distribution also includes Devices. A disparity could be caused if those Persons who only distribute Devices are not currently licensed by the State and, therefore, not subject to regulation

- (1) intracompany Distribution of any Drug between members of an affiliate or within a Manufacturer;
- (2) the Distribution of a Drug or an offer to Distribute a Drug among hospitals or other Health Care Entities that are under common control;
- (3) the Distribution of a Drug or an offer to Distribute a Drug for Emergency Medical Reasons, including a Public Health Emergency declaration made by the Secretary of the United States Department of Health and Human Services, except that, for purposes of this paragraph, a Drug shortage not caused by a Public Health Emergency shall not constitute an Emergency Medical Reason;
- (4) the Dispensing of a Drug pursuant to a Prescription Drug Order;
- (5) the Distribution of minimal quantities of a Drug by a licensed retail Pharmacy to a licensed Practitioner for office use;<sup>45</sup>
- (6) the Distribution of a Drug or an offer to Distribute a Drug by a charitable organization to a nonprofit affiliate of the organization to the extent otherwise permitted by law;
- (7) the purchase or other acquisition by a Dispenser, hospital, or other Health Care Entity of a Drug for use by such Dispenser, hospital, or other Health Care Entity;
- (8) the Distribution of a Drug by the Manufacturer of such Drug;
- (9) the receipt or transfer of a Drug by an authorized Third-Party Logistics Provider, provided that such Third-Party Logistics Provider does not take ownership of the Drug;
- (10) a Common Carrier that transports a Drug, provided that the Common Carrier does not take ownership of the Drug;
- (11) the Distribution of a Drug or an offer to Distribute a Drug by an authorized Repackager that has taken ownership or possession of the Drug and Repackages it in accordance with Federal law;
- (12) salable Drug Returns when conducted by a Dispenser;
- (13) the Distribution of a collection of finished medical Devices, which may include a Drug Product or Biological Product, assembled in kit form strictly for the convenience of the purchaser or user (referred to in this subparagraph as a “medical convenience kit”) if:
  - (i) the medical convenience kit is assembled in an establishment that is registered with FDA as a Device Manufacturer;
  - (ii) the medical convenience kit does not contain a controlled substance;
  - (iii) in the case of a medical convenience kit that includes a Drug Product, the Person that Manufactures the kit:
    - (A) purchased such Drug Product directly from the pharmaceutical Manufacturer or from a Wholesale Distributor that purchased the Drug Product directly from the pharmaceutical Manufacturer; and
    - (B) does not alter the primary container or Label of the Drug Product as purchased from the Manufacturer or Wholesale Distributor; and
  - (iv) in the case of a medical convenience kit that includes a Drug Product, the Drug Product is:
    - (A) an intravenous solution intended for the replenishment of fluids and electrolytes;
    - (B) a Product intended to maintain the equilibrium of water and minerals in the body;
    - (C) a Product intended for irrigation or reconstitution;
    - (D) an anesthetic;
    - (E) an anticoagulant;
    - (F) a vasopressor; or
    - (G) a sympathomimetic.

---

by the Board. Different requirements and standards would exist for these Persons than would apply for Persons who Distribute both Drugs and Devices. It is NABP’s position that Persons who Manufacture and/or Distribute Devices should be licensed with the Board and adhere to the same requirements as those in place for Persons who Manufacture and/or Distribute Drugs. In developing laws and rules, States may need to review their current regulations regarding licensure for Persons who solely Manufacture and/or Distribute Devices in order to determine the applicability of the Model Rules to Persons who Manufacture and/or Distribute Devices.

<sup>45</sup> Excludes Compounded Drugs unless the Pharmacy is registered under Federal law and Distributing such Compounded Drugs as an Outsourcing Facility.



- (14) the Distribution of an intravenous Drug that, by its formulation, is intended for the replenishment of fluids and electrolytes (such as sodium, chloride, and potassium) or calories (such as dextrose and amino acids);
- (15) the Distribution of an intravenous Drug used to maintain the equilibrium of water and minerals in the body, such as dialysis solutions;
- (16) the Distribution of a Drug that is intended for irrigation, or sterile water, whether intended for such purposes or for injection;
- (17) the Distribution of medical gas;
- (18) facilitating the Distribution of a Product by providing solely administrative services, including processing of orders and payments; or
- (19) the transfer of a Product by a hospital or other Health Care Entity, or by a Wholesale Distributor or Manufacturer operating at the direction of the hospital or other Health Care Entity, to a Repackager and registered with FDA for the purpose of Repackaging the Drug for use by that hospital or other Health Care Entity and other Health Care Entities that are under common control, if ownership of the Drug remains with the hospital or other Health Care Entity at all times.

(a8) “Wholesale Distributor” means any Person, which may include a Virtual Wholesale Distributor, (other than a Manufacturer, a Manufacturer’s co-licensed partner, a Third-Party Logistics Provider, or Repackager) engaged in Wholesale Distribution of Prescription Drugs or Devices in or into the State.

## **Section 105(l2). Comment.**

45 CFR §164.514(b) reads:

“requirements for de-identification of protected health information. A covered entity may determine that health information is not individually identifiable health information only if:

(1) A person with appropriate knowledge of and experience with generally accepted statistical and scientific principles and methods for rendering information not individually identifiable:

(i) Applying such principles and methods, determines that the risk is very small that the information could be used, alone or in combination with other reasonably available information, by an anticipated recipient to identify an individual who is a subject of the information; and

(ii) Documents the methods and results of the analysis that justify such determination; or

(2)(i) The following identifiers of the individual or of relatives, employers, or household members of the individual, are removed:

(A) Names;

(B) All geographic subdivisions smaller than a State, including street address, city, county, precinct, zip code, and their equivalent geocodes, except for the initial three digits of a zip code if, according to the current publicly available data from the Bureau of the Census:

(1) The geographic unit formed by combining all zip codes with the same three initial digits contains more than 20,000 people; and

(2) The initial three digits of a zip code for all such geographic units containing 20,000 or fewer people is changed to 000.

(C) All elements of dates (except year) for dates directly related to an individual, including birth date, admission date, discharge date, date of death; and all ages over 89 and all elements of dates (including year) indicative of such age, except that such ages and elements may be aggregated into a single category of age 90 or older;

(D) Telephone numbers;

(E) Fax numbers;

(F) Electronic mail addresses;

(G) Social security numbers;

(H) Medical record numbers;

(I) Health plan beneficiary numbers;

(J) Account numbers;

(K) Certificate/license numbers;

(L) Vehicle identifiers and serial numbers, including license plate numbers;

(M) Device identifiers and serial numbers;

(N) Web Universal Resource Locators (URLs);

(O) Internet Protocol (IP) address numbers;

(P) Biometric identifiers, including finger and voice prints;

(Q) Full face photographic images and any comparable images; and

(R) Any other unique identifying number, characteristic, or code, except as permitted by paragraph (c) of this section; and

(ii) The covered entity does not have actual knowledge that the information could be used alone or in combination with other information to identify an individual who is a subject of the information.”

## **Section 105(z5). Comment.**

45 CFR §162.103 reads as follows:

Electronic media means the mode of electronic transmission. It includes the Internet (wide-open), Extranet (using Internet technology to link a business with information only accessible to collaborating parties), leased lines, dial-up lines, private networks, and those transmissions that are physically moved from one location to another using magnetic tape, disk, or compact disk media.

20 USC §1232g reads as follows:

Sec. 1232g. Family educational and privacy rights

(a) Conditions for availability of funds to educational agencies or institutions; inspection and review of education records; specific information to be made available; procedure for access to education records; reasonableness of time for such access; hearings; written explanations by parents; definitions

(1)(A) No funds shall be made available under any applicable program to any educational agency or institution which has a policy of denying, or which effectively prevents, the parents of students who are or have been in attendance at a school of such agency or at such institution, as the case may be, the right to inspect and review the education records of their children. If any material or document in the education record of a student includes information on more than one student, the parents of one of such students shall have the right to inspect and review only such part of such material or document as relates to such student or to be informed of the specific information contained in such part of such material. Each educational agency or institution shall establish appropriate procedures for the granting of a request by parents for access to the education records of their children within a reasonable period of time, but in no case more than forty-five days after the request has been made.

(B) No funds under any applicable program shall be made available to any State educational agency (whether or not that agency is an educational agency or institution under this section) that has a policy of denying, or effectively prevents, the parents of students the right to inspect and review the education records maintained by the State educational agency on their children who are or have been in attendance at any school of an educational agency or institution that is subject to the provisions of this section.

(C) The first sentence of subparagraph (A) shall not operate to make available to students in institutions of postsecondary education the following materials:

(i) financial records of the parents of the student or any information contained therein;

(ii) confidential letters and statements of recommendation, which were placed in the education records prior to January 1, 1975, if such letters or statements are not used for purposes other than those for which they were specifically intended;

(iii) if the student has signed a waiver of the student's right of access under this subsection in accordance with

subparagraph (D), confidential recommendations –

(I) respecting admission to any educational agency or institution,

(II) respecting an application for employment, and

(III) respecting the receipt of an honor or honorary recognition

(D) A student or a person applying for admission may waive his right of access to confidential statements described in clause (iii) of subparagraph (C), except that such waiver shall apply to recommendations only if

(i) the student is, upon request, notified of the names of all persons making confidential recommendations and

(ii) such recommendations are used solely for the purpose for which they were specifically intended. Such waivers may not be required as a condition for admission to, receipt of financial aid from, or receipt of any other services or benefits from such agency or institution.

(2) No funds shall be made available under any applicable program to any educational agency or institution unless the parents of students who are or have been in attendance at a school of such agency or at such institution are provided an opportunity for a hearing by such agency or institution, in accordance with regulations of the Secretary, to challenge the content of such student's education records, in order to insure that the records are not inaccurate, misleading, or otherwise in violation of the privacy rights of students, and to provide an opportunity for the correction or deletion of any such inaccurate, misleading or otherwise inappropriate data contained therein and to insert into such records a written explanation of the parents respecting the content of such records.

(3) For the purposes of this section the term "educational agency or institution" means any public or private agency or institution which is the recipient of funds under any applicable program.

(4)(A) For the purposes of this section, the term "education records" means, except as may be provided otherwise in subparagraph (B), those records, files, documents, and other materials which –

(i) contain information directly related to a student; and

(ii) are maintained by an educational agency or institution or by a person acting for such agency or institution.

(B) The term "education records" does not include –

(i) records of instructional, supervisory, and administrative personnel and educational personnel ancillary thereto which are in the sole possession of the maker thereof and which are not accessible or revealed to any other person except a substitute;

(ii) records maintained by a law enforcement unit of the educational agency or institution that were created by that law enforcement unit for the purpose of law enforcement;

(iii) in the case of persons who are employed by an educational agency or institution but who are not in attendance at such agency or institution, records made and maintained in the normal course of business which relate exclusively to such person in that person's capacity as an employee and are not available for use for any other purpose; or  
(iv) records on a student who is eighteen years of age or older, or is attending an institution of postsecondary education, which are made or maintained by a physician, psychiatrist, psychologist, or other recognized professional or paraprofessional acting in his professional or paraprofessional capacity, or assisting in that capacity, and which are made, maintained, or used only in connection with the provision of treatment to the student, and are not available to anyone other than persons providing such treatment, except that such records can be personally reviewed by a physician or other appropriate professional of the student's choice.

(5)(A) For the purposes of this section the term "directory information" relating to a student includes the following: the student's name, address, telephone listing, date and place of birth, major field of study, participation in officially recognized activities and sports, weight and height of members of athletic teams, dates of attendance, degrees and awards received, and the most recent previous educational agency or institution attended by the student.

(B) Any educational agency or institution making public directory information shall give public notice of the categories of information which it has designated as such information with respect to each student attending the institution or agency and shall allow a reasonable period of time after such notice has been given for a parent to inform the institution or agency that any or all of the information designated should not be released without the parent's prior consent.

(6) For the purposes of this section, the term "student" includes any person with respect to whom an educational agency or institution maintains education records or personally identifiable information, but does not include a person who has not been in attendance at such agency or institution.

(b) Release of education records; parental consent requirement; exceptions; compliance with judicial orders and subpoenas; audit and evaluation of federally-supported education programs; record keeping

(1) No funds shall be made available under any applicable program to any educational agency or institution which has a policy or practice of permitting the release of education records (or personally identifiable information contained therein other than directory information, as defined in paragraph (5) of subsection (a) of this section) of students without the written consent of their parents to any individual, agency, or organization, other than to the following –

(A) other school officials, including teachers within the educational institution or local educational agency, who have been determined by such agency or institution to have legitimate educational interests, including the educational interests of the child for whom consent would otherwise be required;

(B) officials of other schools or school systems in which the student seeks or intends to enroll, upon condition that the student's parents be notified of the transfer, receive a copy of the record if desired, and have an opportunity for a hearing to challenge the content of the record;

(C)(i) authorized representatives of (I) the Comptroller General of the United States, (II) the Secretary, or (III) State educational authorities, under the conditions set forth in paragraph (3), or

(ii) authorized representatives of the Attorney General for law enforcement purposes under the same conditions as apply to the Secretary under paragraph (3);

(D) in connection with a student's application for, or receipt of, financial aid;

(E) State and local officials or authorities to whom such information is specifically allowed to be reported or disclosed pursuant to State statute adopted –

(i) before November 19, 1974, if the allowed reporting or disclosure concerns the juvenile justice system and such system's ability to effectively serve the student whose records are released, or

(ii) after November 19, 1974, if –

(I) the allowed reporting or disclosure concerns the juvenile justice system and such system's ability to effectively serve, prior to adjudication, the student whose records are released; and

(II) the officials and authorities to whom such information is disclosed certify in writing to the educational agency or institution that the information will not be disclosed to any other party except as provided under State law without the prior written consent of the parent of the student.

(F) organizations conducting studies for, or on behalf of, educational agencies or institutions for the purpose of developing, validating, or administering predictive tests, administering student aid programs, and improving instruction, if such studies are conducted in such a manner as will not permit the personal identification of students and their parents by persons other than representatives of such organizations and such information will be destroyed when no longer needed for the purpose for which it is conducted;

(G) accrediting organizations in order to carry out their accrediting functions;

(H) parents of a dependent student of such parents, as defined in section 152 of title 26;

(I) subject to regulations of the Secretary, in connection with an emergency, appropriate persons if the knowledge of such information is necessary to protect the health or safety of the student or other persons; and

(J)(i) the entity or persons designated in a Federal grand jury subpoena, in which case the court shall order, for good cause shown, the educational agency or institution (and any officer, director, employee, agent, or attorney for such agency or institution) on which the subpoena is served, to not disclose to any person the existence or contents of the subpoena or any information furnished to the grand jury in response to the subpoena; and

(ii) the entity or persons designated in any other subpoena issued for a law enforcement purpose, in which case the court or other issuing agency may order, for good cause shown, the educational agency or institution (and any officer, director, employee, agent, or attorney for such agency or institution) on which the subpoena is served, to not disclose to any person the existence or contents of the subpoena or any information furnished in response to the subpoena. Nothing in clause (E) of this paragraph shall prevent a State from further limiting the number or type of State or local officials who will continue to have access thereunder.

(2) No funds shall be made available under any applicable program to any educational agency or institution which has a policy or practice of releasing, or providing access to, any personally identifiable information in education records other than directory information, or as is permitted under paragraph (1) of this subsection, unless –

(A) there is written consent from the student's parents specifying records to be released, the reasons for such release, and to whom, and with a copy of the records to be released to the student's parents and the student if desired by the parents, or

(B) except as provided in paragraph (1)(J), such information is furnished in compliance with judicial order, or pursuant to any lawfully issued subpoena, upon condition that parents and the students are notified of all such orders or subpoenas in advance of the compliance therewith by the educational institution or agency.

(3) Nothing contained in this section shall preclude authorized representatives of (A) the Comptroller General of the United States, (B) the Secretary, or (C) State educational authorities from having access to student or other records which may be necessary in connection with the audit and evaluation of Federally-supported education programs, or in connection with the enforcement of the Federal legal requirements which relate to such programs: Provided, That except when collection of personally identifiable information is specifically authorized by Federal law, any data collected by such officials shall be protected in a manner which will not permit the personal identification of students and their parents by other than those officials, and such personally identifiable data shall be destroyed when no longer needed for such audit, evaluation, and enforcement of Federal legal requirements.

(4)(A) Each educational agency or institution shall maintain a record, kept with the education records of each student, which will indicate all individuals (other than those specified in paragraph (1)(A) of this subsection), agencies, or organizations which have requested or obtained access to a student's education records maintained by such educational agency or institution, and which will indicate specifically the legitimate interest that each such person, agency, or organization has in obtaining this information. Such record of access shall be available only to parents, to the school official and his assistants who are responsible for the custody of such records, and to persons or organizations authorized in, and under the conditions of, clauses (A) and (C) of paragraph (1) as a means of auditing the operation of the system.

(B) With respect to this subsection, personal information shall only be transferred to a third party on the condition that such party will not permit any other party to have access to such information without the written consent of the parents of the student. If a third party outside the educational agency or institution permits access to information in violation of paragraph (2)(A), or fails to destroy information in violation of paragraph (1)(F), the educational agency or institution shall be prohibited from permitting access to information from education records to that third party for a period of not less than five years.

(5) Nothing in this section shall be construed to prohibit State and local educational officials from having access to student or other records which may be necessary in connection with the audit and evaluation of any federally or State supported education program or in connection with the enforcement of the Federal legal requirements which relate to any such program, subject to the conditions specified in the proviso in paragraph (3).

(6)(A) Nothing in this section shall be construed to prohibit an institution of postsecondary education from disclosing, to an alleged victim of any crime of violence (as that term is defined in section 16 of title 18), or a nonforcible sex offense, the final results of any disciplinary proceeding conducted by such institution against the alleged perpetrator of such crime or offense with respect to such crime or offense.

(B) Nothing in this section shall be construed to prohibit an institution of postsecondary education from disclosing the final results of any disciplinary proceeding conducted by such institution against a student who is an alleged perpetrator of any crime of violence (as that term is defined in section 16 of title 18), or a nonforcible sex offense, if the institution

determines as a result of that disciplinary proceeding that the student committed a violation of the institution's rules or policies with respect to such crime or offense.

(C) For the purpose of this paragraph, the final results of any disciplinary proceeding

(i) shall include only the name of the student, the violation committed, and any sanction imposed by the institution on that student; and

(ii) may include the name of any other student, such as a victim or witness, only with the written consent of that other student.

(7)(A) Nothing in this section may be construed to prohibit an educational institution from disclosing information provided to the institution under section 14071 of title 42 concerning registered sex offenders who are required to register under such section.

(B) The Secretary shall take appropriate steps to notify educational institutions that disclosure of information described in subparagraph

(A) is permitted.

(c) Surveys or data-gathering activities; regulations not later than 240 days after October 20, 1994, the Secretary shall adopt appropriate regulations or procedures, or identify existing regulations or procedures, which protect the rights of privacy of students and their families in connection with any surveys or data-gathering activities conducted, assisted, or authorized by the Secretary or an administrative head of an education agency. Regulations established under this subsection shall include provisions controlling the use, dissemination, and protection of such data. No survey or data-gathering activities shall be conducted by the Secretary, or an administrative head of an education agency under an applicable program, unless such activities are authorized by law.

(d) Students' rather than parents' permission or consent; for the purposes of this section, whenever a student has attained eighteen years of age, or is attending an institution of postsecondary education, the permission or consent required of and the rights accorded to the parents of the student shall thereafter only be required of and accorded to the student.

(e) Informing parents or students of rights; under this section, no funds shall be made available under any applicable program to any educational agency or institution unless such agency or institution effectively informs the parents of students, or the students, if they are eighteen years of age or older, or are attending an institution of postsecondary education, of the rights accorded them by this section.

(f) Enforcement; termination of assistance; the Secretary shall take appropriate actions to enforce this section and to deal with violations of this section, in accordance with this chapter, except that action to terminate assistance may be taken only if the Secretary finds there has been a failure to comply with this section, and he has determined that compliance cannot be secured by voluntary means.

(g) Office and review board; creation; functions the Secretary shall establish or designate an office and review board within the Department for the purpose of investigating, processing, reviewing, and adjudicating violations of this section and complaints which may be filed concerning alleged violations of this section. Except for the conduct of hearings, none of the functions of the Secretary under this section shall be carried out in any of the regional offices of such Department.

## **Article II**

### **Board of Pharmacy**

#### **Introductory Comment to Article II**

*Before it can regulate the Practice of Pharmacy, the State must first establish and empower the Board of Pharmacy. Accordingly, Article II of the Model Act defines and creates the Board of Pharmacy by specifying elements necessary to its formation, organization, and operation.*

*Each of the sections contained in this article covers elements that NABP felt necessary to the proper formation and efficient operation of the Board. Several of these sections, especially those that contain innovative or infrequently utilized provisions, are supplemented by individual explanatory comments.*

*Among the sections of Article II that may be of particular interest to users of the Model Act are the following: Sections 202 and 203(b), pertaining to the inclusion of public members as Board members; Section 207, which provides grounds and procedures for removal of Board members; and Section 213(c)(2), which enables Boards to avail themselves of research and study grants and other non-State monies.*

*It is also important to note that Section 212 specifically empowers the Board to make such rules as are necessary to fully administer and implement the Act. This is a most significant feature of the Model Act. The underlying philosophy of this approach is that the statute should create objectives, guidelines, and policies in general areas, and permit the Board to provide the specifics in its rules. This approach recognizes that it is impossible for State legislatures to enact comprehensive provisions regarding all of the matters with which a Board of Pharmacy may be confronted or to anticipate the rapidly changing conditions of the professions and the delivery of health care. Consequently, NABP recommends that Boards have adequate power to adopt and amend rules with the greatest possible flexibility and autonomy. Section 212 of the Model Act accomplishes this objective.*

*As noted in the findings of the 1990 report on the “State Discipline of Pharmacists” by the Federal Health and Human Services Department, Office of Inspector General (OIG), “The ability of many State Pharmacy Boards to protect the public is hampered by limitations in their legal authorities, administrative processes, and resources.” Based on these findings, the OIG recommended that, “State governments should ensure that State Pharmacy Boards have adequate resources and authority for carrying out their enforcement responsibilities effectively.”*

#### **Section 201. Designation.**

The responsibility for enforcement of the provisions of this Act is hereby vested in the Board of Pharmacy. The Board shall have all of the duties, powers, and authority specifically granted by or necessary for the enforcement of this Act, as well as such other duties, powers, and authority as it may be granted from time to time by applicable law. In the event of a declared State of Emergency, the Board may waive the requirements of this Act in order to protect the public health, safety, or welfare of its citizens and to facilitate the provision of Drugs, Devices, and Pharmacist Care Services to the public.<sup>46</sup>

---

<sup>46</sup> In states where Centralized Prescription Filling or Centralized Prescription Processing are not permitted, states may consider allowing the performance of such activities in a declared State of Emergency.

## Section 202. Membership.

The Board of Pharmacy shall consist of \_\_\_\_\_ members, \_\_\_\_\_ of whom shall be a representative of the public, and the remainder] [each] of whom shall be Pharmacists who possess the qualifications specified in Section 203.<sup>47</sup>

## Section 203. Qualifications.

- (a) Each Pharmacist member of the Board of Pharmacy shall at the time of appointment<sup>48</sup>:
  - (1) be a resident of this State for not less than six months;
  - (2) be currently licensed and in good standing to engage in the Practice of Pharmacy in this State;
  - (3) be actively engaged in the Practice of Pharmacy in this State;
  - (4) have five (5) years of experience in the Practice of Pharmacy in this State after licensure.
- (b) The public member of the Board of Pharmacy shall be a resident of this State who has attained the age of majority and shall not be, nor shall ever have been, a Pharmacist, or the spouse of a Pharmacist, or a Person who has ever had any material financial interest in the provision of Pharmacy services or who has engaged in any activity directly related to the Practice of Pharmacy.<sup>49</sup>

## Section 204. Appointment.

- (a) The Governor shall appoint the members of the Board of Pharmacy in accordance with other provisions of this Section and the State Constitution.
- (b) Nominations for appointment to the Board may be made to the Governor by any individual, association, or any other entity. Such nominations shall be recommendations only and shall not be binding in any manner upon the Governor.<sup>50</sup>

---

<sup>47</sup> The number of Board members should be determined by each individual state according to its particular requirements. Individual states may wish to consider a Board composition that represents the diversity of practice sites and interests within a state. Variable factors, such as state population, number of Pharmacists, number of pharmacies, and other local considerations, may all be relevant in determining the number of Board members needed to most effectively enforce the Act. In the event a state prefers to limit the Board membership to currently licensed Pharmacists, the bracketed language pertaining to a public member should be deleted, as should Section 203(b). In this event, the alternative “each” should be selected, and Section 203(a) should be renumbered as Section 203.

<sup>48</sup> Section 203(a) of the Act requires that a Pharmacist be engaged in the Practice of Pharmacy at the time of his or her appointment as a Board member and that he or she have at least five (5) years of experience in the Practice of Pharmacy in the state prior thereto. Since the Practice of Pharmacy is defined in Section 104 in broad terms, it renders a Pharmacist actively engaged in almost any phase of the practice eligible for appointment. This provides for candidates who have divergent backgrounds and experiences and who are knowledgeable in the affairs of the profession.

However, it should be noted from the definition of Pharmacy Practice in Section 104 that those persons actively engaged in the Practice of Pharmacy would basically be limited to those individuals who are working within settings where medications/Devices are Dispensed and Pharmacist Care Services is provided. To include persons who are in positions related to the practice but who are not engaged in Dispensing and Pharmacist Care Services functions would wrongfully cause the inclusion of individuals, such as personnel employed by Drug Manufacturers, Wholesale Distributors, and the like, who may be licensed to practice but who do not practice Pharmacy under the terms of the applicable Practice Act. The determination as to whether or not an individual is actively engaged in the Practice of Pharmacy will undoubtedly be rendered on a case-by-case basis. The general Criteria described above, however, would most probably be applicable in making the determination. Under the terms of a Practice Act, which includes a definition of Pharmacy practice, only those persons who are actively engaged in the basic functions set forth in the definition would be individuals “actively engaged in the Practice of Pharmacy.”

<sup>49</sup> Specific qualifying Criteria for the public member have been deliberately omitted from this section. Reliance has been placed in the Governor to determine what attributes an individual should possess in order to meaningfully serve on a Board of Pharmacy. In order to help ensure that such a member would be truly independent in his or her judgments, those persons who have a possible substantial relationship with the profession are rendered ineligible by this Section.

<sup>50</sup> The purpose of Section 204(b) is to provide a mechanism through which any interested Person or group may designate a candidate for the Board. Since nominations are recommendations only, the Governor retains complete discretion in regard to the appointees. As an alternative to appointment of Board of Pharmacy members by the Governor, some state laws call for the election of such members by the states’ Pharmacists.



## **Section 205. Terms of Office.**

- (a) Except as provided in subsection (b), members of the Board of Pharmacy shall be appointed for a term of \_\_\_\_\_ years, except that members of the Board who are appointed to fill vacancies that occur prior to the expiration of a former member's full term shall serve the unexpired portion of such term.
- (b) The terms of the members of the Board shall be staggered, so that the terms of no more than \_\_\_\_\_ member(s) shall expire in any year. Each member shall serve until a successor is appointed and qualified.
  - (1) The present members of the Board shall serve the balance of their terms.
  - (2) Any present Board member appointed initially for a term of less than \_\_\_\_\_ years shall be eligible to serve for \_\_\_\_\_ additional full terms.
- (c) No member of the Board shall serve more than \_\_\_\_\_ consecutive full terms. The completion of the unexpired portion of a full term shall not constitute a full term for purposes of this Section.

## **Section 206. Vacancies.**

Any vacancy that occurs in the membership of the Board for any reason, including expiration of term, removal, resignation, death, disability, or disqualification, shall be filled by the Governor in the manner prescribed by Section 204.

## **Section 207. Removal.<sup>51</sup>**

- (a) A Board member may be removed pursuant to the procedures set forth in subsection (b) herein, upon one or more of the following grounds:
  - (1) the refusal or inability for any reason of a Board member to perform his or her duties as a member of the Board in an efficient, responsible, and professional manner;
  - (2) the misuse of office by a member of the Board to obtain personal, pecuniary, or material gain or advantage for himself or herself or another through such office;
  - (3) the violation by any member of the laws governing the Practice of Pharmacy or the Distribution of Drugs and/or Devices.
- (b) Removal of a member of the Board of Pharmacy shall be in accordance with the Administrative Procedures Act of this State, or other applicable laws.

## **Section 208. Organization.**

- (a) The Board of Pharmacy shall elect from its members a President and such other officers as it deems appropriate and necessary to the conduct of its business. The President of the Board of Pharmacy shall preside at all meetings of the Board and shall be responsible for the performance of all of the duties and functions of the Board required or permitted by this Act. Each additional officer elected by the Board shall perform those duties normally associated with his or her position and such other duties assigned to him or her from time to time by the Board.
- (b) Officers elected by the Board shall serve terms of one (1) year commencing with the day of their election and ending upon election of their successors and shall serve no more than \_\_\_\_\_ consecutive full terms in each office to which they are elected.

---

<sup>51</sup> In certain jurisdictions, there may be general statutory provisions that establish the procedures and grounds for the removal of appointed public officials. In these jurisdictions, you may wish to disregard Section 207.

- (c) The Board shall employ a Pharmacist to serve as a full time employee of the Board in the position of Executive Director. The Executive Director shall be responsible for the performance of the administrative functions of the Board and such other duties as the Board may direct.<sup>52</sup>

### **Section 209. Compensation of Board Members.**

Each member of the Board of Pharmacy shall receive as compensation the sum of \$\_\_\_\_\_ per day for each day on which the member is engaged in performance of the official duties of the Board, and shall be reimbursed for all reasonable and necessary expenses incurred in connection with the discharge of such official duties.

### **Section 210. Meetings.**

- (a) The Board of Pharmacy shall meet at least once every \_\_\_\_\_ months to transact its business. The Board shall meet at such additional times as it may determine. Such additional meetings may be called by the President of the Board or by two-thirds (2/3) of the members of the Board.
- (b) The Board shall meet at such place as it may from time to time determine. The place for each meeting shall be determined prior to giving notice of such meeting and shall not be changed after such notice is given without adequate prior notice.
- (c) Notice of all meetings of the Board shall be given in the manner and pursuant to requirements prescribed by the State's Administrative Procedures Act.
- (d) A majority of the members of the Board shall constitute a quorum for the conduct of a Board meeting and, except where a greater number is required by this Act or by any rule of the Board, all actions of the Board shall be by a majority of a quorum.
- (e) All Board meetings and hearings shall be open to the public. The Board may, in its discretion and according to law, conduct any portion of its meeting in executive session, closed to the public.<sup>53</sup>

### **Section 211. Employees.**

The Board of Pharmacy may, at its discretion, employ persons, in addition to the Executive Director, in such other positions or capacities as it deems necessary to the proper conduct of Board business and to the fulfillment of the Board's responsibilities as defined by this Act.<sup>54</sup>

### **Section 212. Rules.**

The Board of Pharmacy shall make, adopt, amend, and repeal such rules as may be deemed necessary by the Board from time to time for the proper Administration and enforcement of this Act. Such rules shall be promulgated in accordance with the procedures specified in the Administrative Procedures Act of this State.

---

<sup>52</sup> NABP urges that every Board have a permanent administrative official, an Executive Director who is a currently licensed Pharmacist, to perform and supervise the administrative duties and functions for which the Board is responsible on a day-to-day basis. The responsibilities of the Executive Director should include the hiring of necessary staff to assist in fulfilling the responsibilities of the Board.

<sup>53</sup> Many states have adopted "sunshine" laws that provide for open meetings. Section 210(e) may not be necessary or may need revision to eliminate or to curtail the use of executive sessions.

<sup>54</sup> Inspectors employed by the Board of Pharmacy may be Pharmacists. Boards may wish to consider whether inspectors must be Pharmacists.

## Section 213. Powers and Responsibilities.

- (a) The Board of Pharmacy shall be responsible for the control and regulation of the Practice of Pharmacy in this State including, but not limited to, the following<sup>55</sup>:
- (1) the licensing by examination or by license transfer of applicants who are qualified to engage in the Practice of Pharmacy under the provisions of this Act;
  - (2) the renewal of licenses to engage in the Practice of Pharmacy;
  - (3) the establishment and enforcement of compliance with professional standards and rules of conduct of Pharmacists engaged in the Practice of Pharmacy;
  - (4) the determination and issuance of standards for recognition and approval of degree programs of schools and colleges of Pharmacy whose graduates shall be eligible for licensure in this State, and the specification and enforcement of requirements for practical training, including Pharmacy practice experience<sup>56</sup>;
  - (5) the enforcement of those provisions of this Act relating to the conduct or competence of Pharmacists practicing in this State; the Revocation, Summary Suspension, Suspension, Probation, Censure, or Reprimand of, or the issuance of Warnings or the assessment of Fines/Civil Penalties or Costs/Administrative Costs against licenses to engage in the Practice of Pharmacy; and the issuance of Cease and Desist orders against any Person or entity;
  - (6) the licensure and regulation of the training, qualifications, and employment of Pharmacy Interns, Certified Pharmacy Technicians, and Certified Pharmacy Technician Candidates;
  - (7) the collection of professional demographic data;
  - (8) the right to seize any such Drugs and Devices found by the Board to constitute an imminent danger to the public health and welfare;
  - (9) establishing minimum specifications for the physical facilities, technical equipment, environment, supplies, personnel, and procedures for the storage, Compounding and/or Dispensing of such Drugs or Devices, for the monitoring of Drug therapy, and for the Manufacture and Distribution of Drugs and Devices;
  - (10) establishing minimum standards for the purity and quality of such Drugs, Devices, and other materials within the Practice of Pharmacy;
  - (11) the issuance and renewal of licenses for Pharmacies located within this State, or outside this State if providing services to patients within this State, that Compound or Dispense Drugs or Devices or provide Pharmacist Care Services.
  - (12) the issuance and renewal of licenses of all Manufacturers and Distributors of Drugs and Devices located within this State, or outside this State if providing such services within this State;
  - (13) inspection at all reasonable hours of the facility and appropriate records of any licensed Person or licensed facility and any Person or facility seeking licensure for the purpose of determining if any provisions of the laws governing licensure, the legal Distribution of Drugs or Devices, or the Practice of Pharmacy are being violated, including the inspection of Protected Health Information. The Board

---

<sup>55</sup> The "Practice of Pharmacy in this State" includes shipping Prescription Drugs into this State from another jurisdiction. However, this is not meant to be construed as a licensure requirement for every Pharmacist that is employed at a Nonresident Pharmacy unless they are specifically engaged in the Practice of Pharmacy and provide services to residents in this State (see Sections 104 and 501(a) of this Act).

<sup>56</sup> Great care should be exercised by the Boards with respect to this Section. Many states have statutes or rules which provide that approved or accredited degree programs of schools or colleges of Pharmacy are those approved by the Accreditation Council for Pharmacy Education (ACPE).

It is a well-established rule of administrative law that any delegation of governmental power must carry with it appropriate limitations and procedural safeguards for affected individuals. Thus, a direct, unequivocal grant of the accreditation function to a private organization, such as ACPE, might be deemed an unauthorized, improper, and invalid delegation of Board or legislative authority. An NABP study of this question discovered at least one case where a court overturned a Board action based upon such invalid delegation to a private body. See *Garces v Department of Registration and Education*, 254 N.E.2d 622 (Ill, 1969).

NABP urges all Boards to adopt, in their Rules, the Standards of Accreditation established from time to time by the ACPE, the nationally recognized accrediting agency for Pharmacy degree programs.

of Pharmacy, its officers, inspectors, and representatives shall cooperate with all agencies charged with the enforcement of the laws of the United States, of this State, and of all other states relating to Drugs, Devices, and the Practice of Pharmacy;

(14) establishing minimum standards for maintaining the integrity and confidentiality of prescription information and other patient health care information; and<sup>57</sup>

(15) the approval of Pharmacy practice initiatives that improve the quality of or access to Pharmacist Care Services, but which fall outside the scope of present regulations. This subsection shall not be construed to expand the definition of the Practice of Pharmacy as defined in this Act.

(b) Centralized Performance Database

(1) The Board of Pharmacy shall utilize a Centralized Performance Database. The Centralized Performance Database shall be maintained in such a way as to permit an evaluator to apply Criteria and Standards to data from one pharmacy, and determine whether, over time, outcomes from that pharmacy compare favorably with outcomes from other pharmacies.

(2) The Board of Pharmacy shall conduct a Practice Accountability Audit at least once every six months to identify pharmacies that consistently violate Criteria and/or standards. The Board of Pharmacy shall require that pharmacies so identified provide an explanation of the reason for their consistent violation of Criteria and/or standards.

(c) The Board of Pharmacy shall have such other duties, powers, and authority as may be necessary to the enforcement of this Act and to the enforcement of Board rules made pursuant thereto, which shall include, but are not limited to, the following:

(1) The Board may join such professional organizations and associations organized exclusively to promote the improvement of the standards of the Practice of Pharmacy for the protection of the health and welfare of the public and/or whose activities assist and facilitate the work of the Board.

(2) The Board may receive and expend funds, in addition to its [annual/biennial] appropriation, from parties other than the State, provided:

(i) such funds are awarded for the pursuit of a specific objective which the Board is authorized to accomplish by this Act, or which the Board is qualified to accomplish by reason of its jurisdiction or professional expertise;

(ii) such funds are expended for the pursuit of the objective for which they are awarded;

(iii) activities connected with or occasioned by the expenditures of such funds do not interfere with the performance of the Board's duties and responsibilities, and do not conflict with the exercise of the Board's powers as specified by this Act;

(iv) such funds are kept in a separate, special account; and

(v) periodic reports are made concerning the Board's receipt and expenditure of such funds.

(3) The Board may establish a Bill of Rights for patients concerning the health care services a patient may expect in regard to Pharmacist Care Services.<sup>58</sup>

---

<sup>57</sup> Under this Act, "Protected Health Information" may be used or disclosed without acknowledgement, authorization, or opportunity to agree or object in the situations described in 45 CFR 164.512(a) – (l), and which include:

- As required by law
- For certain public health activities
- For certain health oversight activities
- Pursuant to judicial or administrative proceedings
- For law enforcement purposes
- For military or national security purposes
- As necessary to comply with worker compensation laws
- In situations presenting a serious threat to health or safety

Investigative activities of the Boards of Pharmacy are considered health oversight activities and, therefore, fall under this disclosure exemption.

<sup>58</sup> A Patient's Bill of Rights establishes the professional services that a patient may expect when obtaining Drugs or Devices from a Pharmacist. The Bill of Rights would normally contain patient expectations that could translate into standards of professional practice and/or codes of conduct for the Pharmacist. Accordingly, if a Board should choose to establish a Patient's Bill of Rights, the Bill should be consistent with standards of practice, codes of ethics, and

- (4) Any investigation, inquiry, or hearing which the State Board of Pharmacy is empowered to hold or undertake may be held or undertaken by or before any member or members of the Board and the finding or order of such member or members shall be deemed to be the order of said Board when approved and confirmed as noted in Section 210(d).
- (5) Embargo.<sup>59</sup>
- (i) Notwithstanding anything in this Act to the contrary, whenever a duly authorized representative of the Board finds, or has probable cause to believe, that any Drug or Device is adulterated or misbranded within the meaning of the (State) Food and Drug Act, he or she shall affix to such Drug or Device a tag or other appropriate marking giving notice that such article is or is suspected of being adulterated or misbranded, has been detained or embargoed, and Warning all Persons not to remove or dispose of such article by sale or otherwise until provision for removal or disposal is given by the Board, its agent, or the Court. No Person shall remove or dispose of such embargoed Drug or Device by sale or otherwise without the permission of the Board or its agent or, after summary proceedings have been instituted, without permission from the Court.
  - (ii) When a Drug or Device detained or embargoed under Paragraph (i) of this subsection (5) has been declared by such representative to be adulterated or misbranded, the Board shall, as soon as practical thereafter, petition the Judge of the \_\_\_\_\_ Court in which jurisdiction the article is detained or embargoed for an order for condemnation of such article. If the judge determines that the Drug or Device so detained or embargoed is not adulterated or misbranded, the Board shall direct the immediate removal of the tag or other marking.
  - (iii) If the Court finds the detained or embargoed Drug or Device is adulterated or misbranded, such Drug or Device, after entry of the decree, shall be destroyed at the expense of the owner under the supervision of a Board representative and all court costs and fees, storage, and other proper expense shall be borne by the owner of such Drug or Device. When the adulteration or misbranding can be corrected by proper Labeling or processing of the Drug or Device, the Court, after entry of the decree and after such costs, fees, and expenses have been paid and a good and sufficient bond has been posted, may direct that such Drug or Device be Delivered to the owner thereof for such Labeling or processing under the supervision of a Board representative. Expense of such supervision shall be paid by the owner. Such bond shall be returned to the owner of the Drug or Device on representation to the Court by the Board that the Drug or Device is no longer in violation of the embargo and the expense of supervision has been paid.
  - (iv) It is the duty of the Attorney General [State's Attorney] to whom the Board reports any violation of Section 213(c)(5) to cause appropriate proceedings to be instituted in the proper court without delay and to be prosecuted in the manner required by law. Nothing in this subparagraph (iv) shall be construed to require the Board to report violations whenever the Board believes the public's interest will be adequately served in the circumstances by a suitable written notice or Warning.
- (6) The Board may place under seal all Drugs or Devices that are owned by or in the possession, custody, or control of a licensee at the time his or her license is Suspended or Revoked or at the time the Board refuses to renew his license. Except as otherwise provided in this section, Drugs or Devices so sealed shall not be disposed of until appeal rights under the Administrative Procedures Act have expired, or an appeal filed pursuant to that Act has been determined. The court involved in an appeal filed pursuant to the Administrative Procedures Act may order the Board, during the

---

regulations that the Board has adopted under the Pharmacy Practice Act. If care is not taken, a Board could inadvertently expand the role and the responsibilities of the Pharmacist through the establishment of a Patient's Bill of Rights.

<sup>59</sup> The purpose of this subsection is to ensure quality, purity, and correct Labeling of Drugs, Devices, and other materials.

pendency of the appeal, to sell sealed Drugs that are perishable. The proceeds of such a sale shall be deposited with that court.

- (7) Except as otherwise provided to the contrary, the Board shall exercise all of its duties, powers, and authority in accordance with the State Administrative Procedures Act.
- (8) In addition to the fees specifically provided for herein, the Board may assess additional reasonable fees for services rendered to carry out its duties and responsibilities as required or authorized by this Act or Rules adopted hereunder. Such services rendered shall include, but not be limited to, the following:
  - (i) issuance of duplicate certificates or identification cards;
  - (ii) mailing lists or reports of data maintained by the Board;
  - (iii) copies of any documents;
  - (iv) certification of documents;
  - (v) notices of meetings;
  - (vi) licensure transfer;
  - (vii) examination Administration to a licensure applicant; and
  - (viii) examination materials.
- (9) Cost Recovery.<sup>60</sup>
  - (i) If any order issues in resolution of a disciplinary proceeding before the Board of Pharmacy, the Board may request the \_\_\_\_\_ to direct any licensee found guilty of a charge involving a violation of any Drug laws or rules, to pay to the Board a sum not to exceed the reasonable costs of the investigation and prosecution of the case and, in any case, not to exceed twenty-five thousand dollars (\$25,000).
  - (ii) In the case of a Pharmacy or Wholesale Distributor, the order may be made as to the corporate owner, if any, and as to any Pharmacist, officer, owner, or partner of the Pharmacy or Wholesale Distributor who is found to have had knowledge of or have knowingly participated in one or more of the violations set forth in this section.
  - (iii) The costs to be assessed shall be fixed by the \_\_\_\_\_ and shall not be increased by the Board; where the Board does not adopt a proposed decision and remands the case to a(n) \_\_\_\_\_, the \_\_\_\_\_ shall not increase any assessed costs.
  - (iv) Where an order for recovery of costs is made and timely payment is not made as directed in the Board's decision, the Board may enforce the order for payment in the Court in the county where the administrative hearing was held. This right of enforcement shall be in addition to any other rights the Board may have as to any Person directed to pay costs.
  - (v) In any action for recovery of costs, proof of the Board's decision shall be conclusive proof of the validity of the order of payment and the terms for payment.

---

<sup>60</sup> The "\_\_\_\_\_" interspersed throughout this section may be filled with the terms: "administrative law judge," "hearing officer," or "presiding officer," as determined by individual states.

## Article III Licensing

### Introductory Comment to Article III

*Article III of the Model Act specifies the requirements for initial licensure of Pharmacists, transfer of licensure, and renewal of licenses and registrations. In each of these areas, the Act sets forth basic Criteria and delegates to the Board the authority for implementing those Criteria. The Board does this by utilizing appropriate administrative enforcement mechanisms and by the issuance of specific rules.*

*Section 301 establishes the basis for this Article by making it unlawful for any unlicensed Person to engage in the Practice of Pharmacy, and by enabling the Board to exact penalties for unlawful practice.*

*In the area of initial licensure (Section 302), the Board must implement the Act by approving degree programs of Pharmacy, by specifying the examination to be employed (Section 302[b]), by establishing Pharmacy practice experience standards (Section 302[c]), and by ensuring that all other prerequisites are met by each applicant to whom it issues a license.*

*The Act also reflects the efforts of NABP to continue uniform standards for transfer of licensure (Section 303).*

### Section 301. Unlawful Practice.

- (a) Except as otherwise provided in this Act, it shall be unlawful for any individual, whether located in or outside this State, to engage in the Practice of Pharmacy in this State unless currently licensed to practice under any facet of the provisions of this Act.
- (b) The provision of Pharmacist Care Services to an individual in this State, through the use of Telepharmacy Technologies, regardless of the location of the Pharmacist, shall constitute the Practice of Pharmacy and shall be subject to regulation.<sup>61</sup>
  - (1) Licensed Pharmacies located outside this State that provide Pharmacist Care Services to individuals in this State must be licensed within this State under Article V of this Act.
  - (2) Pharmacists located outside this State who are providing Pharmacist Care Services outside of a licensed Pharmacy to individuals located in this State must register with this State to engage in the nonresident Practice of Pharmacy.
- (c) Licensed Practitioners authorized under the laws of this State to Compound Drugs and to Dispense Drugs to their patients in the practice of their respective professions shall meet the same standards, record-keeping requirements, and all other requirements for the Dispensing of Drugs applicable to Pharmacists.<sup>62</sup>

---

<sup>61</sup> NABP recognizes that protection of the public health should extend across State borders. Accordingly, the NABP *Model Act* incorporates the Practice of Telepharmacy within the scope of the "Practice of Pharmacy" and requires an independently practicing Pharmacist located outside this State to obtain full licensure for providing Pharmacist Care Services from outside the State to patients within the State.

<sup>62</sup> Boards of Pharmacy are often confronted with the problem of preventing unlicensed individuals from engaging in one or more facets of the Practice of Pharmacy. The regulation of the Practice of Pharmacy, including the control of unlicensed practice in the profession, has a reasonable and rational relationship to public health, safety, and welfare. See, for example, *State v Wakeen*, 57 N.W.2d 364 (Wis, 1953). cf. *State v VanKeegan*, 113 A.2d 141 (Conn, 1955) and *Williamson v Lee Optical of Oklahoma*, 348 U.S. 483 (1955), concerning prohibitions on the unlicensed practice of ophthalmology. For this reason, vesting the power in the Board to regulate the illicit practice would not appear to violate the constitutional due process requirements. Monetary fines are another enforcement action Boards can utilize to protect the public health. Although monetary fines are not generally considered criminal sanctions, it can be forcibly argued that there are no constitutional barriers impeding the imposition of fines by a Board of Pharmacy. See, for example, *Helvering v Mitchell*, 303 U.S. 376 (1938); *City of Waukegan v Pollution Control Board*, 311 N.E.2d 146 (Ill, 1974); *County Council for Montgomery County v Investors Funding Corp*, 312 A.2d 225 (Md, 1973); and *Rody v Hollis*, 500 P.2d 97 (Wash, 1972).

One area that could present a serious question of law in regard to Section 301(b), however, involves the constitutional limitation on the delegation of authority to administrative agencies. It is likely that the delegation contemplated in Article III of the *Model Act* would be valid in a majority of jurisdictions. See, for example, *Jordan v Board of Insurance*, 334 S.W. 2d 278 (Tex, 1960); *Sutherland v Ferguson*, 397 P. 2d 335 (Kan, 1964); and *Kovack v Licensing Board of City of Waterville*, 173 A. 2d 554 (Me, 1961); see generally L. Davis, *Administrative Law Treatise*, Section 2.10 (1970 Suppl.). Be cautioned, however, that certain

- (d) It shall be unlawful for any individual to perform the activities of a Certified Pharmacy Technician or Certified Pharmacy Technician Candidate unless currently licensed to do so under the provisions of this Act.
- (e)
  - (1) The Board may in its own name issue a Cease and Desist order to stop an individual from engaging in an unauthorized Practice of Pharmacy.
  - (2) Except as otherwise indicated in this Act, any individual who, after due process, shall be found by the Board to have unlawfully engaged in the Practice of Pharmacy shall be subject to a Fine to be imposed by the Board not to exceed \$\_\_\_\_\_ for each offense. Each such violation of this Act or the rules promulgated hereunder pertaining to unlawfully engaging in the Practice of Pharmacy shall also constitute a misdemeanor punishable upon conviction as provided in the criminal code of this State.
  - (3) Except as otherwise indicated in this Act, any individual who, after due process, shall be found by the Board to have unlawfully engaged in the Practice of Pharmacy that resulted in harm to an individual shall be subject to a Fine to be imposed by the Board not to exceed \$\_\_\_\_\_ for each offense. Each such violation of this Act or the rules promulgated hereunder pertaining to unlawfully engaging in the Practice of Pharmacy that resulted in harm to an individual shall also constitute a felony punishable upon conviction as provided in the criminal code of this State.

### **Section 302. Qualifications for Licensure by Examination.**

- (a) To obtain a license to engage in the Practice of Pharmacy, an applicant for licensure by examination shall:
  - (1) have submitted an application in the form prescribed by the Board of Pharmacy;
  - (2) have attained the age of majority;
  - (3) have graduated and received the first professional degree from a college or school of Pharmacy that has been approved by the Board of Pharmacy;<sup>63</sup>
  - (4) have graduated from a foreign college of Pharmacy, completed a transcript verification program, taken and passed a college of Pharmacy equivalency examination program, and completed a process of communication-ability testing as defined under Board of Pharmacy regulations so that it is ensured that the applicant meets standards necessary to protect public health and safety;<sup>64</sup>
  - (5) have completed a Pharmacy practice experience program or other program that has been approved by the Board of Pharmacy, or demonstrated to the Board's satisfaction that experience in the Practice of Pharmacy which meets or exceeds the minimum Pharmacy practice experience requirements of the Board;
  - (6) have successfully passed an examination or examinations approved by the Board of Pharmacy;
  - (7) have undergone a state and federal fingerprint-based criminal background check as specified by Board rule; and
  - (8) have paid the fees specified by the Board of Pharmacy for the examination and any related materials, and have paid for the issuance of the license.
- (b) Examinations.
  - (1) The examinations for licensure, which include a pharmacy practice examination and a jurisprudence examination, required under Section 302(a)(7) of the Act, shall be administered by an NABP

---

jurisdictions require very specific standards in a delegation of authority that could render Section 301(b) constitutionally suspect. See, for example, *People v Tibbits*, 305 N.E. 2d 152 (Ill, 1973); *Sarasota County v Barg*, 302 So. 2d 737 (Fla, 1974). In these jurisdictions, revisions of Article III may be necessary.

<sup>63</sup> It is contemplated that Boards will approve those programs whose standards are at least equivalent to the standards required by the ACPE. This would include college-structured pharmacy practice experience programs and continuing education programs. See the footnote to Section 213(a)(4) above for further discussion of the Board's proper role in the accreditation process.

<sup>64</sup> Boards should avoid mention of specific examinations or other contracted services in their practice act and regulations to avoid challenges related to an unconstitutional delegation of authority. It is contemplated that Boards will utilize the Foreign Pharmacy Graduate Equivalency Examination® (FPGEE®) as part of their assessment of pharmacy education equivalence.



contracted testing provider. If applicable, state-specific compounding examinations shall be administered by the Board. NABP will determine the content and subject matter of the pharmacy practice examination and the Board shall determine the content and subject matter of each state-specific compounding and jurisprudence examination.

- (2) The examinations shall be prepared to measure the competence of the applicant to engage in the Practice of Pharmacy. NABP may employ, cooperate, and contract with any organization or consultant in the preparation and grading of an examination, but the Board shall retain the sole discretion and responsibility for determining which applicants are eligible for licensure.

(c) Pharmacy Practice Experience Programs and Other Training Programs.<sup>65</sup>

- (1) All applicants for licensure by examination shall obtain practical experience in the Practice of Pharmacy concurrent with or after college attendance, or both, under such terms and conditions as the Board shall determine.<sup>66</sup>
- (2) The Board shall establish such licensure requirements for Pharmacy Interns and standards for Pharmacy practice experiences, or any other experiential program necessary to qualify an applicant for the licensure examination, and shall also determine the qualifications of Preceptors used in practical experience programs.<sup>67</sup>

### Section 303. Qualifications for Licensure Transfer.<sup>68</sup>

- (a) In order for a Pharmacist currently licensed in another jurisdiction to obtain a license as a Pharmacist by license transfer in this State, an applicant shall:<sup>69</sup>
  - (1) have submitted an application in the form prescribed by the Board of Pharmacy;
  - (2) have attained the age of majority;
  - (3) have possessed at the time of initial licensure as a Pharmacist all qualifications necessary to have been eligible for licensure at that time in this State;
  - (4) have engaged in the Practice of Pharmacy for a period of at least one (1) year or have met the Pharmacy practice experience requirements of this State within the one (1) year period immediately previous to the date of such application;

---

<sup>65</sup> As college-based Pharmacy practice experience programs become uniform under the most recent revision of the ACPE *Standards and Guidelines for the Professional Program in Pharmacy Leading to the Doctor of Pharmacy Degree* (effective July 1, 2007), and when boards of pharmacy are convinced that schools and colleges of pharmacy are meeting these Accreditation Standards and Guidelines and the competency requirements set out by Boards, Boards should begin to broadly accept and recognize college-based Pharmacy practice experience programs completed by students in other jurisdictions and eliminate requirements that such students obtain additional Pharmacy practice experience hours in addition to those obtained as part of the college of pharmacy curriculum.

Because of the potential lack of uniformity among non-college-based Pharmacy practice experience programs, it is recommended that Boards exercise their prerogative to accept only at their discretion non-college based Pharmacy practice experiences completed by Pharmacy Interns in other jurisdictions.

<sup>66</sup> Although Boards of Pharmacy mandate a specified number of hours of Pharmacy practice experiences as a prerequisite to licensure, Boards of Pharmacy are also encouraged to deem those requirements met if Boards find that the college-based Pharmacy practice experiences meet or exceed the hourly Pharmacy practice experience requirements.

As indicated in the Model Rules for Pharmacy Interns, applicants for licensure as Pharmacists shall submit evidence that they have satisfactorily completed: (1) an objective assessment mechanism intended to evaluate achievement of desired competencies as delineated in the ACPE Accreditation Standards and Guidelines and (2) not less than 1,740 hours of Pharmacy practice experience credit under the instruction and supervision of a Preceptor. Boards may consider moving away from requiring a specific number of contact hours should it be determined that the ACPE Accreditation Standards and Guidelines result in appropriate preparation for students and objective assessment mechanisms demonstrate such.

<sup>67</sup> Boards of Pharmacy are strongly encouraged to utilize the ACPE Accreditation Standards and Guidelines as a basis for establishment and revision of Board standards for Pharmacy practice experiences. These Accreditation Standards and Guidelines also contain additional guidance on the desired behaviors, qualities, and values of preceptors.

<sup>68</sup> See the NABP Model Rules for Public Health Emergencies for language that addresses the temporary recognition of nonresident pharmacist licensure in the case of a declared State of Emergency issued due to a Public Health Emergency.

<sup>69</sup> It is intended that NABP's National Disciplinary Clearinghouse would be utilized by state Boards for verifying information provided by applicants.

- (5) have presented to the Board proof of initial licensure by examination and proof that such license is in good standing;
  - (6) have presented to the Board proof that any other license granted to the applicant by any other state has not been Suspended, Revoked, or otherwise restricted for any reason, except nonrenewal or for the failure to obtain the required continuing education credits, in any state where the applicant is currently licensed but not engaged in the Practice of Pharmacy; and
  - (7) have paid the fees specified by the Board.
- (b) No applicant shall be eligible for license transfer unless the state in which the applicant was initially licensed as a Pharmacist also grants licensure transfer to Pharmacists duly licensed by examination in this State, under like circumstances and conditions.<sup>70</sup>

#### **Section 304. Renewal of Licenses.**

- (a) Each Pharmacist, Pharmacy Intern, and Certified Pharmacy Technician shall apply for renewal of his or her license annually [or at such interval determined by the Board], no later than the first day of \_\_\_\_\_. A Pharmacist or Pharmacy Intern who desires to continue in the Practice of Pharmacy in this State shall file with the Board an application in such form and containing such data as the Board may require for renewal of the license. If the Board finds that the applicant has been licensed, and that such license has not been Revoked or placed under Suspension, that the applicant has attested that he or she has no criminal convictions or arrests, has paid the renewal fee, has continued his or her Pharmacy education in accordance with the rules of the Board, and is entitled to continue in the Practice of Pharmacy, the Board shall issue a license to the applicant.
- (b) If a Pharmacist fails to make application to the State Board of Pharmacy for renewal of his or her license within a period of three years from the expiration of his or her license, he or she must pass an examination for license renewal; except that a Person who has been licensed under the laws of this State and after the expiration of his or her license, has continually practiced Pharmacy in another State under a license issued by the authority of such State, may renew his or her license upon payment of the designated fee.
- (c) Certified Pharmacy Technician Candidates must complete requirements for Certified Pharmacy Technician licensure within 12 months. For good cause shown, the Board may approve one 12-month extension.

#### **Section 305. Continuing Pharmacy Education.**

The Board shall, by rule, establish requirements for continuing education in Pharmacy, including the determination of acceptable program content and fees. The Board shall adopt rules necessary to carry out the stated objectives and purposes, to enforce the provisions of this Section, and to ensure continued competence.

#### **Section 306. Pharmacy Practice Experience Program Standards; Pharmacy Intern Licensure.**

The Board of Pharmacy shall establish standards for Pharmacy practice experience programs for the purpose of providing the practice experience necessary for licensure as a Pharmacist. The Board shall grant a Pharmacy Intern license to Pharmacy students, authorizing those students to engage in the Practice of Pharmacy under the

---

<sup>70</sup> Endorsement states may wish to consider the removal of Subparagraph (b) in this Section.

supervision of a Pharmacist. The Board of Pharmacy shall adopt rules regarding the licensure of Pharmacy Interns and the standards for Pharmacy practice experience programs.<sup>71</sup>

### **Section 307. Licensure of Certified Pharmacy Technicians.**

- (a) In order to be licensed as a Certified Pharmacy Technician in this State, an applicant shall:
  - (1) have submitted an application in the form prescribed by the Board of Pharmacy;
  - (2) have attained the age of \_\_\_\_\_;
  - (3) have graduated from high school or obtained a Certificate of General Educational Development (GED) or equivalent;
  - (4) have:<sup>72</sup>
    - (i) graduated from a site-specific training program or a competency-based pharmacy technician education and training program that includes experiential training approved by the Board of Pharmacy;<sup>73</sup>
    - (ii) completed a minimum number of pharmacy technician practice experience hours approved by the Board of Pharmacy;<sup>74</sup>
  - (5) have successfully passed an examination developed using nationally recognized and validated psychometric and pharmacy practice standards approved by the Board of Pharmacy;
  - (6) have undergone a state and federal fingerprint-based criminal background check as specified by Board rule; and
  - (7) have paid the fees specified by the Board of Pharmacy for the examination and any related materials, and have paid for the issuance of the license.
- (b) No Pharmacist whose license has been denied, Revoked, Suspended, or restricted for disciplinary purposes shall be eligible to be licensed as a Certified Pharmacy Technician.<sup>75</sup>
- (c) The Board of Pharmacy shall, by rule, establish requirements for licensure of Certified Pharmacy Technicians.

### **Section 308. Licensure of Certified Pharmacy Technician Candidates.**

- (a) In order to be licensed as a Certified Pharmacy Technician Candidate in this State, an applicant shall:
  - (1) have submitted an application in the form prescribed by the Board of Pharmacy;
  - (2) have attained the age of \_\_\_\_\_;
  - (3) have undergone a state and federal fingerprint-based criminal background check as specified by Board rule;
  - (4) have paid the fees specified by the Board; and

---

<sup>71</sup> Boards of Pharmacy are strongly encouraged to utilize the ACPE Accreditation Standards and Guidelines as a basis for establishment and revision of board standards for Pharmacy practice experiences. These Accreditation Standards and Guidelines also contain additional guidance on the desired behaviors, qualities, and values of preceptors.

<sup>72</sup> Boards should avoid mention of specific examinations or other contracted services in their practice act and regulations to avoid challenges related to an unconstitutional delegation of authority. It is contemplated that Boards will utilize the Certified Pharmacy Technician Candidate Certification Board examination as part of their assessment of technician competence to assist in the practice of pharmacy.

<sup>73</sup> It is recommended that states adopt this requirement, if not currently required, through a process that incorporates provisions for grandfathering.

<sup>74</sup> It is contemplated that Boards will approve those Certified Pharmacy Technician Candidate training programs whose standards are at least equivalent to the minimum standards developed by an accrediting organization recognized by state Boards, such as ACPE and ASHP. See the footnote to Section 213(a)(4) above for further discussion of the Board's proper role in the accreditation process.

<sup>75</sup> The Board may specifically authorize a Pharmacist whose license has been disciplined to register as a Certified Pharmacy Technician or Certified Pharmacy Technician Candidate under terms and conditions deemed appropriate.

- (5) have enrolled in a site-specific training program or a competency-based pharmacy technician education and training program that includes experiential training approved by the Board of Pharmacy and an objective assessment mechanism prepared in accordance with any rules established by the Board.
- (b) No Pharmacist whose license has been denied, Revoked, Suspended, or restricted for disciplinary purposes shall be eligible to be licensed as a Certified Pharmacy Technician Candidates.<sup>76</sup>
- (c) The Board of Pharmacy shall, by rule, establish requirements for licensure of Certified Pharmacy Technician Candidates.

---

<sup>76</sup> The Board may specifically authorize a Pharmacist whose license has been disciplined to register as a Certified Pharmacy Technician or Certified Pharmacy Technician Candidate under terms and conditions deemed appropriate.

## Article IV Discipline

### Introductory Comment to Article IV

*At the very heart of any Pharmacy Act is the enforcement power of the Board of Pharmacy. The Board must have authority to discipline and/or prohibit Pharmacies, Pharmacists, Pharmacy Interns, Certified Pharmacy Technicians, or Certified Pharmacy Technician Candidates who violate this Act or Rules from continuing to threaten the public if it is to fulfill its responsibilities. The Board must have the ability to stop wrongdoers, either permanently or temporarily, discipline them, and, where appropriate, to guide and assist errant licensees in rehabilitating themselves.*

*The Model Act disciplinary provisions are contained in Article IV. They were drafted with the purpose of granting to the Board the widest possible scope within which to perform its disciplinary functions. Standardized disciplinary action terms and definitions were developed to facilitate the accurate reporting of disciplinary actions taken by Boards of Pharmacy and to avoid confusion associated with state-to-state variations in terms and definitions. The grounds for disciplinary action were developed to ensure protection of the public, while reserving to the Board the power to expand upon them and adapt them to changing or local conditions as necessary. The penalties permitted under the Model Act will afford the Board the flexibility to conform and relate discipline to offenses.*

### Section 401. Disciplinary Action Terms.

The following is a list of disciplinary actions that may be taken, issued, or assessed by the Board of Pharmacy: Revocation, Summary Suspension, Suspension, Probation, Censure, Reprimand, Warning, Cease and Desist, Fine/Civil Penalty, Costs/Administrative Costs.<sup>77</sup>

### Section 402. Grounds, Penalties, and Reinstatement.<sup>78</sup>

- (a) The Board of Pharmacy may refuse to issue or renew, or may Revoke, Summarily Suspend, Suspend, place on Probation, Censure, Reprimand, issue a Warning against, or issue a Cease and Desist order against, the licenses or the registration of, or assess a Fine/Civil Penalty or Costs/Administrative Costs against any Person Pursuant to the procedures set forth in Section 403 herein below, upon one or more of the following grounds:
- (1) unprofessional conduct as that term is defined by the rules of the Board;<sup>79</sup>
  - (2) incapacity that prevents a licensee from engaging in the Practice of Pharmacy or a registrant from assisting in the Practice of Pharmacy, with reasonable skill, competence, and safety to the public;<sup>80</sup>

---

<sup>77</sup> Guidelines for the imposition of sanctions for certain designated offenses can be found in Appendix C: Guidelines for Disciplinary Sanctions of the *Model Act*.

<sup>78</sup> The penalties provided in Section 402 give the Board wide latitude to make the disciplinary action fit the offense. The “reasonable intervals” in 402(c) would be determined by the Board.

<sup>79</sup> It is particularly important to emphasize the need for specificity in defining the grounds upon which a Pharmacist’s or Pharmacy Intern’s license to practice Pharmacy, or a Certified Pharmacy Technician’s or Certified Pharmacy Technician Candidate’s registration to assist in the Practice of Pharmacy, may be Revoked or Suspended. The term “unprofessional conduct” is particularly susceptible to judicial challenge for being unconstitutionally vague. Each offense included within the meaning of this term must be capable of being understood with reasonable precision by the Persons regulated so that it can be readily enforced and relied upon during disciplinary proceedings, and so that those regulated by it may easily conform their professional conduct to its meaning(s).

These potential problems make it essential for Boards to issue appropriate rules making the grounds for disciplinary action specific, understandable, and reasonable. In addition, the Boards must ensure that such rules are published for the benefit of all licensees within their jurisdiction. Only by doing so can Boards be assured of authority to take successful and meaningful disciplinary actions that will not later be overturned by the courts.

This section must be examined in light of other state laws since some states, for example, restrict the circumstances under which a license may be denied to an individual because of the commission of a felony. In addition, an individual who has been convicted of a felony and who has paid his debt to society has restored constitutional protections that may curtail a strict application of Section 402(a)(3).

<sup>80</sup> Boards need to consider the issue of impairment if a registrant or licensee tests positive for a substance of misuse and/or abuse.

- (3) being guilty of one (1) or more of the following:
  - (i) a felony; or
  - (ii) violations of the Pharmacy or Drug laws of this State or rules and regulations pertaining thereto; or of laws, rules, and regulations of any other state; or of the Federal government;<sup>81</sup>
- (4) disciplinary action taken by another state or jurisdiction against a license or other authorization to Practice Pharmacy based upon conduct by the licensee similar to conduct that would constitute grounds for actions as defined in this section, which involves or may result in direct patient impact or harm in states other than that of the initiating Board;
- (5) failure to report to the Board any adverse action taken by another licensing jurisdiction (United States or foreign), government agency, law enforcement agency, or court for conduct that would constitute grounds for action as defined in this section;
- (6) failure to report to the Board one's surrender of a license or authorization to Practice Pharmacy in another state or jurisdiction while under disciplinary investigation by any of those authorities or bodies for conduct that would constitute grounds for action as defined in this section;
- (7) failure to report to the Board any adverse judgment, settlement, or award arising from a malpractice claim arising related to conduct that would constitute grounds for action as defined in this section;
- (8) knowing or suspecting that a Pharmacist or Pharmacy Intern is incapable of engaging in the Practice of Pharmacy or that a Certified Pharmacy Technician or Certified Pharmacy Technician Candidate is incapable of assisting in the Practice of Pharmacy, with reasonable skill, competence, and safety to the public, and failing to report any relevant information to the Board of Pharmacy;
- (9) misrepresentation of a material fact by a licensee in securing the issuance or renewal of a license or registration;
- (10) fraud by a licensee in connection with the Practice of Pharmacy;
- (11) affiliating with websites that may deceive or defraud patients or that violate Pharmacy or Drug laws of this State or rules and regulations pertaining thereto; or of laws, rules, and regulations of any other state; or of the Federal government;
- (12) engaging, or aiding and abetting an individual to engage in the Practice of Pharmacy without a license; assisting in the Practice of Pharmacy or aiding and abetting an individual to assist in the Practice of Pharmacy without being licensed by the Board of Pharmacy; or falsely using the title of Pharmacist, Pharmacy Intern, Certified Pharmacy Technician, or Certified Pharmacy Technician Candidate;
- (13) requiring Pharmacy personnel to meet production and/or performance metrics and/or quotas that negatively impact patient safety.<sup>82</sup>
- (14) failing to pay the costs assessed in a disciplinary hearing pursuant to Section 213(c)(9);
- (15) engaging in any conduct that subverts or attempts to subvert any licensing examination or the Administration of any licensing examination;<sup>83</sup>

---

<sup>81</sup> It is contemplated that Boards of Pharmacy will consider state and federal law, including any discrepancies between state and federal law, when evaluating complaints against a registrant or licensee related to a positive result on a cannabinoid Drug test. It is also contemplated that any complaint of this nature will be assessed on a case-by-case basis.

<sup>82</sup> This is not intended to include performance metrics that may be related to the ability and competency of Pharmacy personnel.

<sup>83</sup> It is recommended that the following rule be adopted defining subversion or the attempt to subvert any licensing examination.

- (a) Conduct which subverts or attempts to subvert any licensing examination or the administration of any examination shall include, but not be limited to, the following:
  - (1) Conduct which violates the security of the examination materials; removing from the examination room any examination materials without authorization; the unauthorized reproduction by any means of any portion of the actual licensing examination; aiding by any means the unauthorized reproduction of any portion of the actual licensing examination; paying or using professional or paid examination takers for the purpose of reconstructing any portion of the licensing examination; obtaining examination questions or other examination

- (16) being found by the Board to be in violation of any of the provisions of this Act or rules adopted pursuant to this Act;
- (17) illegal use or disclosure of Protected Health Information;
- (18) failure to furnish to the Board, its investigators, or representatives any information legally requested by the Board.

(b)

- (1) The Board may defer action with regard to an impaired licensee who voluntarily signs an agreement, in a form satisfactory to the Board, agreeing not to practice Pharmacy and to enter an approved treatment and monitoring program in accordance with this Section, provided that this Section should not apply to a licensee who has been convicted of, pleads guilty to, or enters a plea of nolo contendere to a felonious act prohibited by \_\_\_\_\_ or a conviction relating to a controlled substance in a court of law of the United States or any other state, territory, or country. A licensee who is physically or mentally impaired due to addiction to Drugs or alcohol may qualify as an impaired Pharmacist and have disciplinary action deferred and ultimately waived only if the Board is satisfied that such action will not endanger the public and the licensee enters into an agreement with the Board for a treatment and monitoring plan approved by the Board, progresses satisfactorily in such treatment and monitoring program, complies with all terms of the agreement and all other applicable terms of subsection (b)(2). Failure to enter such agreement or to comply with the terms and make satisfactory progress in the treatment and monitoring program shall disqualify the licensee from the provisions of this Section and the Board shall activate an immediate investigation and disciplinary proceedings. Upon completion of the rehabilitation program in accordance with the agreement signed by the Board, the licensee may apply for permission to resume the Practice of Pharmacy upon such conditions as the Board determines necessary.
- (2) The Board may require a licensee to enter into an agreement that includes, but is not limited to, the following provisions:
  - (i) Licensee agrees that his or her license shall be Suspended or Revoked indefinitely under subsection (b)(1).
  - (ii) Licensee will enroll in a treatment and monitoring program approved by the Board.
  - (iii) Licensee agrees that failure to satisfactorily progress in such treatment and monitoring program shall be reported to the Board by the treating professional, who shall be immune from any liability for such reporting made in good faith.
  - (iv) Licensee consents to the treating physician or professional of the approved treatment and monitoring program reporting to the Board on the progress of licensee at such intervals as the Board deems necessary and such Person making such report will not be liable when such reports are made in good faith.
- (3) The ability of an impaired Pharmacist to practice shall only be restored and charges dismissed when the Board is satisfied by the reports it has received from the approved treatment program that licensee can resume practice without danger to the public.
- (4) Licensee consents, in accordance with applicable law, to the release of any treatment information from anyone within the approved treatment program.
- (5) The impaired licensee who has enrolled in an approved treatment and monitoring program and entered into an agreement with the Board in accordance with subsection (b)(1) hereof shall have his

---

materials, except by specific authorization either before, during, or after an examination; or selling, Distributing, buying, receiving, or having unauthorized possession of any portion of a future, current, or previously administered licensing examination.

- (2) Unauthorized communication of examination information with any other examinee during the administration of a licensing examination; copying answers from another examinee or permitting one's answers to be copied by another examinee; having in one's possession during the administration of the licensing examination any books, equipment, notes, written or printed materials, or data of any kind other than the examination materials Distributed, or otherwise authorized to be in one's possession during the examination; or impersonating any examinee or having an impersonator take the licensing examination on one's behalf.

license Suspended or Revoked, but enforcement of this Suspension or Revocation shall be stayed by the length of time the licensee remains in the program and makes satisfactory progress, and complies with the terms of the agreement and adheres to any limitations on his practice imposed by the Board to protect the public. Failure to enter into such agreement or to comply with the terms and make satisfactory progress in the treatment and monitoring program shall disqualify the licensee from the provisions of this Section and the Board shall activate an immediate investigation and disciplinary proceedings.

- (6) Any Pharmacist who has substantial evidence that a licensee has an active addictive disease for which the licensee is not receiving treatment under a program approved by the Board pursuant to an agreement entered into under this Section, is diverting a controlled substance, or is mentally or physically incompetent to carry out the duties of his or her license, shall make or cause to be made a report to the Board. Any Person who reports pursuant to this Section in good faith and without malice shall be immune from any civil or criminal liability arising from such reports. Failure to provide such a report within a reasonable time from receipt of knowledge may be considered grounds for disciplinary action against the licensee so failing to report.
- (c) Any Person whose license to practice Pharmacy in this State has been Revoked, Summarily Suspended, Suspended, placed on Probation, Censured, Reprimanded, issued a Warning against, or issued a Cease and Desist order against, the licenses or the registration of, or assessed a Fine/Civil Penalty or Costs/Administrative Costs against pursuant to this Act, whether voluntarily or by action of the Board, shall have the right, at reasonable intervals, to petition the Board for reinstatement of such license.<sup>84</sup> Such petition shall be made in writing and in the form prescribed by the Board. Upon investigation and hearing, the Board may, at its discretion, grant or deny such petition, or it may modify its original finding to reflect any circumstances that have changed sufficiently to warrant such modifications. The Board, also at its discretion, may require such Person to pass an examination(s) for reentry into the Practice of Pharmacy.
- (d) Nothing herein shall be construed as barring criminal prosecutions for violations of this Act.
- (e) All final decisions by the Board shall be subject to judicial review pursuant to the Administrative Procedures Act.
- (f) Any individual or entity whose license to practice Pharmacy, or registration to assist in the Practice of Pharmacy, is Revoked, Suspended, or not renewed shall return his or her license or registration certificate to the offices of the State Board of Pharmacy within 10 days after receipt of notice of such action.

### **Section 403. Procedure.<sup>85</sup>**

- (a) Notwithstanding any provisions of the State Administrative Procedures Act, the Board may, without a hearing, Summarily Suspend a license for not more than 60 days if the Board finds that a Pharmacist, Pharmacy Intern, Certified Pharmacy Technician, or Certified Pharmacy Technician Candidate has violated a law or rule that the Board is empowered to enforce, and if continued practice by the Pharmacist, Pharmacy Intern, Certified Pharmacy Technician, or Certified Pharmacy Technician Candidate would create an imminent risk of harm to the public. The Suspension shall take effect upon written notice to the Pharmacist, Pharmacy Intern, Certified Pharmacy Technician, or Certified Pharmacy Technician Candidate specifying the statute or rule violated. At the time it issues the Suspension notice,

---

<sup>84</sup> A Pharmacist who is under investigation or who has been charged with a violation of the Pharmacy Practice Act may agree to voluntarily surrender his or her license. When this occurs, the Board should formally enter stipulated findings and an order describing the terms and conditions of the surrender including any agreed upon time limitations. This establishes statutory grounds that would support disciplinary action, and prevents a Pharmacist who has surrendered a license from applying for reinstatement within a time frame unacceptable to the Board.

<sup>85</sup> The procedures which must be followed before disciplinary action can be taken in many of the states are determined by the Administrative Procedures Act. The *Model Act* was drafted on the assumption that such an Act was in effect.



the Board shall schedule a disciplinary hearing to be held under the Administrative Procedures Act within 20 days thereafter. The Pharmacist, Pharmacy Intern, Certified Pharmacy Technician, or Certified Pharmacy Technician Candidate shall be provided with at least 10 days notice of any hearing held under this subsection.

- (b) Notwithstanding any provisions of the State Administrative Procedures Act, the Board may, in its own name, issue a Cease and Desist order to stop an individual from engaging in an unauthorized Practice of Pharmacy or violating or threatening to violate a statute, rule, or order that the Board has issued or is empowered to enforce. The Cease and Desist order must state the reason for its issuance and give notice of the individual's right to request a hearing under applicable procedures as set forth in the Administrative Procedures Act. Nothing herein shall be construed as barring criminal prosecutions for violations of this Act.

## Article V

### Licensing of Facilities

#### Introductory Comment to Article V

*The fifth and last substantive Article of the Model Act concerns licensure of Pharmacies, Manufacturers, Wholesale Distributors, Repackagers, Third-Party Logistics Providers, and the like. The licensure requirements of this Article will provide a Board with knowledge of all facilities involved in the storage, Distribution, and sale of Drugs or Devices within the state and those located outside the state that are shipping Drugs or Devices into the state. They will permit a Board to verify compliance with federal requirements and better ensure against Drug or Device diversion from the legitimate channels of commerce and provide the necessary data for effective recalls and the dissemination of information.*

#### Section 501. Licensing.

- (a) The following Persons located within this State, and the following Persons located outside this State that provide services to patients within this State, shall be licensed by the Board of Pharmacy and shall annually renew their license with the Board:<sup>86</sup>
- (1) persons engaged in the Practice of Pharmacy (including Telepharmacy);
  - (2) dispensing Practitioners and Practitioner's facilities including those engaged in nonsterile<sup>87</sup> Compounding;<sup>88</sup>
  - (3) persons engaged in the Manufacture or Repackaging of Drugs or Devices;
  - (4) persons engaged in the Wholesale Distribution of Drugs or Devices;
  - (5) persons engaged in Third-Party Logistics Provider activities of Drugs or Devices;
  - (6) pharmacies where Drugs or Devices are Dispensed, or Compounded, or Pharmacist Care Services are provided;
  - (7) Outsourcing Facilities;
  - (8) Pharmacy Benefits Managers; and
  - (9) Repository Programs
- Where operations are conducted at more than one location, each such location shall be licensed by the Board of Pharmacy.
- (b) The Board shall establish by rule, under the powers granted to it under Section 212 and 213 of this Act and as may be required from time to time, under federal law, the Criteria that each Person must meet to qualify for licensure in each classification, as well as the required practice standards applicable to each type of activity and/or facility. The Board shall adopt definitions in addition to those provided in Article I, Section 105, where necessary to carry out the Board's responsibilities. The Board may issue licenses with varying restrictions to such Persons where the Board deems it necessary.<sup>89</sup>
- (c) Each Pharmacy shall have a Pharmacist-in-Charge. Whenever an applicable rule requires or prohibits action by a Pharmacy, responsibility shall be that of the owner and/or pharmacy permit holder and the

---

<sup>86</sup> State may require additional licensing/registration requirements.

<sup>87</sup> It is contemplated that dispensing Practitioners and Practitioners' facilities should only compound nonsterile human drug products and that sterile compounded human drug products should be obtained from Outsourcing Facilities.

<sup>88</sup> Licensed Practitioners authorized under the laws of this State to Compound Drugs and to Dispense Drugs to their patients in the practice of their respective professions shall meet the same standards, record-keeping requirements, counseling, and all other requirements for the Dispensing of Drugs applicable to Pharmacists.

<sup>89</sup> Section 501(b) contemplates that the Criteria for licensure, beyond minimum requirements for all Persons and Pharmacies, established in an individual entity classification could differ. For example, the Criteria that must be met by a nuclear Pharmacy will certainly differ from that of the community Pharmacy. This type of latitude places the responsibility on the Board to adopt appropriate rules to meet the situation at hand. It also provides a forum for change to meet the changing concepts of professional practice and the Distribution of Drugs and/or Devices.

Pharmacist-in-Charge of the Pharmacy, whether the owner and/or pharmacy permit holder is a sole proprietor, partnership, association, corporation, or otherwise.

- (d) Each licensed Person located outside of this State who ships, mails, Distributes, Wholesale Distributes, or Delivers Drugs or Devices in this State, or Pharmacy located outside of this State who ships, mails, Distributes, or Delivers Drugs or Devices in this State, shall comply with the laws of patients' domicile, and shall designate a registered agent in this state for service of process. Any such licensed Person or Pharmacy who does not so designate a registered agent shall be deemed to have designated the Secretary of State of this State to be its true and lawful attorney, upon whom may be served all legal process in any action or proceeding against such licensed Person growing out of or arising from such Delivery. A copy of any such service of process shall be mailed to such Person or Pharmacy by the Board by certified mail, return receipt requested, postage prepaid, at the address such licensed Person has designated on its application for licensure in this State. If any such Person is not licensed in this State, service on the Secretary of State only shall be sufficient service.<sup>90</sup>
- (e) The Board may enter into agreements with other states or with third parties for the purpose of exchanging information concerning the licensure and inspection of entities located in this jurisdiction and those located outside this State.
- (f) The Board of Pharmacy may deny or refuse to renew a license if it determines that the granting or renewing of such license would not be in the public interest.
- (g) The Board shall establish the standards that a Person must meet for initial and continued licensure under Article V.
- (h) For facilities that Compound and/or Repackage Sterile Pharmaceuticals, an initial inspection shall be required prior to initial licensure or upon initiation of sterile Compounding activity. Thereafter, an annual inspection shall be required for licensure renewal. For facilities that do not Compound Sterile Pharmaceuticals, an initial inspection, and thereafter an inspection that takes place not less than every 24 months, shall be required for purposes of licensure or licensure renewal<sup>91</sup>. Such inspection shall be performed by the following:
  - (1) the Board or its duly authorized agent;
  - (2) a duly authorized agent of a third party approved by the Board, such as the NABP Verified Pharmacy Program (VPP) (see Appendix A for the Multistate Pharmacy Inspection Blueprint); or
  - (3) for Nonresident Pharmacies, the resident state Board of Pharmacy, if the resident Board's inspection is substantially equivalent to inspection in this State, or a VPP inspection.
- (i) Agents duly authorized to conduct inspections, whether agents of the Board or an approved third party such as VPP, must be competent to inspect the facilities they are assigned to inspect to include training on any applicable State, Federal, and USP standards.
- (j) The Board may consider exempting facilities engaged solely in the Distribution of dialysate, Drugs, or Devices necessary to perform home renal dialysis to patients with chronic kidney failure from pharmacy licensure, provided that the following criteria are met:
  - (1) The dialysate, Drugs, or Devices are approved by Food and Drug Administration, as required by federal law.
  - (2) The dialysate, Drugs, or Devices are lawfully held by a manufacturer (or a manufacturer's agent) that is properly registered with the Board as a Manufacturer and/or Wholesale Drug Distributor
  - (3) The dialysate, Drugs, or Devices are held and delivered in their original, sealed labeled packaging from the Manufacturing facility.
  - (4) The dialysate, Drugs, or Devices are delivered only by the Manufacturer (or the Manufacturer's agent) and only upon receipt of a physician's order.

---

<sup>90</sup> This section provides for service of process on any Person who Dispenses, Distributes, or Delivers Drugs or Devices within the State.

<sup>91</sup> State resources may have to be considered when evaluating inspection scheduling in combination with risk assessment consideration.

- (5) The Manufacturer (or Manufacturer's agent) delivers the dialysate, Drugs, or Devices directly to:
  - (i) a patient with chronic kidney failure, or his/her designee, for the patient's self-administration of dialysis therapy, or
  - (ii) a health care provider or institution for administration or delivery of the dialysis therapy to a patient with chronic kidney failure.
- (6) Records of all sales and Distribution of dialysate, Drugs, or Devices to home dialysis patients must be retained and readily available for inspection and copying by the Board for \_\_\_\_\_ years.

## Section 502. Application.<sup>92</sup>

- (a) The Board shall specify by rule the licensure procedures to be followed, including but not limited to, specification of forms for use in applying for such licensure and times, places, and applicable fees.
- (b) Applicants for licensure to Distribute, Wholesale Distribute, Manufacture, sell, purchase, transfer, and/or produce Drugs or Devices, and applicants for licensure as a Pharmacy Benefits Manager, shall file with the Board of Pharmacy a verified application containing such information as the Board requires of the applicant relative to the qualifications for a license.
- (c) Licenses issued by the Board pursuant to this Act shall not be transferable or assignable.
- (d) The Board shall specify by rule minimum standards for responsibility of any Person, Pharmacy, or Pharmacy Benefits Manager that has employees or personnel engaged in the Practice of Pharmacy, or Manufacture, Distribution, Wholesale Distribution, production, sale, or use of Drugs or Devices in the conduct of their business. If the licensed Person is a Pharmacy located in this State, that portion of the facility to which such license applies shall be operated only under the direct supervision of a Pharmacist licensed to practice in this State. If that Person is an Outsourcing Facility, all Compounding at the facility shall be under the direct supervision of a licensed Pharmacist and comply with Federal requirements applicable to Outsourcing Facilities.
- (e) A "surety" bond of not less than \$100,000, or other equivalent means of security acceptable to the Board or a third party recognized by the Board such as insurance, an irrevocable letter of credit, or funds

---

<sup>92</sup> Boards may want to consider requesting the following information on applications for Pharmacy and Wholesale Distributor licensure:

- (a) personal information;
- (b) marital information;
- (c) family information (parents, siblings, in-laws);
- (d) education;
- (e) military information;
- (f) arrests, detentions, litigations, and arbitrations;
- (g) residences (past 25 years);
- (h) employment (back to age 18);
- (i) character references;
- (j) safe deposit box or other depository information;
- (k) privileged, occupational, or professional licensure;
- (l) out-of-state business, venture, or industry licensure or financial interest in such;
- (m) appearances before any licensing agency or similar authority in or outside the state;
- (n) denials of a personal license, permit, certificate, or registration for a privileged, occupational, or professional activity;
- (o) denials of a business or industry license or related finding of suitability, or participation in a group that has been denied a business or industry license or related finding of suitability;
- (p) Administrative actions or proceedings related to the pharmaceutical industry or participation in a group that has been the subject of such administrative actions or proceedings;
- (q) guilty findings or pleadings or pleas of nolo contendere to any offense, federal or state, related to prescription Drugs and/or controlled substances or participation in a group that has been found or pled guilty or that has pled nolo contendere to any such offense;
- (r) surrender, voluntary or otherwise, of licensure, permit, or certificate of registration relating to the pharmaceutical industry, or participation in a group that has surrendered, voluntary or otherwise, any such licensure, permit, or certificate of registration; and
- (s) any relatives within the fourth degree of consanguinity associated with or employed in the pharmaceutical or Drug-related industry.

deposited in a trust account or financial institution, to secure payment of any administrative penalties imposed by the Board and any fees or costs incurred by the Board regarding that licensee when those penalties, fees, or costs are authorized under state law and the licensee fails to pay thirty (30) days after the penalty, fee, or costs becomes final. A separate surety bond or other equivalent means of security is not required for each company's separate locations or for affiliated companies/groups when such separate locations or affiliated companies/groups are required to apply for or renew their Wholesale Distributor license with the Board. The Board may make a claim against such bond or other equivalent means of security until one year after the Wholesale Distributor's license ceases to be valid or until sixty (60) days after any administrative or legal proceeding before or on behalf of the Board that involves the Wholesale Distributor is concluded, including any appeal, whichever occurs later. Manufacturers shall be exempt from securing a "surety" bond or other equivalent means of security acceptable to the Board or a third party recognized by the Board. The Board may waive the bond requirement, if the Wholesale Distributor:

- (1) has previously obtained a comparable surety bond or other equivalent means of security for the purpose of licensure in another state, where the Wholesale Distributor possesses a valid license in good standing; or
- (2) is a publicly held company.

### **Section 503. Notifications.**

- (a) All licensed Persons shall report to the Board of Pharmacy the occurrence of any of the following:
  - (1) permanent closing;
  - (2) change of ownership, management, location, or Pharmacist-in-Charge of a Pharmacy;
  - (3) any theft or loss of Drugs or Devices;
  - (4) any conviction of any employee of any State or Federal Drug laws;
  - (5) any criminal conviction or pleas of guilty or nolo contendere of all licensed or registered personnel;
  - (6) disasters, accidents, or any theft, destruction, or loss of records required to be maintained by State or Federal law;
  - (7) occurrences of Significant Quality-Related Events;
  - (8) Significant Adverse Drug Reaction associated with Compounded Drugs;
  - (9) recalls of Compounded Drugs;
  - (10) recalls of sterile Repackaged Drugs;
  - (11) illegal use or disclosure of Protected Health Information; or
  - (12) any and all other matters and occurrences as the Board may require by rule.
- (b) All licensed Persons shall report to the Board of Pharmacy, or its authorized agent, if they are engaging in any sterile Compounding activity conducted at a licensed facility prior to commencing of any sterile Compounding activity and at least in a manner determined by the Board. The Board may establish by rule additional reporting requirements for sterile and nonsterile Compounding activities.
- (c) All licensed Persons shall report to the Board of Pharmacy, or its authorized agent, the occurrence of any Pharmacy or Pharmacy-related inspection conducted by any State or Federal regulatory agency or authorized agent thereof and shall provide a copy of the report of such inspection, including applicable documents relating to corrective actions<sup>93</sup>.

### **Section 504. Grounds, Penalties, and Reinstatement.**

- (a) No Person, Pharmacy, or Pharmacy Benefits Manager designated in Section 501 of this Act shall operate until a license has been issued to said Person, Pharmacy, or Pharmacy Benefits Manager by the Board.

---

<sup>93</sup> This includes any report or inspectional observations and any related correspondence with the Federal or State agency.

- (b) Except where otherwise permitted by law, it shall be unlawful for a Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor to Distribute or Deliver Drugs or Devices to any Person in this State not licensed under this statute. Any Person who shall Distribute or Deliver Drugs or Devices to a Person not licensed shall be subject to a fine to be imposed by the Board not to exceed one thousand dollars (\$1,000) for each offense in addition to such other disciplinary action the Board may take under this Act. Except as otherwise indicated in this Act, each such violation shall also constitute a misdemeanor punishable upon conviction as provided in the criminal code of this State.
- (c) The Board may Suspend, Revoke, deny, or refuse to renew the license of any Person, Manufacturer, Repackager, Third-Party Logistics Provider, Wholesale Distributor, Pharmacy, or Pharmacy Benefits Manager on any of the following grounds:<sup>94</sup>
- (1) the finding by the Board of violations of any Federal, State, or local laws relating to the Practice of Pharmacy, Drug samples, Wholesale or retail Drug or Device Distribution, or Distribution of controlled substances;
  - (2) any felony convictions under Federal, State, or local laws;
  - (3) the furnishing of false or fraudulent material in any application made in connection with Drug or Device Manufacturing or Distribution;
  - (4) suspension or Revocation by Federal, State, or local government of any license currently or previously held by the applicant for the Manufacture or Distribution of any Drugs or Devices, including controlled substances;
  - (5) willfully and knowingly submitting false or fraudulent information or omitting information on due diligence questionnaires and/or attestation documents regarding the purchase or receipt of Drugs from Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors.
  - (6) obtaining any remuneration by fraud, misrepresentation, or deception;
  - (7) affiliating with websites that may deceive or defraud patients or that violate Pharmacy or Drug laws of this State or rules and regulations pertaining thereto; or of laws, rules, and regulations of any other state; or of the Federal government;
  - (8) dealing with Drugs or Devices that he or she knows or should have known are Counterfeit, Contraband, or stolen Drugs or Devices;<sup>95</sup>
  - (9) purchasing or receiving of a Drug or Device from a source other than a Person or pharmacy licensed under the laws of the State, except where otherwise provided;
  - (10) the transfer during any consecutive twelve (12)-month period by a Pharmacy to a Wholesale Distributor or to another Pharmacy of more than five percent (5%) of the total amount of Prescription Drugs or Devices purchased by the Pharmacy in the immediately preceding twelve (12)-month period. The following are not subject to the provisions of this subsection:
    - (i) Prescription Drugs or Devices that are Returned by a pharmacy for credit, in an amount equal to or less than the actual purchase price, to the Wholesale Distributor or Manufacturer from which those Products were purchased;
    - (ii) Intracompany sales;
    - (iii) The sale, purchase, or trade of a Drug or an offer to sell, purchase, or trade a Drug among hospitals or other health care entities that are under common control;
    - (iv) The sale, purchase, or trade of a Drug or the offer to sell, purchase, or trade a Drug by a charitable organization described in 503(c)(3) of the Internal Revenue Code of 1954 to a nonprofit affiliate of the organization to the extent otherwise permitted by law;

---

<sup>94</sup> The Prescription Drug Marketing Act of 1987 (PDMA) requires that the state licensing laws provide for the Suspension or Revocation of licenses upon conviction for violation of Federal, State, or local Drug laws or rules pertaining to the unlawful Distribution of Drugs at wholesale. The PDMA defines fines, imprisonment, or civil penalties.

<sup>95</sup> This section restricts Distribution of Drugs or Devices to licensed entities to help ensure against clandestine Distribution to unauthorized and unlicensed Persons.

- (v) The purchase or other acquisition by a hospital or other similar health care entity that is a member of a group purchasing organization of a Drug for its own use from the group purchasing organization or from other hospitals or similar health care entities that are members of these organizations; and
- (vi) The transfer of Prescription Drugs or Devices between Pharmacies pursuant to a Centralized Prescription Processing or Filling agreement.
- (11) the transfer during any consecutive twelve (12)-month period by a Wholesale Distributor to a Wholesale Distributor of more than five percent (5%) of the total amount of prescription Drugs or Devices purchased by Wholesale Distributor in the immediately preceding twelve (12)-month period;
- (12) Wholesale Drug Distributors other than pharmacies Dispensing or Distributing Drugs or Devices directly to patients;
- (13) violations of any of the provisions of this Act or of any of the Rules adopted by the Board under this Act; or
- (14) illegal use or disclosure of Protected Health Information.
- (d) Reinstatement of a license that has been Suspended, Revoked, or restricted by the Board may be granted in accordance with the procedures specified by Section 401 of this Act.

#### **Section 505. Criminal Offense; Forfeiture of Property.**

- (a) Violation of any of the provisions of Article V of this Act by any person engaged in the Wholesale Distribution of Drugs and Devices shall constitute a Class three felony, provided that any such violation that results in the death of a Person shall constitute a Class one felony.
- (b) A Person engaged in the Wholesale Distribution of Drugs and Devices convicted by a criminal court of this State of violating any of the provisions of Article V may be ordered by the court to forfeit to the State any real or personal property:
  - (1) used or intended to be used to commit, to facilitate, or to promote the commission of such offense; or
  - (2) constituting, derived from, or traceable to the gross proceeds that the defendant obtained directly or indirectly as a result of the offense. Any property or assets subject to forfeiture under this section may be seized pursuant to a warrant obtained in the same manner as a search warrant or as otherwise permitted by law, and held until the case against the defendant is adjudicated. Monies ordered forfeited, or proceeds from the sale of other assets ordered forfeited, shall be equitably divided between the Board and other agencies involved in the investigation and prosecution that led to the conviction. Other property ordered forfeited after conviction of the defendant may, at the discretion of the investigating agencies, be placed into official use by the Board or the agencies involved in the investigation and prosecution that led to the conviction.

## **Article VI**

### **Other**

#### **Section 601. Severability.**

If any provision of this Act is declared unconstitutional or illegal, or the applicability of this Act to any Person, Pharmacy, or circumstance is held invalid by a court of competent jurisdiction, the constitutionality or legality of the remaining provisions of this Act and the application of this Act to other Persons, Pharmacies, and circumstances shall not be affected and shall remain in full force and effect without the invalid provision or application.

#### **Section 602. Effective Date.**

This Act shall be in full force and effect on \_\_\_\_\_.



# National Association of Boards of Pharmacy

## Model Rules

### Model Rules for Pharmacy Interns

#### Section 1. Licensure.

Every individual shall be licensed by the Board of Pharmacy before beginning Pharmacy practice experiences in this State.<sup>96</sup> A license to practice Pharmacy as a Pharmacy Intern shall be granted only to those individuals who:

- (a) are enrolled in a professional degree program of a school or college of pharmacy that has been approved by the Board and satisfactorily progressing toward meeting the requirements for licensure as a Pharmacist; or
- (b) are graduates of an approved professional degree program of a school or college of Pharmacy or are graduates who have established educational equivalency by obtaining a Foreign Pharmacy Graduate Examination Committee™ (FPGEC®) Certificate, who are currently licensed by the Board of Pharmacy for the purpose of obtaining practical experience as a requirement for licensure as a Pharmacist; or
- (c) are qualified applicants awaiting examination for licensure or meeting Board requirements for re-licensure;
- (d) are participating in a residency or fellowship program; or
- (e) have undergone a state and federal fingerprint-based criminal background check as specified by Board rule.

#### Section 2. Identification.

The Pharmacy Intern shall be so designated in his or her professional relationships, and shall in no manner falsely assume, directly or by inference, to be a Pharmacist. The Board shall issue to the Pharmacy Intern a license for purposes of identification and verification of his or her role as a Pharmacy Intern, which license shall be surrendered to the Board upon discontinuance of Pharmacy practice experiences for any reason including licensure as a Pharmacist. No individual not properly licensed by the Board as a Pharmacy Intern shall take, use, or exhibit the title of Pharmacy Intern, or any other term of similar like or import.

#### Section 3. Supervision.

A Pharmacy Intern shall be allowed to engage in the Practice of Pharmacy provided that such activities are under the supervision of a Pharmacist. A Pharmacist shall be in contact with, and actually giving instructions to, the Pharmacy Intern during all professional activities throughout the entire Pharmacy practice experience period. The Pharmacist

---

<sup>96</sup> The ACPE Accreditation Standards and Guidelines for the Professional Program in Pharmacy Leading to the Doctor of Pharmacy Degree (effective July 1, 2007) requires schools and colleges of Pharmacy seeking and maintaining ACPE accreditation to incorporate introductory Pharmacy practice experiences within their professional curricula, and such experiences must account for not less than 5% of the total curricular length (not less than 300 contact hours). Under the supervision of a Preceptor and usually taken throughout the first three academic years of the professional program, these introductory Pharmacy practice experiences expose students to and allow students to participate in activities such as processing/Dispensing Medication Orders, conducting Patient interviews, or presenting Patient cases in an organized format.

It is also encouraged that Boards of Pharmacy allow Pharmacy students to be registered as Pharmacy Interns as early as initial enrollment in a Board-approved professional program as long as the Pharmacy student has begun to take professional degree courses.

is responsible for supervising all the Practice of Pharmacy activities performed by the Pharmacy Intern, including but not limited to the accurate Dispensing of the Drug.<sup>97</sup>

#### **Section 4. Change of Address.**

All Pharmacy Interns shall notify the Board immediately upon change of employment and residential address.

#### **Section 5. Evidence of Completion.**

Applicants for licensure as Pharmacists shall submit evidence that they have satisfactorily completed: (1) an objective assessment mechanism intended to evaluate achievement of desired competencies; and (2) not less than 1,740 hours of Pharmacy practice experience credit under the instruction and supervision of a Preceptor.<sup>98</sup>

---

<sup>97</sup> According to the ACPE Accreditation Standards and Guidelines, most Pharmacy practice experiences must be under the supervision of a qualified Pharmacist Preceptor licensed in the United States. Realizing that in some cases non-Pharmacist Preceptors can also provide valuable learning opportunities, it is hoped that Boards of Pharmacy recognize these experiences and that schools and colleges of pharmacy ensure, in most cases through faculty, that the desired competencies are being met.

Supervision includes an actual review of the Prescription Drug Order and the dispensed Drug or Product to ensure public protection.

<sup>98</sup> These requirements coincide with the ACPE Accreditation Standards and Guidelines. Boards of pharmacy are strongly encouraged to utilize these Accreditation Standards and Guidelines as a basis for the establishment and revision of Board standards for Pharmacy practice experiences.

Introductory Pharmacy practice experiences, which are not less than 300 contact hours, are in addition to the advanced practice experiences taken during the final professional year, which account for not less than 25 % of the curricular length or 1,440 contact hours. The total Pharmacy practice experience hour requirement, therefore, is not less than 1,740 hours. Boards may consider moving away from requiring a specific number of contact hours should it be determined that the ACPE Accreditation Standards and Guidelines result in appropriate preparation for students and objective assessment mechanisms demonstrate such.

## **Model Standards for Pharmacy Practice Experience Programs**

### **Section 1. Preceptor.**

- (a) The Pharmacy Intern, excluding those who are currently enrolled in a professional degree program of a school or college of pharmacy approved by the Board and satisfactorily progressing toward meeting the requirements for licensure as a Pharmacist, shall notify the Board of Pharmacy within two weeks of beginning practice as a Pharmacy Intern, on a form provided by the Board, of the identity of the Pharmacy practice experience site and of the Preceptor.
- (b) A Preceptor may be responsible for the training of more than one Pharmacy Intern. The number of Pharmacy Interns engaged in the Practice of Pharmacy at any time is limited to the number of Pharmacy Interns the Pharmacist can appropriately precept as approved by the Board.

### **Section 2. Pharmacy Practice Experience Programs.<sup>99</sup>**

- (a) The Pharmacy at which a Pharmacy Intern is being trained shall provide an environment that is conducive to the learning of the Practice of Pharmacy by a Pharmacy Intern. Pharmacy practice experience sites shall meet the standards approved by the Board.
- (b) Pharmacy practice experience in non-traditional practice sites (eg, industry-sponsored programs) must be approved by the Board of Pharmacy prior to granting of credit.
- (c) When a Pharmacy Intern desires to obtain credit for training received in a state other than this State, the Pharmacy Intern shall abide by all the provisions of the Pharmacy practice experience rules in that state, and shall provide evidence from that state's Board of Pharmacy of the number of clock hours of experience actually participated in by the Pharmacy Intern.

### **Section 3. Global Exchange Pharmacy Students.**

A Global Exchange Pharmacy Student may participate in observation-only clinical learning experiences, not to exceed \_\_\_\_\_, provided:

- (a) the Global Exchange Pharmacy Student has been reviewed and qualified by the ACPE-accredited or Board-approved school or college of pharmacy as exists for Introductory Pharmacy Practice Experience (IPPE) and Advanced Pharmacy Practice Experience (APPE) experiential rotations; and
- (b) he or she is under the direct in-person supervision of a Pharmacist.

---

<sup>99</sup> Boards of pharmacy are strongly encouraged to utilize the ACPE Accreditation Standards and Guidelines for the Professional Program in Pharmacy Leading to the Doctor of Pharmacy Degree (effective July 1, 2007) as a basis for the establishment and revision of Board standards for Pharmacy practice experiences.

## **Model Rules for Institutional Pharmacy**

### **Section 1. Applicability.**

The following Rules are applicable to all Institutional Facilities and Institutional Pharmacies as defined in Section 105 of the Model State Pharmacy Act.

### **Section 2. Absence of Pharmacist at a Pharmacy Located Within an Institutional Facility.**

- (a) During such times as when a Pharmacy, which is located within an Institutional Facility, may be unattended by a Pharmacist, arrangements shall be made in advance by the Pharmacist-in-Charge for provision of Drugs to the medical staff and other authorized personnel of the Institutional Facility by use of night cabinets and, in emergency circumstances, by access to the Pharmacy. A Pharmacist must be “on call” during all absences.
- (b) In the absence of a Pharmacist, Drugs shall be stored in a locked cabinet, Automated Pharmacy System, or other enclosure constructed and located outside of the Pharmacy area, to which only specifically authorized personnel may obtain access by key or combination, and which is sufficiently secure to deny access to unauthorized persons. The Pharmacist-in-Charge shall, in conjunction with the appropriate committee of the Institutional Facility, develop inventory listings of those Drugs to be included in such cabinet(s) and determine who may have access, and shall ensure that:
  - (1) Drugs are properly Labeled;
  - (2) only prepackaged Drugs are available, in amounts sufficient for immediate therapeutic requirements;
  - (3) whenever access to the cabinet occurs, written Practitioner’s orders and proofs-of-use are provided;
  - (4) all Drugs therein are inventoried no less than once per week unless stored in an Automated Pharmacy System;
  - (5) a complete audit of all activity concerning such cabinet is conducted no less than once per month; and
  - (6) written policies and procedures are established to implement the requirements of this Section.
- (c) Whenever any Drug is not available from floor supplies or night cabinets, and such Drug is required to treat the immediate needs of a patient whose health would otherwise be jeopardized, such Drug may be obtained from the Pharmacy in accordance with the requirements of this Section. One supervisory nurse in any given shift is responsible for obtaining Drugs from the Pharmacy. The responsible nurse shall be designated in writing by the appropriate committee of the Institutional Facility. Removal of any Drug from the Pharmacy by an authorized nurse must be recorded on a suitable form showing the patient name, room number, name of Drug, strength, amount, date, time, and signature of nurse. The form shall be left with the container from which the Drug was removed.

### **Section 3. Emergency Kit Use by Institutional Facilities.**

- (a) Emergency kit Drugs may be provided for use by authorized personnel of the Institutional Facility provided, however, such kits meet the following requirements:
  - (1) Emergency kit Drugs are those Drugs which may be required to meet the immediate therapeutic needs of patients and which are not available from any other authorized source in sufficient time to prevent risk of harm to patients by delay resulting from obtaining such Drugs from such other sources.
  - (2) All emergency kit Drugs shall be provided and sealed by a Pharmacist or his or her designee in accordance with applicable security and inventory control policies and procedures.
  - (3) The supplying Pharmacist and the medical staff of the Institutional Facility shall jointly determine the Drugs, by identity and quantity, to be included in emergency kits.

- (4) Emergency kits shall be stored in secured areas to prevent unauthorized access, and to ensure a proper environment for preservation of the Drugs within them.
- (5) The exterior of each emergency kit shall be labeled so as to clearly indicate that it is an emergency Drug kit and that it is for use in emergencies only. The label shall contain a listing of the Drugs contained in the kit, including name, strength, quantity, and expiration date of the contents, and the name, address(es), and telephone number(s) of the supplying Pharmacy.
- (6) Drugs shall be removed from emergency kits only pursuant to a valid Chart Order.
- (7) Whenever an emergency kit is opened, the supplying Pharmacy shall be notified and the Pharmacy shall restock and reseal the kit within a reasonable time so as to prevent risk of harm to patients.<sup>100</sup>
- (8) The expiration date of an emergency kit shall be the earliest date of expiration of any Drug supplied in the kit. Upon the occurrence of the expiration date, the supplying Pharmacy shall replace the expired Drug.
- (9) The Pharmacy that supplies controlled substances for emergency kits must comply with applicable state and federal requirements.

#### **Section 4. Drug Distribution and Pharmacist Care Services.**

- (a) The Pharmacist-in-Charge shall establish written procedures for the safe and efficient acquisition, handling, storage, and Dispensing of Drugs, including investigational Drugs, and for the provision of Pharmacist Care Services. An annual updated copy of such procedures shall be on hand for inspection by the Board of Pharmacy.
- (b) A Pharmacist may engage in Therapeutic Interchange or formulary substitution as authorized by the facility's interdisciplinary committee<sup>101</sup> of health care providers, at a minimum to include a Practitioner and a Pharmacist. Proper record of this authority should be maintained at the Pharmacy.
- (c) To ensure continuous patient care, the facility's director of nursing or his or her documented licensed health care designee may transmit the Chart Order to a Pharmacy.<sup>102</sup>
- (d) The Pharmacist shall assess each patient's medication regimen based on a review of the health record, either remotely or on site, in a timely manner that promotes improving patient clinical outcomes, medication safety and education, and appropriate care management.
- (e) If the Institutional Pharmacy is not located within an Institutional Facility, the Pharmacist-in-Charge may designate a licensed nurse to restock an Automated Pharmacy System using verification technology such as bar code scanning, electronic, or other technology.
- (f) Institutional Pharmacies either located within or not within Institutional Facilities may Dispense medication to patients upon discharge in order to ensure a transition of care between settings until a new Prescription Drug Order is issued.

#### **Section 5. Shared Pharmacy Services Utilization for Immediate Need.<sup>103</sup>**

- (a) In accordance with the Model Rules for the Practice of Pharmacy and Shared Pharmacy Services, an Institutional Pharmacy may outsource services to another Pharmacy for the limited purpose of ensuring that Drugs or Devices are attainable to meet the immediate needs of patients and residents of the

---

<sup>100</sup> When the Pharmacist restocks and reseals the emergency kit Drugs, it is recommended that a lock or other similar device be used to assure that unauthorized access to the kit is minimized.

<sup>101</sup> This is often referred to as the pharmacy and therapeutics committee or the quality assessment and assurance committee.

<sup>102</sup> Federal law may restrict who can transmit a Chart Order for a controlled substance.

<sup>103</sup> Although Institutional Pharmacies primarily outsource services to another Pharmacy for the purposes of meeting the immediate needs of patients and residents when the Institutional Pharmacy is closed, it is also recognized that other services may be outsourced that the Institutional Pharmacy is not able to provide on an ongoing basis.

Institutional Facility or when the Institutional Pharmacy cannot provide services on an ongoing basis, provided that the Institutional Pharmacy:

- (1) has obtained approval from the Institutional Facility to outsource Shared Pharmacy Services for its inpatients and residents; and
- (2) shares a valid Chart Order with the Pharmacy it has contracted with for the Shared Pharmacy Services without the need to transfer the order.

#### **Section 6. Packaging of Previously Dispensed Medication.**

- (a) At a patient's or patient's caregiver's request, a Pharmacy may change the packaging of a Drug previously Dispensed to the patient.
- (b) Any Pharmacy providing packaging services shall have in place policies and procedures to:
  - (1) assess whether the medication may be Adulterated or Misbranded; and
  - (2) package and label the medication in compliance with state and federal requirements and USP standards.
- (c) The Pharmacy that packages a previously Dispensed medication shall retain all original prescription information in accordance with state record-keeping requirements.

#### **Section 7. Institutional Pharmacy Delivery Room.**

Prescription Drugs, Devices, and other Products restricted to sale or Dispensing by, or under the supervision of, a Pharmacist must be stored in the Pharmacy and must not be sold, Delivered, or otherwise removed from a Pharmacy unless a Pharmacist is present, under the following:

- (a) Institutional Pharmacies that are not located within an Institutional Facility may accept Returns or otherwise Deliver fulfilled, verified, and packaged prescription medication in the absence of a Pharmacist or when the Pharmacy is closed for business if the Pharmacy and the Pharmacist-in-Charge maintain written policies and procedures for secured Delivery area storage and removal of prescriptions.
- (b) A Pharmacist or a Pharmacy, by means of its Delivery personnel, may accept the Return of the following Drugs or Devices to the secured Delivery area:
  - (1) emergency kits;
  - (2) prescription medications that were unsuccessfully Delivered by the Pharmacy personnel or Delivery personnel; and
  - (3) prescription medications eligible for Return pursuant to applicable state and federal law.

# Model Rules for the Practice of Pharmacy

## Introductory Comment

*The Board finds that in the interest of protecting the public health and welfare, in order to ensure optimum effect of Drug therapy, and to maximize the quality of Pharmacist Care Services, the following rules are essential.*

## Section 1. Facility.

- (a) To obtain a license for a Pharmacy, an applicant shall:
  - (1) have submitted an application in the form prescribed by the Board of Pharmacy;
  - (2) have attained the age of majority; and
  - (3) have paid the fees specified by the Board of Pharmacy for the issuance of the license.
- (b) The facility owner, if an individual, shall have undergone a state and federal fingerprint-based criminal background check as specified by Board rule;
- (c) The facility shall have undergone a Pharmacy inspection by the Board or authorized agent thereof; and
- (d) Possess the following minimum requirements for a Pharmacy:
  - (1) Each Pharmacy shall be of sufficient size, as determined by the Board, to allow for the safe and proper storage of Prescription Drugs and for the safe and proper Compounding and/or preparation of Prescription Drug Orders.
  - (2) Each Pharmacy shall maintain an area designated for the provision of Patient Counseling services. This area shall be designed to provide a reasonable expectation of privacy of Protected Health Information.
  - (3) Each Pharmacy shall have ready access to references, to include at least one current reference<sup>104</sup> in each of the following in each of the following categories, if applicable to the services provided:
    - (i) State and Federal Drug laws relating to the Practice of Pharmacy and the legal Distribution of Drugs and any rules or regulations adopted pursuant thereto;
    - (ii) pharmacology;
    - (iii) dosage and toxicology;
    - (iv) veterinary Drugs<sup>105</sup>; and
    - (v) general.
  - (4) Each Pharmacy shall maintain patient-oriented reference material for guidance in proper Drug usage.<sup>106</sup>
  - (5) Each Person involved in the development, maintenance, or use of a Drug formulary shall maintain a currently accepted reference containing guidelines for a sound Drug formulary system.
  - (6) All areas where Drugs and Devices are stored shall be dry, well lighted, well ventilated, and maintained in a clean and orderly condition. Storage areas shall be maintained at temperatures which will ensure the integrity of the Drugs prior to their Dispensing as stipulated by the United States Pharmacopeia–National Formulary (USP-NF) and/or the Manufacturer’s or Distributor’s Product Labeling unless otherwise indicated by the Board.
  - (7) Each Pharmacy shall have access to a sink with hot and cold running water that is convenient to the Compounding area for the purpose of hand scrubs prior to Compounding.
  - (8) Equipment/Supplies.

---

<sup>104</sup> Boards may wish to give examples in each of these categories of reference texts.

<sup>105</sup> Such as Plumb’s Veterinary Drug Handbook.

<sup>106</sup> Patient-oriented reference material can include publications such as Facts and Comparisons’ Patient Drug Facts, or the United States Pharmacopeia Dispensing Information (USPDI).

The Pharmacy shall carry and utilize the equipment and supplies necessary to conduct a Pharmacy in a manner that is in the best interest of the patients served and to comply with all State and Federal laws.

- (9) The Pharmacy shall provide a means for patients to prevent disclosure of Confidential Information or personally identifiable information that was obtained or collected by the Pharmacist or Pharmacy incidental to the Delivery of Pharmacist Care Services other than as authorized by law or rules of the Board.
- (10) The Pharmacy, if conducting business over the Internet, shall be accredited by a program approved by the Board.
- (e) Upon renewal, the licensee shall provide to the Board the NABP e-Profile ID of the Pharmacy and the Pharmacist-in-Charge

## **Section 2. Security.**

- (a) Facility
  - (1) Each Pharmacist, while on duty, shall be responsible for the security of the Pharmacy, including provisions for effective control against theft or diversion of Drugs and/or Devices.
  - (2) The Pharmacy shall be secured by either a physical barrier with suitable locks and/or an electronic barrier to detect entry at a time when the Pharmacist is not present. Such barrier shall be approved by the Board of Pharmacy before being put into use. In the event of separation of employment of an employee due to any confirmed Drug-related reason, including diversion, or other acts involving dishonesty, suitable action shall be taken to ensure the security of the pharmacy.
  - (3) Prescription and other patient health care information shall be maintained in a manner that protects the integrity and confidentiality of such information as provided by the rules of the Board.
  - (4) The Pharmacy shall implement and maintain technologies that will aid in theft prevention and suspect apprehension that may include:
    - (i) Video equipment positioned to identify individuals who may be involved in diversion or theft, if utilized, shall have adequate recording, storage, and retrieval capabilities; and
    - (ii) monitored alarm system with backup mechanism.
- (b) Internal Theft/Diversion
  - (1) the Pharmacist-in-Charge and owner/licensee (facility permit holder) shall ensure policies and procedures are in place that address the following:
    - (i) inspection of shipments;
    - (ii) receipt Verification oversight and checking in shipments;
    - (iii) reconciliation of orders; and
    - (iv) inventory management including:
      - (A) determination of Medications that need to be monitored and controlled beyond existing systems such as controlled substances and Drugs of Concern; and
      - (B) conducting quarterly reconciliations at a minimum but shall be more frequent up to perpetual, depending on the potential for or incidence of diversion for a particular Drug.

## **Section 3. Personnel.**

- (a) Duties and Responsibilities of the Pharmacist-in-Charge
  - (1) No Person shall operate a Pharmacy without a Pharmacist-in-Charge. The Pharmacist-in-Charge of a Pharmacy shall be designated in the application of the Pharmacy for license, and in each renewal thereof. A Pharmacist may not serve as Pharmacist-in-Charge unless he or she is physically present in the Pharmacy a sufficient amount of time to provide supervision and control. A Pharmacist may



not serve as Pharmacist-in-Charge for more than one Pharmacy at any one time except upon obtaining written permission from the Board.

- (2) The Pharmacist-in-Charge has the following responsibilities:
- (i) Developing or adopting, implementing, and maintaining:<sup>107</sup>
    - (A) Policies and procedures addressing the following:
      - (-a-) the provision of Pharmacy services;<sup>108</sup>
      - (-b-) the procurement, storage, security, and disposition of Drugs and Devices, particularly controlled substances and Drugs of Concern;
      - (-c-) computerized record-keeping systems;
      - (-d-) Automated Pharmacy Systems;
      - (-e-) preventing the illegal use or disclosure of Protected Health Information, or verifying the existence thereof and ensuring that all employees of the Pharmacy read, sign, and comply with such established policies and procedures;
      - (-f-) operation of the Pharmacy in the event of a fire, flood, pandemic, or other natural or man-made disaster or emergency, to the extent that the Pharmacy can be safely and effectively operated and the Drugs contained therein can be safely stored and Dispensed. Such policies and procedures shall include reporting to the Board the occurrence of any fire, flood, or other natural or man-made disaster or emergency within 10 days of such occurrence<sup>109</sup>;
      - (-g-) the proper management of Drug recalls which may include, where appropriate, contacting patients to whom the recalled Drug Product(s) have been Dispensed;
      - (-h-) the duties to be performed by Certified Pharmacy Technicians and Certified Pharmacy Technician Candidates. The duties and responsibilities of these personnel shall be consistent with their education, training, and experience and shall address the method and level of necessary supervision specific to the practice site. These policies and procedures shall, at a minimum, specify that Certified Pharmacy Technicians and Certified Pharmacy Technician Candidates are not assigned duties that may be performed only by a Pharmacist. Such policies and procedures shall also specify that Certified Pharmacy Technician Candidates shall not be assigned duties that may be performed only by Certified Pharmacy Technicians.
      - (-i-) actions to be taken to prevent and react to pharmacy robberies and thefts, including but not limited to coordinating with law enforcement, training, mitigation of harm, and protecting the crime scene.
      - (-j-) the PIC shall have policies and procedures in place that restrict and monitor control over and access to the locks, barriers, and systems used to secure the pharmacy and pharmacy systems in accordance with state laws and regulations.
    - (B) Policies and procedures that address the following activities related to prescription medication shipment by mail or common carrier:
      - (-a-) properly transferring prescription information to an alternative Pharmacy of the patient's choice in situations where the medication is not Delivered or Deliverable;

---

<sup>107</sup> The owner and/or pharmacy permit holder, along with the Pharmacist-in-Charge, are responsible for these policies and procedures.

<sup>108</sup> The Pharmacist-in-Charge, as part of the responsibilities to manage as effectively as possible a patient's therapy to avoid a harmful interruption of therapy because of a shortage or limited Distribution of medications, can proactively improve Pharmacy operations by developing a systematic approach to address such circumstances. References such as the American Society of Health-System Pharmacists (ASHP) Guidelines in Managing Drug Product Shortages could be used as resources for developing policies and procedures if appropriate. Additionally, Food and Drug Administration maintains a list of current and resolved Drug shortages, as well as discontinued Drugs on the agency's Drug Shortages Web page at [www.fda.gov/cder/drug/shortages](http://www.fda.gov/cder/drug/shortages).

<sup>109</sup> States should recognize that hospitals, in order to prepare for a disaster or emergency, may be stocking emergency supplies of medications in areas outside the licensed pharmacy. Hospitals should be encouraged to expand the space allotted to the licensed pharmacy area to accommodate the need to store emergency supplies.

- (-b-) verifying that common carriers have in place security provisions, such as criminal background checks and random drug screens on its employees who have access to prescription medications;
  - (-c-) tracking all shipments; and
  - (-d-) ensuring that Drugs do not become adulterated in transit
- (C) Quality assurance programs addressing the following:
  - (-a-) Pharmacy services. The quality assurance program should be designed to objectively and systematically monitor and evaluate the quality and appropriateness of patient care, pursue opportunities to improve patient care, and resolve identified problems;
  - (-b-) Automated Pharmacy Systems. The quality assurance program should monitor the performance of the Automated Pharmacy System, ensure the Automated Pharmacy System is in good working order and accurately Dispenses the correct strength, dosage form, and quantity of the Drug prescribed, while maintaining appropriate record keeping and security safeguards; and
  - (-c-) The prevention and detection of Drug diversion.<sup>110</sup>
- (ii) Ensuring that:
  - (A) all Pharmacists, Pharmacy Interns, Certified Pharmacy Technicians, and Certified Pharmacy Technician Candidates employed at the Pharmacy are currently licensed by the Board of Pharmacy.
- (iii) Notifying the Board of Pharmacy, immediately and in writing, of any of the following<sup>111</sup> changes:
  - (A) change of employment or responsibility as the Pharmacist-in-Charge;
  - (B) the separation of employment of any Pharmacist, Pharmacy Intern, Certified Pharmacy Technician Candidate, or Certified Pharmacy Technician for any confirmed Drug-related reason, including but not limited to, Adulteration, abuse, theft, or diversion, and shall include in the notice the reason for the termination: if it is the employment of the Pharmacist-in-Charge that is terminated, the owner and/or pharmacy permit holder shall notify the Board of Pharmacy;
  - (C) change of ownership of the Pharmacy;
  - (D) change of address of the Pharmacy;
  - (E) permanent closing of the Pharmacy;
  - (F) Significant Quality-Related Events;
  - (G) the installation or removal of Automated Pharmacy Systems. Such notice must include, but is not limited to:
    - (-a-) the name and address of the Pharmacy;
    - (-b-) the location of the Automated Pharmacy System; and
    - (-c-) the identification of the responsible Pharmacist.

---

<sup>110</sup> As part of a quality assurance program designed to prevent and detect drug diversion, the Pharmacist-in-Charge, if the practice setting warrants, may also consider implementing diversion prevention and detection policies and procedures that address the following: periodic reviews of employee access to any secure controlled substance storage areas, which may include:

- alarm codes and lock combinations;
- passwords; and
- keys and access badges.

<sup>111</sup> If states require the Pharmacist-in-Charge or other Person in charge of the Pharmacy to submit information regarding the separation of employment of licensees, especially in circumstances of suspected or confirmed abuse, theft, or diversion of Drugs, states should also be aware of confidentiality and employment laws that may restrict the release of information and be cautioned that the release of such information may create a liability for the reporting Pharmacy.

In instances where the Pharmacist-in-Charge and the owner and/or pharmacy permit holder are the same person and that person is no longer employed or designated as the Person in charge, then the Board must take action to cease operation of the Pharmacy.

- (-d-) Such notice must be must occur prior to the installation or removal of the system.
  - (iv) Making or filing any reports required by state or federal laws and rules.
  - (v) Reporting any theft, suspected theft, diversion, or other Significant Loss of any Prescription Drug within one business day of discovery to the Board of Pharmacy and as required by Drug Enforcement Administration (DEA) or other State or federal agencies for Prescription Drugs and controlled substances.
  - (vi) Responding to the Board of Pharmacy regarding any minor violations brought to his or her attention.
- (3) The Pharmacist-in-Charge shall be assisted by a sufficient number of Pharmacists, Certified Pharmacy Technicians, and Certified Pharmacy Technician Candidates as may be required to competently and safely provide Pharmacy services.
  - (i) The Pharmacist-in-Charge shall maintain and file with the Board of Pharmacy, on a form provided by the Board, a current list of all Certified Pharmacy Technicians and Certified Pharmacy Technician Candidates assisting in the provision of Pharmacy services.
  - (ii) The Pharmacist-in-Charge shall develop or adopt, implement, and maintain written policies and procedures to specify the duties to be performed by Certified Pharmacy Technicians and Certified Pharmacy Technician Candidates. The duties and responsibilities of these personnel shall be consistent with their education, training, and experience and shall address the method and level of necessary supervision specific to the practice site. These policies and procedures shall, at a minimum, specify that Certified Pharmacy Technicians and Certified Pharmacy Technician Candidates are not assigned duties that may be performed only by a Pharmacist. Such policies and procedures shall also specify that Certified Pharmacy Technician Candidates shall not be assigned duties that may be performed only by Certified Pharmacy Technicians.
  - (iii) The Pharmacist-in-charge shall develop or adopt, implement, and maintain a Certified Pharmacy Technician training program that is site-specific to the practice setting of which he or she is in charge for all individuals employed by the Pharmacy who will assist in the Practice of Pharmacy. The Pharmacist-in-Charge shall utilize a Certified Pharmacy Technician training manual as part of the training program. The Pharmacist-in-Charge shall be responsible for maintaining a record of all Certified Pharmacy Technicians and Certified Pharmacy Technician Candidates successfully completing a site-specific training program and an objective assessment mechanism. The Pharmacist-in-Charge shall attest to the Board of Pharmacy, in a timely manner, those persons who, from time to time, have met the training requirements necessary for licensure by the Board.<sup>112</sup>
- (b) Professional Performance Evaluation
 

Each Pharmacist who performs any of the acts described within the definition of “Practice of Pharmacy” is responsible for ensuring that he or she is the subject of a Professional Performance Evaluation at least once each year. Each Pharmacy is responsible for ensuring that every Pharmacist who practices at the Pharmacy for more than 40 hours during any twelve (12)-month period and who performs any of the acts described within the definition of “Practice of Pharmacy” is the subject of a Professional Performance Evaluation at least once each year.
- (c) If any action of the Pharmacy is deemed to contribute to or cause a violation of any provision of this section, the Board may hold the owner and/or Pharmacy permit holder responsible and/or absolve the Pharmacist-in-Charge from the responsibility of that action.

---

<sup>112</sup> All training programs should be subject to approval by the Board of Pharmacy.

## Section 4. Prescription Drug Order Processing.

### (a) Prescription Drug Order

A Prescription Drug Order shall contain the following information at a minimum:

- (1) full name, date of birth, and street address of the patient;
- (2) name, prescribing Practitioner's license designation, address, and, if required by law or rules of the Board, DEA registration number of the prescribing Practitioner;
- (3) date of issuance;
- (4) name, strength, dosage form, and quantity of Drug prescribed;
- (5) directions for use;
- (6) refills authorized, if any;
- (7) if a written Prescription Drug Order, prescribing Practitioner's signature;
- (8) if an electronically transmitted Prescription Drug Order, prescribing Practitioner's electronic or digital signature;
- (9) if a hard copy Prescription Drug Order generated from electronic media, prescribing Practitioner's electronic or manual signature. For those with electronic signatures, such Prescription Drug Orders shall be applied to paper that utilizes security features<sup>113</sup> that will ensure the Prescription Drug Order is not subject to any form of copying and/or alteration.

### (b) Manner of Issuance of a Prescription Drug Order

A Prescription Drug Order, to be valid, must be issued for a legitimate medical purpose by a Practitioner acting within the course of legitimate professional practice. The responsibility for the proper prescribing and Dispensing of controlled substances is upon the prescribing Practitioner, but a corresponding responsibility rests with the Pharmacist who fills the prescription.<sup>114</sup>

- (1) A Prescription Drug Order must be communicated to a Pharmacist, or when recorded in such a way that the Pharmacist may review the Prescription Drug Order as transmitted, to a Pharmacy Intern or a Certified Pharmacy Technician, in a licensed Pharmacy. This may be accomplished in one of the following ways. A Prescription Drug Order, including that for a controlled substance listed in Schedules II through V, may be communicated in written form. A Prescription Drug Order, including that for a controlled substance listed in Schedules III through V, and, in certain situations, that for a controlled substance listed in Schedule II, may be communicated orally (including telephone voice communication)<sup>115</sup> or issued electronically.<sup>116</sup>
- (2) The Pharmacist shall not dispense a Prescription Drug if the Pharmacist knows or reasonably should know that the Prescription Drug Order was issued solely on the basis of an Internet-based questionnaire, an Internet-based consultation, or a telephonic consultation, all without a valid Patient-Practitioner relationship.
- (3) If communicated orally, the Prescription Drug Order shall be immediately reduced to a form by the Pharmacist, the Pharmacy Intern, or the Certified Pharmacy Technician that may be maintained for the time required by laws or rules.

---

<sup>113</sup> Examples of security features for prescription paper include those that prevent copying, such as hidden background words or darker-colored areas of the paper (which, when photocopied appear as black), those that prevent adulteration, such as solvent dye and brownstain features, and those that verify authenticity, such as the incorporation of fluorescent threads or watermarks.

<sup>114</sup> While Pharmacists have a corresponding responsibility to ensure that a controlled substance is Dispensed only pursuant to a valid Prescription Drug Order written for a legitimate medical purpose, this should not impede patients from receiving legitimately prescribed controlled substances or non-controlled substances, as patient care should be the primary consideration.

<sup>115</sup> Policies and procedures should also provide guidance for properly identifying agents of the prescribing Practitioner who are trained and competent in communicating Prescription Drug Orders.

<sup>116</sup> Electronically transmitted prescriptions should be transmitted from prescriber to Pharmacy with no intervening Persons making illegal alterations that may be considered as engaging in the Practice of Pharmacy without the authority to do so or without being Licensed to do so to such prescriptions. Evolving technologies and systems have alleviated previous concerns regarding the routes by which electronic prescriptions are transmitted, but any attempts to return to illegal prescription altering practices will be halted.

- (4) A Prescription Drug Order for a Schedule II controlled substance may be communicated orally only in the following situations and/or with the following restrictions. Otherwise, a Prescription Drug Order for a Schedule II controlled substance must be communicated in written form or issued electronically.
- (i) A Prescription Drug Order for a Schedule II controlled substance may be communicated by the Practitioner or the Practitioner's agent by way of Electronic Transmission, provided the original written, signed Prescription Drug Order is presented to the Pharmacist for review prior to the actual Dispensing of the controlled substance, except as noted in paragraph (ii) or (iii) of this Section 3(b)(3). The original, written Prescription Drug Order shall be maintained in accordance with state and federal record-keeping requirements.
  - (ii) In the case of an Emergency Situation, a Prescription Drug Order for a Schedule II controlled substance may be communicated by the Practitioner orally, provided that:
    - (A) the quantity prescribed and Dispensed is limited to the amount adequate to treat the patient during the emergency period (Dispensing beyond the emergency period must be pursuant to a Prescription Drug Order either written and signed or electronically issued by the prescribing Practitioner);
    - (B) the orally communicated Prescription Drug Order shall be immediately reduced to writing by the Pharmacist, Pharmacy Intern, or Certified Pharmacy Technician, if necessary, and shall contain the information required by state and federal law;
    - (C) if the prescribing Practitioner is not known to the Pharmacist, Pharmacy Intern, or Certified Pharmacy Technician, he or she must make a reasonable effort to determine that the oral authorization came from a registered Practitioner, which may include a callback to the Practitioner using the Practitioner's phone number as listed in the telephone directory and/or other good faith efforts to ensure his or her identity; and
    - (D) within seven days after authorizing an emergency oral Prescription Drug Order, the Practitioner shall cause a written Prescription Drug Order for the emergency quantity prescribed to be delivered to the Dispensing Pharmacist. The Prescription Drug Order shall have written on its face "Authorization for Emergency Dispensing," and the date of the orally transmitted Prescription Drug Order. The written Prescription Drug Order may be delivered to the Pharmacist in person or by mail, but if delivered by mail, it must be postmarked within the seven (7)-day period. Upon receipt, the Dispensing Pharmacist shall attach this written Prescription Drug Order to the emergency oral Prescription Drug Order, which had earlier been reduced to writing. The Pharmacist shall notify the nearest office of the DEA if the prescribing Practitioner fails to deliver a written Prescription Drug Order.
  - (iii) The prescribing Practitioner may authorize his or her agent to communicate a Prescription Drug Order orally or by way of Electronic Transmission via facsimile to a Pharmacist, Pharmacy Intern, or Certified Pharmacy Technician in a licensed Pharmacy, provided that the identity of the transmitting agent is included in the order. In an Institutional Facility, the prescribing Practitioner's agent must be authorized by and in accordance with written policies and procedures of the Facility and applicable state and federal laws.
- (5) A Prescription Drug Order for a Schedule II narcotic substance to be Compounded for the direct Administration to a patient by parenteral, intravenous, intramuscular, subcutaneous, or intraspinal infusion may be communicated by the Practitioner or the Practitioner's agent to the Home Infusion Pharmacy via facsimile. The hard copy of such faxed prescription serves as the original, written Prescription Drug Order and it shall be maintained in accordance with state and federal record-keeping requirements.
- (6) A Prescription Drug Order for a Schedule II controlled substance for a resident of a Long-Term Care Facility may be communicated by the Practitioner or the Practitioner's agent via facsimile. The hard

copy of such faxed prescription serves as the original, written Prescription Drug Order and it shall be maintained in accordance with state and federal record-keeping requirements.

- (7) All Prescription Drug Orders for a Schedule III-V controlled substance communicated by way of Electronic Transmission via facsimile shall:
  - (i) be transmitted to a Pharmacist, Pharmacy Intern, or Certified Pharmacy Technician in a licensed Pharmacy of the patient's choice;
  - (ii) identify the transmitter's phone number or any other suitable means to contact the transmitter for verbal and/or written confirmation, the time and date of transmission, and the identity of the Pharmacy intended to receive the transmission, as well as any other information required by federal or state law;
  - (iii) be transmitted by an authorized Practitioner or his or her designated agent; and
  - (iv) be deemed the original Prescription Drug Order, provided it meets the requirements of this subsection.
- (8) All Prescription Drug Orders for a Schedule II-V controlled substance issued and processed electronically shall be in compliance with existing federal or state laws and rules.
- (9) The Pharmacist shall exercise professional judgment regarding the accuracy, validity, and authenticity of the Prescription Drug Order issued electronically or by facsimile to ensure it is consistent with existing federal or state laws and rules.
- (10) All electronic equipment for receipt of Prescription Drug Orders issued electronically or by facsimile shall be maintained so as to ensure against unauthorized access.
- (11) Persons other than those bound by a confidentiality agreement shall not have access to Pharmacy records containing Protected Health Information concerning the Pharmacy's patients.

(c) Transfer of a Prescription Drug Order

Pharmacies utilizing automated data-processing systems shall satisfy all information requirements of a manual mode for Prescription Drug Order transferal, except as noted below for those Pharmacies accessing a common electronic file. The transfer of original Prescription Drug Order information for the purpose of refill Dispensing is permissible between Pharmacies subject to the following requirements:

- (1) The information for a prescription, other than for a controlled substance,<sup>117</sup> is communicated directly between Pharmacists or Certified Pharmacy Technicians and the transferring Pharmacist or Certified Pharmacy Technician records the following information:
  - (i) write the word "VOID" on the face of the invalidated Prescription Drug Order;
  - (ii) record on the reverse side of the invalidated Prescription Drug Order the name and address of the Pharmacy to which it was transferred and the name of the Pharmacist or Certified Pharmacy Technician receiving the Prescription Drug Order;
  - (iii) record the date of the transfer and the name of the Pharmacist or Certified Pharmacy Technician transferring the information; and
  - (iv) the computer record shall reflect the fact that the original Prescription Drug Order has been voided and shall contain all the other information required above.
- (2) The Pharmacist or Certified Pharmacy Technician receiving the transferred Prescription Drug Order information shall reduce to writing the following:
  - (i) Write the word "TRANSFER" on the face of the transferred Prescription Drug Order.
  - (ii) Provide all information required to be on a Prescription Drug Order pursuant to state and federal laws and rules, and include:
    - (A) date of issuance of original Prescription Drug Order;
    - (B) original number of refills authorized on original Prescription Drug Order;
    - (C) date of original Dispensing;
    - (D) number of valid refills remaining and date of last refill;

---

<sup>117</sup> According to 21 CFR §1306.25 (b)(1), the transfer of original prescription information for a controlled substance listed in Schedule III, IV, or V for the purpose of refill Dispensing must be communicated directly between two licensed Pharmacists.

- (E) Pharmacy's name, address, and original prescription number from which the Prescription Drug Order information was transferred; and
  - (F) name of transferring Pharmacist or Certified Pharmacy Technician.
- (iii) Systems providing for the electronic transfer of information shall not infringe on a patient's freedom of choice as to the provider of Pharmacist Care Services.
- (3) Both the original and transferred Prescription Drug Order shall be maintained for a period of five years from the date of last refill.
- (4) Pharmacies accessing a common electronic file or database used to maintain required Dispensing information are not required to transfer Prescription Drug Orders or information for Dispensing purposes between or among Pharmacies participating in the same common prescription file, provided, however, that any such common file shall contain complete records of each Prescription Drug Order and refill Dispensed, and, further, that a hard copy record of each Prescription Drug Order transferred or accessed for purposes of refilling shall be generated and maintained at the Pharmacy refilling the Prescription Drug Order or to which the Prescription Drug Order is transferred and shall protect against the illegal use or disclosure of Protected Health Information.
- (5) In an emergency, a Pharmacy may transfer original Prescription Drug Order information for a non-controlled substance to a second Pharmacy for the purpose of Dispensing up to a 72-hour supply without voiding the original Prescription Drug Order.
- (d) Drug Product Selection by the Pharmacist
  - (1) A Pharmacist Dispensing a Prescription Drug Order for a Drug Product prescribed by its brand name may select any Equivalent Drug Product provided that the Manufacturer or Distributor holds, if applicable, either an approved New Drug Application (NDA) or an approved Abbreviated New Drug Application (ANDA), unless other approval by law or from the Federal Food and Drug Administration is required.
  - (2) The Pharmacist shall not select an Equivalent Drug Product if the Practitioner instructs otherwise, either orally or in writing, on the Prescription Drug Order.
  - (3) The Pharmacist shall notify the patient or patient's agent if a Drug other than the brand name Drug prescribed is Dispensed.
  - (4) Whenever Drug Product selection is performed by a Pharmacist, the Pharmacist shall Dispense the Equivalent Drug Product in a container Labeled in accordance with Section 3(e) (Labeling).
- (e) Labeling
  - (1) All Drugs Dispensed to ambulatory or outpatients, including Drugs Dispensed by Practitioners shall have a Label affixed to the container in which such Drug is Dispensed. The Label shall include the following:<sup>118</sup>
    - (i) Critical Information for Patients – Critical information must appear on the Label with emphasis (highlighted or bolded), in a sans serif typeface (such as "arial"), minimum 12-point size, and in "sentence case." Field size and font size may be increased in the best interest of patient care.<sup>119</sup> Critical information text should never be truncated and shall include:
      - (A) patient name:
        - (-a-) legal name of the patient; or
        - (-b-) if patient is an animal, include the last name of the owner, name of the animal, and animal species;
      - (B) directions for use:

<sup>118</sup> Electronically transmitted prescriptions should be transmitted from prescriber to Pharmacy with no intervening Persons making illegal alterations that may be considered as engaging in the Practice of Pharmacy without the authority to do so or without being licensed to do so to such prescriptions. Evolving technologies and systems have alleviated previous concerns regarding the routes by which electronic prescriptions are transmitted, but any attempts to return to illegal prescription altering practices will be halted.

<sup>119</sup> Alternative-access methods may be utilized to address visual impairment in patients or caregivers.

- (-a-) directions for use as indicated by the prescriber and medication purpose/indication if included on Prescription Drug Order;<sup>120</sup> and
  - (-b-) language should be simplified, avoiding unfamiliar words and medical jargon; when applicable, use numeric instead of alphabetic characters;<sup>121</sup>
- (C) drug name:
  - (-a-) if written for a brand name and a generic Drug is Dispensed, include phrase “Generic for [brand name]” or similar wording;<sup>122</sup> and
  - (-b-) include Drug name suffixes, such as CD, SR, XL, XR, etc;
- (D) drug strength, expressed in the metric system whenever possible;
- (E) oral liquid medication dosage, expressed in milliliters; and
- (F) “use by” date:
  - (-a-) date after which Drug should not be used; not expiration date of Drug or expiration date of prescription;<sup>123</sup> and
  - (-b-) format as – “Use by: MM/DD/YY.”
- (ii) Important information for patients – Must appear on the Label but should not supersede critical information for patients and shall include: <sup>124</sup>
  - (A) Pharmacy name or Dispensing practitioner’s entity name;<sup>125</sup>
  - (B) Pharmacy telephone number;<sup>126</sup>
  - (C) prescriber name:
    - (-a-) format as – “Prescriber: [prescriber name]”;
  - (D) “fill date”;<sup>127</sup>
    - (-a-) format as – “Date filled: MM/DD/YY”;
  - (E) prescription number;
  - (F) Drug quantity:
    - (-a-) format as – “Qty: [number]”;
  - (G) number of remaining refills:
    - (-a-) format as – “Refills: [number remaining]” or “No refills,” using whole numbers only and managing partial fills through the Pharmacy record-keeping system;

<sup>120</sup> Boards of Pharmacy and licensees should recognize that “take as directed” may not provide sufficient information for the appropriate use of the medication. “Take as directed” is appropriate when specific directions are included on a unit-of-use package or Dispensed package or in situations when directions are not able to be included on the Label and the Pharmacist presents directions to the patient and documents that such directions were given. “Take as directed” should not be used in lieu of Patient Counseling.

It is understood that Prescription Drug Orders often do not include the indication for use.

<sup>121</sup> Consider adhering to the universal medication schedule (UMS). The UMS shifts medication-taking into four standardized time periods (morning, noon, evening, bedtime) and uses simplified language and formatting to promote understanding (eg, “take 1 tablet in the morning and 1 tablet at bedtime”).

<sup>122</sup> If an Interchangeable Product is Dispensed, include the phrase “interchangeable for [Reference Product].”

<sup>123</sup> Boards of Pharmacy may determine that this “use by” date does not apply to all Drugs (for example epinephrine auto-injectors) and may allow the Manufacturer’s expiration date to be used if the Drug is kept in the Manufacturer’s original, unopened packaging, provided that the Pharmacist uses professional judgement to assess the continued need for the Drug and counsels the patient on proper storage.

<sup>124</sup> Information traditionally included on the patient Label must continue to be maintained and safeguarded by the record-keeping system. Boards of Pharmacy should require that record-keeping systems prohibit any alteration or modification of these data unless an appropriate audit trail and justification exists. Record-keeping systems should also prohibit any deletion of information except in accordance with state and federal requirements for data management and retention.

<sup>125</sup> Boards of Pharmacy should recognize that some pharmacies “do business as” a name other than the corporate name.

<sup>126</sup> Include phone number of the Dispensing pharmacy, recognizing that a pharmacy providing shared services may be involved in the filling process; Boards of Pharmacy should not require more than one telephone number on the Label.

<sup>127</sup> “Fill date” and “use by” date should be the only dates appearing on the prescription Label. Other dates often found on Labels, such as the original and expiration dates of the prescription drug order can be misunderstood by patients and clutter the Label with unnecessary information.



- (H) written or graphic product description;
- (I) auxiliary information;<sup>128</sup>
- (J) any cautions and other provisions which may be required by federal or state law.
- (iii) The following additional information for patients – may appear on the label:
  - (A) bar codes;
  - (B) Pharmacy address; and
  - (C) store number.<sup>129</sup>

## Section 5. Record Keeping.

### (a) Patient Records<sup>130</sup>

- (1) A patient record system shall be maintained by all Pharmacies and dispensing Practitioners for patients for whom Prescription Drug Orders are Dispensed. The patient record system shall provide for the immediate retrieval of information necessary for the Dispensing Pharmacist to identify previously Dispensed Drugs at the time a Prescription Drug Order is presented for Dispensing, and be created and stored in a manner to protect against illegal use or disclosure of Protected Health

<sup>128</sup> Auxiliary information, including auxiliary Labels, should be evidence based, standardized, and demonstrated to complement the prescription Label.

<sup>129</sup> Boards of Pharmacy may consider utilizing these suggested Labeling formats provided below.

Pharmacy Name: Phone:	Date Filled: MM/DD/YY Rx No.:	Cautions:
<b>Purpose:</b> <b>Patient Q. Name</b> Prescriber: <b>Take 1 tablet in the morning and 2 tablets at bedtime.</b> <b>Drug Name and Strength</b> <b>Generic for:</b> <b>Discard after: MM/DD/YY</b>		Description:
	Qty: Refills:	

Pharmacy Name: Phone:	<b>Purpose:</b> <b>Take 1 tablet in the morning and 2 tablets at bedtime.</b>	
<b>Patient Q. Name</b> Rx No.: Date Filled: MM/DD/YY Prescriber: <b>Drug Name and Strength</b> <b>Generic for:</b> Qty: <b>Discard after: MM/DD/YY</b>	Cautions:	Description:
	Refills:	

<sup>130</sup> The Pharmacist should have access to clinical and laboratory data concerning each patient, and should monitor each patient's response to his or her Drug therapy. Any unexpected or untoward response should be reported to the prescribing physician. If the Pharmacist is not doing this monitoring, the identity of the health care provider that has assumed this responsibility should be documented in the patient's profile.

It is acceptable for new Prescription Drug Order data to be added to the patient profile, but original entries may not be altered.

Information. The Pharmacist shall make a reasonable effort to obtain, record, and maintain the following information:

- (i) full name of the patient for whom the Drug is intended;
  - (ii) street address and telephone number of the patient;
  - (iii) patient's age or date of birth;
  - (iv) patient's gender;
  - (v) a list of the patient's medications taken during the preceding 24 months; and
  - (vi) Pharmacist comments relevant to the individual's Drug therapy, including any other information peculiar to the specific patient or Drug.
- (2) The Pharmacist shall make a reasonable effort to obtain from the patient or the patient's agent and shall record any known allergies, Drug reactions, idiosyncrasies, and chronic conditions or disease states of the patient and the identity of any other Drugs, including over-the-counter Drugs or Devices currently being used by the patient which may relate to Prospective Drug Review.
  - (3) A patient record shall be maintained for a period of not less than ten years from the date of the last entry in the profile record. This record may be a hard copy or a computerized form.
  - (4) Protected Health Information may be used or disclosed as allowed under state and federal privacy rules.
  - (5) Significant Adverse Drug Reactions shall be reported to the Practitioner and an appropriate entry shall be made in the patient's record.

(b) Records of Dispensing/Delivery

- (1) Records of receipt, Dispensing, Delivery, Distribution, or other disposition of all Drugs or Devices are to be made and kept by Pharmacies for five years<sup>131</sup> and shall include, but not be limited to:
  - (i) quantity Dispensed for original and refills, if different from original;
  - (ii) date of receipt, Dispensing, Delivery, Distribution, or other disposition;
  - (iii) serial number (or equivalent if an institution);
  - (iv) the identification of the Pharmacist, Certified Pharmacy Technician, or Certified Pharmacy Technician Candidate responsible for Dispensing;
  - (v) name and Manufacturer of Drug Dispensed if Drug Product selection occurs; and
  - (vi) records of refills to date.
- (2) Pharmacies that ship medications by mail, common carrier, or other type of Delivery service shall implement a mechanism to verify that a patient or caregiver has actually received the Delivered medication.<sup>132</sup>

(c) Electronic Record Keeping

(1) Systems Policies and Procedures

An up-to-date policy and procedure manual shall be developed by the Pharmacist-in-Charge that explains the operational aspects of the computerized record-keeping system and shall:

- (i) include examples of all required output documentation provided by the computerized record-keeping system;
- (ii) outline steps to be followed when the computerized record-keeping system is not operational due to scheduled or unscheduled system interruption;
- (iii) outline regular and routine backup file procedure and file maintenance;
- (iv) outline audit procedures, personnel code assignments, and personnel responsibilities; and
- (v) provide a quality assurance mechanism for data entry validation.

---

<sup>131</sup> States should check federal laws and ensure that the number of years the state requires Dispensing records to be maintained are at least as many as federal requirements.

<sup>132</sup> States that require pharmacies that ship medication by mail, common carrier, or other type of Delivery service to implement a mechanism to verify that the patient or caregiver has actually received the Delivered medication may want to consider allowing the mechanism to include a waiver provision that allows the patient or caregiver to request Delivery without Verification and advises the patient or caregiver of the possible consequences of receiving Delivery without Verification.

(2) Data Storage and Retrieval.

- (i) the system shall have the capability of producing sight-readable information on all original and refill Prescription Drug Orders. The term “sight-readable” means that an authorized individual shall be able to examine the record and read the information from the cathode ray tube (CRT), microfiche, microfilm, printout, or other method acceptable to the Board of Pharmacy; and
- (ii) the system shall provide online retrieval (via CRT display or hard copy printout) of original Prescription Drug Order information. Such information shall include, but not be limited to, the Prescription Drug Order requirements and records of Dispensing as indicated in Section 3 of this Rule; and
- (iii) the storage and retrieval of refill information for original paper prescription orders for controlled substances in Schedule III and IV are subject to the following conditions:
  - (A) the system must provide online retrieval (via computer monitor or hard-copy printout) of original prescription order information for those prescription orders that are currently authorized for refilling. This shall include, but is not limited to, data such as the original prescription number; date of issuance of the original prescription order by the Practitioner; full name and address of the patient; name, address, and DEA registration number of the Practitioner; and the name, strength, dosage form, quantity of the controlled substance prescribed (and quantity dispensed if different from the quantity prescribed), and the total number of refills authorized by the prescribing Practitioner;
  - (B) the system must also provide online retrieval (via computer monitor or hard copy printout) of the current refill history for Schedule III or IV controlled substance prescription orders (those authorized for refill during the past six months). This refill history shall include, but is not limited to, the name of the controlled substance, the date of refill, the quantity Dispensed, the identification code, or name or initials of the Dispensing Pharmacist for each refill and the total number of refills Dispensed to date for that prescription order;
  - (C) Documentation of the fact that the refill information entered into the computer each time a Pharmacist refills an original paper, fax, or oral prescription order for a Schedule III or IV controlled substance is correct must be provided by the individual pharmacist who makes use of such an application. If such an application provides a hard copy printout of each day's controlled substance prescription order refill data, that printout shall be verified, dated, and signed by the individual Pharmacist who refilled such a prescription order. The individual Pharmacist must verify that the data indicated are correct and then sign this document in the same manner as he would sign a check or legal document ( eg, J.H. Smith, or John H. Smith). This document shall be maintained in a separate file at that Pharmacy for a period of two years from the Dispensing date. This printout of the day's controlled substance prescription order refill data must be provided to each Pharmacy using such a computerized application within 72 hours of the date on which the refill was Dispensed. It must be verified and signed by each Pharmacist who is involved with such dispensing. In lieu of such a printout, the Pharmacy shall maintain a bound logbook, or separate file, in which each individual Pharmacist involved in such Dispensing shall sign a statement (in the manner previously described) each day, attesting to the fact that the refill information entered into the computer that day has been reviewed by him or her and is correct as shown. Such a book or file must be maintained at the pharmacy employing such an application for a period of \_\_\_\_\_ years after the date of Dispensing the appropriately authorized refill;
  - (D) the electronic record-keeping system shall have the capability of producing a printout of any Prescription Drug Order data. The system shall provide a refill-by-refill audit trail for any specified strength and dosage form of any Drug. Such an audit trail shall be by

- printout, and include the name of the prescribing Practitioner, name and location of the patient, quantity Dispensed on each refill, date of Dispensing of each refill, name or identification code of the Dispensing Pharmacist, and unique identifier of the Prescription Drug Order; and
- (E) any facility maintaining centralized prescription records shall be capable of sending a requested printout to the Pharmacy within 48 hours.
- (iv) if a prescription is created, signed, transmitted, and received electronically, all records related to that prescription must be retained electronically and are subject to the following:
    - (A) records must be maintained electronically for \_\_\_\_\_ years from the date of their creation or receipt;
    - (B) records regarding controlled substances prescriptions must be readily retrievable from all other records. Electronic records must be easily readable or easily rendered into a format that a person can read;
    - (C) records required by this section part must be made available to the state and federal agencies upon request;
    - (D) if the Pharmacy discontinues or changes the electronic prescription service provider or transfers the electronic Prescription Drug Order records to another Pharmacy, the Pharmacy must ensure that the records are stored in a format that can be retrieved, displayed, and printed in a readable format; and
    - (E) digitally signed prescription records must be transferred or migrated with the digital signature.
- (3) Security
- To maintain the confidentiality of patient records, the system shall have adequate security and systems safeguards designed to prevent and detect unauthorized access, modification, or manipulation of patient records. Once the Drug has been Dispensed, any alterations in Prescription Drug Order data shall be documented, including the identification of the Pharmacist responsible for the alteration.
- (4) System Backup (Auxiliary Records Maintenance)
- (i) In the event of an unscheduled system interruption, sufficient patient data and Prescription Drug Order data should be available to permit reconstruction of such data within a two-hour time period for the Pharmacist to Dispense Drugs with sound professional judgment.
  - (ii) An auxiliary system shall be established for the documentation of refills if the automated data processing system is inoperative for any reason and to ensure that all refills are authorized by the original Prescription Drug Order and that the maximum number of refills is not exceeded.
  - (iii) The auxiliary system shall be in place to provide for the maintenance of all necessary patient Drug information (as outlined in this rule) until the automated system becomes operational. However, nothing in this section shall preclude the Pharmacist from using professional judgment for the benefit of a patient's health and safety.
  - (iv) When the automated system is restored to operation, the information regarding Prescription Drug Orders Dispensed and refilled during the inoperative period shall be entered into the automated system within 96 hours.
  - (v) Routine backup systems and procedures (hard copy, copy, disk, etc) shall be in place and operational to ensure against loss of patient data.
  - (vi) In the event that permanent Dispensing information is lost due to unscheduled system interruption, the Board of Pharmacy shall be notified within 24 hours.

## Section 6. Pharmacist Care Services.<sup>133</sup>

### (a) Prospective Drug Utilization Review (DUR)<sup>134</sup>

A Pharmacist shall obtain and review the patient records and medical history for each Prescription Drug Order for:

- (1) known allergies;
- (2) rational therapy contraindications;
- (3) reasonable dose, duration of use, and route of Administration, considering age, gender, and other patient factors;
- (4) reasonable directions for use;
- (5) potential or actual adverse Drug reactions;
- (6) Drug-Drug interactions;
- (7) Drug-food interactions;
- (8) Drug-disease contraindications;
- (9) therapeutic duplication;
- (10) proper utilization (including over- or under-utilization), and optimum therapeutic outcomes; and
- (11) abuse/misuse.

Upon recognizing any of the above, which may also include information obtained from reviewing data found in the prescription monitoring program, the Pharmacist shall take appropriate steps to avoid or resolve the problem which, if necessary, includes consultation with the Practitioner.

### (b) Patient Counseling<sup>135</sup>

- (1) Upon receipt of a Prescription Drug Order and following a review of the patient's record, a Pharmacist shall personally initiate discussion of matters which will enhance or optimize Drug therapy with each patient or caregiver of such patient. Such discussion shall be in Person, whenever practicable, or by telephone and shall include appropriate elements of Patient Counseling. Such elements may include the following:
  - (i) the name and description of the Drug;
  - (ii) the dosage form, dose, route of Administration, and duration of Drug therapy;
  - (iii) intended use of the Drug and expected action;
  - (iv) special directions and precautions for preparation, Administration, and use by the patient;
  - (v) common severe side or adverse effects or interactions and therapeutic contraindications that may be encountered, including their avoidance, and the action required if they occur;
  - (vi) techniques for self-monitoring Drug therapy;
  - (vii) proper storage and appropriate disposal method(s) of unwanted or unused medication;
  - (viii) prescription refill information;
  - (ix) action to be taken in the event of a missed dose; and
  - (x) Pharmacist comments relevant to the individual's Drug therapy, including any other information peculiar to the specific patient or Drug.

---

<sup>133</sup> Additional Pharmacist Care Services may include, but are not limited to, Patient assessment and evaluation; assessing health plan and medication eligibility and coverage; Administering Drugs, vaccines, or biologicals; performing Peer Review and peer consultations; reviewing, selecting, and developing formularies or plan /practice guidelines; consulting with other health care professionals; providing patient referrals; performing Medication Therapy Management; ordering lab tests; and performing lab tests as provided by State and Federal law.

<sup>134</sup> Pharmacists should be permitted to use computer software, if available, to accomplish this review.

<sup>135</sup> The intent of this Section is to require that the Pharmacist personally initiate counseling for all new Prescriptions and to exercise his or her professional judgment for refills. Situations may arise, however, where the prescriber specifically indicates that a patient should not be counseled. In such circumstances, it is the responsibility of the Pharmacist to provide the best patient care through appropriate communication with the prescriber and to document the reason(s) for not providing counseling to the patient.

- (2) An offer for Patient Counseling can be made by a Certified Pharmacy Technician or Certified Pharmacy Technician Candidate when it is not required by law or deemed necessary that it be done by the Pharmacist.
  - (3) Alternative forms of patient information shall be used to supplement Patient Counseling when appropriate. Examples include written information leaflets, pictogram labels, video programs, etc.
  - (4) Patient Counseling, as described above and defined in this Act, shall not be required for inpatients of a hospital or institution where other licensed health care professionals are authorized to Administer the Drug(s).
  - (5) A Pharmacist shall not be required to counsel a patient or caregiver when the patient or caregiver refuses such consultation.
- (c) Medication Adherence Monitoring Services and Intervention Programs
- Medication Adherence Monitoring Services and Intervention Programs designed to promote improved medication use behaviors, such as compliance and adherence, appropriate monitoring and self-reporting, increased patient knowledge, and improved therapy options, shall comply with established Guidelines for the Appropriate Use and Disclosure of Protected Health Information in Medication Adherence Monitoring Services and Patient Intervention Programs.
- (d) Collaborative Pharmacy Practice
- (1) Collaborative Pharmacy Practice Agreement
- A Pharmacist planning to engage in Collaborative Pharmacy Practice shall have on file at his or her place of practice the written Collaborative Pharmacy Practice Agreement. The initial existence and subsequent termination of any such agreement and any additional information the Board may require concerning the Collaborative Pharmacy Practice Agreement, including the agreement itself, shall be made available to the Board for review upon request. The Agreement may allow the Pharmacist, within the Pharmacist's Scope of Practice Pursuant to the Collaborative Pharmacy Practice Agreement, to conduct activities approved by the Practitioner, and as defined by law and by the Rules of the Board. The collaboration that the Practitioner agrees to conduct with the Pharmacist must be within the scope of the Practitioner's current practice. Patients or caregivers shall be advised of such agreement.
- (2) Contents
- The Collaborative Pharmacy Practice Agreement shall include:
- (i) identification of the Practitioner(s) and Pharmacist(s) who are parties to the Agreement;
  - (ii) the types of decisions that the Pharmacist is allowed to make;
  - (iii) a process for generating any necessary Medical Orders, including prescription orders, required to initiate allowed activities;
  - (iv) a method for the Practitioner to monitor compliance with the Agreement and clinical outcomes and to intercede where necessary;
  - (v) a description of the Continuous Quality Improvement Program used to evaluate effectiveness of patient care and ensure positive patient outcomes;
  - (vi) a provision that allows the Practitioner to override a Collaborative Practice decision made by the Pharmacist whenever he or she deems it necessary or appropriate;
  - (vii) a provision that allows either party to cancel the Agreement by written notification;
  - (viii) an effective date;
  - (ix) signatures of all collaborating Pharmacists and Practitioners who are party to the Agreement, as well as dates of signing; and
  - (x) a procedure for periodic review and renewal within a time frame that is clinically appropriate.
- (3) Amendments to a Collaborative Pharmacy Practice Agreement must be documented, signed, and dated.

(4) Documentation of Pharmacist Activities

Documentation of allowed activities must be kept as part of the patient's permanent record and be readily available to other health care professionals who are providing care to that patient and who are authorized to receive it. Documentation of allowed activities shall be considered Protected Health Information.

(e) Emergency-Use Dispensing

Prescribing and Dispensing Drugs for emergency-use pursuant to a Pharmacist-issued Prescription<sup>136</sup> and appropriate Patient Counseling, including but not limited to:

- (1) Opioid overdose reversal agents, such as naloxone;
- (2) Epinephrine;
- (3) Antidote kits;
- (4) Short-acting beta agonist inhalers; and
- (5) Medication-assisted Treatment for the purpose of initiating therapy for opioid use disorder. The Pharmacist must:
  - (i) obtain a DEA registration and a state controlled substance license or registration, if required; and
  - (ii) use professional judgment to assess the clinical appropriateness of the patient's request and the length of time until the patient obtains treatment from an authorized Practitioner.<sup>137</sup>

## Section 7. Continuous Quality Improvement Program.

(a) Continuous Quality Improvement Program

- (1) Compliance with this section may be considered by the Board as a mitigating factor in the investigation and evaluation of a Quality-Related Event (QRE).
- (2) Each Pharmacy shall establish a Continuous Quality Improvement (CQI) Program for the purpose of detecting, documenting, assessing, and preventing QREs. At a minimum, a CQI Program shall include provisions to:
  - (i) designate an individual or individuals responsible for implementing, maintaining, and monitoring the CQI Program, which is managed in accordance with written policies and procedures maintained in the pharmacy in an immediately retrievable form;
  - (ii) initiate documentation of QREs as soon as possible, but no more than three days, after determining their occurrence;
  - (iii) analyze data collected in response to QREs to assess causes and any contributing factors such as staffing levels, workflow, and technological support;
  - (iv) use the findings of the analysis to formulate an appropriate response and develop pharmacy systems and workflow processes designed to prevent QREs and increase good outcomes for patients;
  - (v) provide ongoing CQI education at least annually to all pharmacy personnel;
  - (vi) for those Persons utilizing a Drug formulary, a periodic review of such formulary shall be undertaken to ensure that appropriate medications are being offered/selected in the best interest of patients.
- (3) As a component of its CQI Program, each Pharmacy shall ensure that periodic meetings are held, at least annually, by staff members of the Pharmacy to consider the effects on quality of the Pharmacy system due to staffing levels, workflow, and technological support. Such meetings shall review data showing evidence of the quality of care for patients served by the Pharmacy and shall develop plans for improvements in the system of Pharmacy Practice so as to increase good outcomes for patients.

---

<sup>136</sup> Pharmacist may prescribe pursuant to specific statewide protocols or standing orders.

<sup>137</sup> It is contemplated that for long-term treatment, Pharmacists should be prescribing under a collaborative practice agreement rather than under an emergency-use provision.

- (4) Appropriately-blinded incidents of QREs shall be reported to a nationally recognized error reporting program designated by the Board.
- (5) **Quality Self-Audit**  
Each Pharmacy shall conduct a Quality Self-Audit at least quarterly to determine whether the occurrence of QREs has decreased and whether there has been compliance with preventative procedures, and to develop a plan for improved adherence with the CQI Program in the future. Each pharmacy shall conduct a Quality Self-Audit upon change of Pharmacist-in-Charge to familiarize that Person with the Pharmacy's CQI Program.
- (6) **Consumer Survey**  
As a component of its CQI Program, each Pharmacy should conduct a Consumer Survey of patients who receive pharmaceutical Products and services at the Pharmacy. A Consumer Survey should be conducted at least once per year. A statistically valid sampling technique may be used in lieu of surveying every patient. Each Pharmacy should use the results of its Consumer Survey to evaluate its own performance at a particular time and over a period of time.
- (7) **Protection from Discovery**<sup>138</sup>  
All information, communications, or data maintained as a component of a pharmacy CQI Program are privileged and confidential. This shall not prevent review of a pharmacy's CQI Program and records maintained as part of a system by the Board, pursuant to subpoena, as necessary to protect the public health and safety. All information, communications, or data furnished to any Peer Review Committee, and any findings, conclusions, or recommendations resulting from the proceedings of such committee, board, or entity are privileged. The records and proceedings of any Peer Review Committee, are confidential and shall be used by such committee, and the members thereof, only in the exercise of the proper functions of the committee and shall not be public records nor be available for court subpoena or for discovery proceedings. The disclosure of confidential, privileged Peer Review Committee information during advocacy, or as a report to the Board of Pharmacy, or to the affected Pharmacist or Pharmacy auxiliary personnel under review does not constitute a waiver of either confidentiality or privilege.
- (8) **Compliance with Subpoena**  
All persons shall comply fully with a subpoena issued by the Board for documents or information as otherwise authorized by law. The disclosure of documents or information under subpoena does not constitute a waiver of the privilege associated with a CQI Program. Failure to comply with the subpoena is grounds for disciplinary action against the Person by the appropriate licensing board.

## **Section 8. Shared Pharmacy Services.**

- (a) **General Requirements**<sup>139, 140</sup>
  - (1) The Pharmacy must possess a resident or nonresident permit issued by the Board prior to engaging in Shared Pharmacy Services.<sup>141</sup>

<sup>138</sup> Boards of pharmacy may have more or less authority to inspect CQI records, depending on state law. When authorizing the implementation of CQI Programs the extent of authority needed to obtain these materials must be determined.

<sup>139</sup> The Board may want to consider the extent to which this General Requirements Section is applicable to institutional-based Shared Pharmacy Services Pharmacies, as such application may be subject to interpretation of existing state and federal law governing Institutional Facilities.

<sup>140</sup> In order to ensure accountability, the Pharmacist-in-Charge of a Pharmacy engaging in Shared Pharmacy Services must possess a license to practice Pharmacy in all jurisdictions that he/she is engaging in such series until such a time in which provisions for multistate practice exist.

<sup>141</sup> Often the terms "licensure," "registration," and "permit" are used interchangeably throughout the *Model Act*. In the case of Shared Pharmacy Services Pharmacies that utilize Automated Pharmacy Systems, Boards may determine that it is appropriate to issue a permit for the Automated Pharmacy System but not for the physical site where the Automated Pharmacy System is located.



- (2) A Pharmacy may provide or utilize Shared Pharmacy Services only if the Pharmacies involved:
  - (i) have the same owner; or
  - (ii) have a written contract or agreement that outlines the services provided and the shared responsibilities of each party in complying with federal and state pharmacy laws and rules; and
  - (iii) share a common electronic file or technology that allows access to information necessary or required to perform Shared Pharmacy Services in conformance with the pharmacy act and the Board's rules.
- (3) A Pharmacy engaged in Shared Pharmacy Services shall comply with appropriate federal and state controlled substance registrations for each Pharmacy if controlled substances are maintained.
- (4) A Pharmacy engages in Shared Pharmacy Services shall notify the Board in writing within 10 days of a change of location, discontinuance of service, or closure of a Pharmacy.

(b) Operations

- (1) Pharmacies engaging in Shared Pharmacy Services, or a Pharmacist acting independently of a Pharmacy and participating in Shared Pharmacy Services shall:
  - (i) maintain records identifying, individually, for each Prescription Drug Order processed, the name of each Pharmacist or Pharmacy Intern who took part in the Drug Utilization Review, refill authorization, or therapeutic intervention functions performed at that Pharmacy and the name of any Certified Pharmacy Technician or Certified Pharmacy Technician Candidate if they assisted in any of those functions;
  - (ii) maintain records identifying individually, for each Prescription Drug Order filled or Dispensed, the name of each Pharmacist or Pharmacy Intern who took part in the filling, Dispensing, and counseling functions performed at that Pharmacy and the name of any Certified Pharmacy Technician or Certified Pharmacy Technician Candidate if they assisted in any of those functions;
  - (iii) report to the Board as soon as practical the results of any disciplinary action taken by another state's Board of Pharmacy involving Shared Pharmacy Services;
  - (iv) maintain a mechanism for tracking the Prescription Drug Order during each step of the processing and filling procedures performed at the Pharmacy;
  - (v) maintain a mechanism for the patient to identify all Pharmacies involved in filling the Prescription Drug Order; and
  - (vi) be able to obtain for inspection any required record or information within 72 hours of any request by the Board or its designee.
- (2) Notification to Patients
  - (i) Pharmacies engaging in Shared Pharmacy Services shall notify patients that their Prescription Drug Orders may be processed or filled by another Pharmacy unless the Prescription Drug is delivered to patients in Institutional Facilities where a licensed health care professional is responsible for administering the Prescription Drug to the patient.

(c) Drug Storage and Security

- (1) Drugs shall be stored in compliance with state and federal laws and in accordance with these Rules, including those addressing temperature, proper containers, and the handling of outdated drugs.
- (2) Drugs stored at Shared Pharmacy Services Pharmacies shall be stored in an area that is:
  - (i) separate from any other Drugs used by the health care facility; and
  - (ii) secured, so as to prevent access by unauthorized personnel.
- (3) Access to the area where Drugs are stored at the Shared Pharmacy Services Pharmacy must be limited to:
  - (i) Pharmacists, Certified Pharmacy Technicians, Certified Pharmacy Technician Candidates, or Pharmacy Interns who are employed by the Shared Pharmacy Services Pharmacy; or

- (ii) Personnel employed at the Institutional Facility or clinic where the Shared Pharmacy Services Pharmacy is located who:
  - (A) are licensed health care providers;
  - (B) are designated in writing by the Pharmacist-in-Charge or the Person responsible for the supervision and on-site operation of the facility where the Automated Pharmacy System is located; and
  - (C) have completed documented training concerning their duties associated with the Shared Pharmacy Services Pharmacy.
- (4) Shared Pharmacy Services Pharmacies shall have adequate security to:
  - (i) comply with federal and state laws and regulations; and
  - (ii) Protect the confidentiality and integrity of Protected Health Information.

(d) Policies and Procedures

- (1) Each participant in Shared Pharmacy Services shall jointly develop, implement, review, revise, and comply with joint policies and procedures for Shared Pharmacy Services. Each participant is required to maintain this portion of the joint policies and procedures that relate to that participant's operations. The policies and procedures shall:
  - (i) outline the responsibilities of each of the pharmacies;
  - (ii) include a list of the name, address, telephone numbers, and all license and permit numbers of the pharmacies involved in Shared Pharmacy Services; and
  - (iii) include policies and procedures for:
    - (A) notifying patients that their Prescription Drug Orders may be processed or filled by another Pharmacy and providing the name of the Pharmacy;
    - (B) protecting the confidentiality and integrity of Protected Health Information;
    - (C) dispensing Prescription Drug Orders when the filled Prescription Drug Order is not received or the patient comes in before the Prescription Drug Order is received;
    - (D) maintaining required manual or electronic records to identify the name, initials or identification code and specific activity or activities of each Pharmacist, Certified Pharmacy Technician, Certified Pharmacy Technician Candidate, or Pharmacy Intern who performed any Shared Pharmacy Services;
    - (E) complying with federal and state laws; and
    - (F) operating a Continuous Quality Improvement Program for Shared Pharmacy Services, designed to objectively and systematically monitor and evaluate the quality and appropriateness of patient care, pursue opportunities to improve patient care, and resolve identified problems.

(e) Individual Practice

- (1) Nothing in this Section shall prohibit an individual Pharmacist licensed in the state, who is an employee of or under contract with a Pharmacy, or a licensed Certified Pharmacy Technician, Certified Pharmacy Technician Candidate, or Pharmacy Intern, working under the supervision of the Pharmacist, from accessing that Pharmacy's electronic database from inside or outside the Pharmacy and performing the Prescription Drug Order processing functions permitted by the Pharmacy Act, if both of the following conditions are met:
  - (i) the Pharmacy establishes controls to protect the confidentiality and integrity of Protected Health Information; and
  - (ii) no part of the database is duplicated, downloaded, or removed from the Pharmacy's electronic database.

(f) Practice of Telepharmacy – Remote Dispensing Site Requirements <sup>142</sup>

A Remote Dispensing Site:

- (1) Shall submit an Application to the Board.
- (2) The Pharmacist-in-Charge of the Shared Pharmacy Services Pharmacy shall be responsible for all operations of the Remote Dispensing Site.
- (3) Shall have a written contract or agreement that outlines the services provided and the responsibilities of each party in complying with Federal and State pharmacy laws and rules.
- (4) The Pharmacist-in-Charge shall oversee monthly inspections, maintenance and reconciliation of all controlled substances, including maintaining a perpetual inventory for all Schedule II controlled substances.
- (5) A Pharmacist must be designated to be available within ( ) hours, in case of emergency.
- (6) A functioning video and audio communication system that provides for effective communication between the Shared Pharmacy Services Pharmacy and the Remote Dispensing Site personnel and patients, and their agents or caregivers, must be maintained. The system must provide an adequate number of views of the entire site, facilitate adequate pharmacist supervision, and allow the appropriate exchanges of visual, verbal, and written communications for patient counseling and other matters involved in the lawful transaction or Delivery of Drugs. The Remote Dispensing Site must retain a recording of the facility surveillance, excluding patient communications, for a minimum of ( ) days.
- (7) Unless a Pharmacist is present, a Remote Dispensing Site must not be open or its employees allowed access during times the Shared Pharmacy Services Pharmacy is closed or during a system outage.

**Section 9. Automated Pharmacy Systems.**

- (a) Automated Pharmacy Systems can be utilized in licensed pharmacies and Shared Pharmacy Services Pharmacies located within an Institutional Facility or clinic. A Pharmacist is not required to be physically present at the site of the Automated Pharmacy System if the system is supervised electronically by a Pharmacist. Automated Pharmacy Systems shall comply with the following provisions.
- (1) Documentation as to type of equipment, serial numbers, content, policies and procedures, and Shared Pharmacy Services Pharmacy location shall be maintained in the Pharmacy for review . Such documentation shall include, but is not limited to:
    - (i) name and address of the Pharmacy and the Shared Pharmacy Services Pharmacy where the Automated Pharmacy System (s) is being used;
    - (ii) Manufacturer’s name and model;
    - (iii) description of how the Automated Pharmacy System is used;
    - (iv) quality assurance procedures to determine continued appropriate use of the Automated Pharmacy System;
    - (v) policies and procedures for system operation, safety, security, accuracy, patient confidentiality, access, and malfunction; and
    - (vi) documentation evidencing that the Automated Pharmacy System has been tested prior to initial use and on a periodic basis at each location to ensure that the Automated Pharmacy System is operating properly.
  - (2) Automated Pharmacy Systems should be used only in settings where there is an established program of Pharmacist Care Services that ensures medication orders or Prescription Drug Orders

---

<sup>142</sup> For Boards of Pharmacy that have yet to add rules for the Practice of Telepharmacy and/or require more specificity, see Appendix F Model Rules for the Practice of Telepharmacy.

are reviewed by a Pharmacist in accordance with established policies and procedures and good Pharmacist Care Services.<sup>143</sup>

- (i) A Pharmacist shall be accessible to respond to inquiries or requests pertaining to Drugs Dispensed from the Automated Pharmacy System.<sup>144</sup>
  - (ii) Any Pharmacy that maintains an Automated Pharmacy System for the purposes of remote Dispensing to outpatients<sup>145</sup> shall maintain a video/auditory communication system to provide for effective communication between the patient and the Pharmacist; the video/auditory communication system shall allow for the appropriate exchange of oral and written communication and Patient Counseling; if the video/auditory communication system malfunctions, then all operations of the Automated Pharmacy System shall cease until the system is fully functional.
- (3) All policies and procedures must be maintained in the Pharmacy responsible for the Automated Pharmacy System and, if the Automated Pharmacy System is being used at a different location, at that location as well.
  - (4) Automated Pharmacy Systems shall have adequate security systems and procedures, evidenced by written policies and procedures<sup>146</sup>, to:
    - (i) prevent unauthorized access;
    - (ii) comply with federal and state regulations; and
    - (iii) prevent the illegal use or disclosure of Protected Health Information.
  - (5) Records and/or electronic data kept by Automated Pharmacy Systems shall meet the following requirements.
    - (i) All events involving the contents of the Automated Pharmacy System must be recorded electronically.
    - (ii) Records must be maintained by the Pharmacy and must be readily available to the Board. Such records shall include:
      - (A) identity of system accessed;
      - (B) identification of the individual accessing the system;
      - (C) type of Transaction;
      - (D) name, strength, dosage form, and quantity of the Drug accessed;
      - (E) name of the patient for whom the Drug was ordered; and
      - (F) such additional information as the Pharmacist-in-Charge may deem necessary.
  - (6) Access to and limits on access (eg, security levels) to the Automated Pharmacy System must be defined by policy and procedures and must comply with state and federal regulations.<sup>147</sup>
  - (7) The Pharmacist-in-Charge shall have the responsibility to:
    - (i) assign, discontinue, or change access to the system;

---

<sup>143</sup> Each state should determine whether or not the Dispensing of a “first dose” or an “emergency dose” may take place without prior order review by a Pharmacist but with appropriate security and patient medication management controls in place.

<sup>144</sup> In order to facilitate communication between the Pharmacy and the site where the Automated Pharmacy System is located, a Pharmacy should provide a toll-free telephone number so that the Pharmacist is accessible at all times the Automated Pharmacy System is operational.

<sup>145</sup> Although an “outpatient” generally refers to a Person who receives Drugs for use outside of an Institutional Facility, the definition of “outpatient” must be defined by each state. For example, although the *Model Act* classifies penal institutions as a type of Institutional Facility and therefore its inmates as inpatients, the Pharmacist is exempt from providing Patient Counseling. However, some states may consider inmates of penal institutions as outpatients and therefore should decide if a video/audio communication system is required in such facilities so that the Pharmacist is able to provide Patient Counseling.

<sup>146</sup> The use of Automated Pharmacy Systems requires written policies and procedures in place prior to installation to ensure safety, accuracy, security, and patient confidentiality and to define access and limits to access to equipment and medications.

<sup>147</sup> This section anticipates that decisions regarding which health care professionals may access the Automated Pharmacy System and the level of access allowed (eg, access to medications, access to patient profiles for viewing only, access to patient profiles for modification) will be left up to the individual(s) responsible for the Automated Pharmacy System; however, states may decide to take on this responsibility and define those who may have access to the system and the levels of access allowed.

- (ii) ensure that access to the medications comply with state and federal regulations;
- (iii) ensure that the Automated Pharmacy System is filled/stocked accurately and in accordance with established, written policies and procedures.
- (8) The filling/stocking of all medications in the Automated Pharmacy System shall be accomplished by qualified personnel under the supervision of a licensed Pharmacist.
- (9) A record of medications filled/stocked into an Automated Pharmacy System shall be maintained and shall include identification of the persons filling/stocking and checking for accuracy.<sup>148</sup>
- (10) All containers of medications stored in the Automated Pharmacy System shall be packaged and labeled in accordance with federal and state laws and regulations.
- (11) All aspects of handling controlled substances shall meet the requirements of all state and federal laws and regulations.
- (12) The Automated Pharmacy System shall provide a mechanism for securing and accounting for medications removed from and subsequently returned to the Automated Pharmacy System, all in accordance with existing state and federal law.<sup>149</sup>
- (13) The Automated Pharmacy System shall provide a mechanism for securing and accounting for wasted medications or discarded medications in accordance with existing state and federal law.

## **Section 10. Return and Reuse of Prescription Drugs.**

- (a) Prescription Drugs may only be returned and reused providing that the Prescription Drugs were packaged in:
  - (1) the original, sealed, and tamper-evident bulk, unit-of-use,<sup>150</sup> or unit dose packaging; or
  - (2) the Dispensing Pharmacy's original packaging that maintains the Product quality.
- (b) All returned packaging must indicate that the Prescription Drug's integrity and stability has been maintained.
- (c) All returned Prescription Drugs must be evaluated by appropriate Pharmacy staff to ensure such Prescription Drugs are not adulterated or misbranded.
- (d) A state-licensed Pharmacist must verify compliance with all of the above elements.

## **Section 11. Prescription Drug Repository Programs.**

- (a) Repository Programs must have written policies and procedures, which include at a minimum:
  - (1) Qualifications of acceptable medications for reuse. Such qualifications must include the following provisions:
    - (i) only non-controlled medications will be accepted;<sup>151</sup>
    - (ii) all medications will be inspected and determined to be:
      - (A) unadulterated;
      - (B) unexpired; and
      - (C) in unopened unit dose or manufacturer's tamper-evident original packaging, or otherwise approved by the Board of Pharmacy;
    - (iii) maintenance of a separate physical inventory;

---

<sup>148</sup> This section anticipates that states will allow non-Pharmacist personnel to fill/stock Automated Pharmacy Systems under a Pharmacist's supervision; however, the state may decide to only allow a Pharmacist to perform this function. Should the State allow non-Pharmacist personnel to perform this function, it should define the level of Pharmacist supervision necessary (eg, immediate, direct, or general).

<sup>149</sup> The State may require that each licensed Pharmacy or facility have in place written policies and procedures to address situations in which medications removed from the system remain unused and must be secured and accounted for.

<sup>150</sup> Unit-of-use is not intended to include co-mingled, multi-medication unit-of-use packages also known as compliance packs.

<sup>151</sup> Except for federally scheduled controlled substance medications that may be prescribed for substance use disorders and as allowed by federal and state laws and regulations.

- (iv) completion of a monthly expiration date review for all medications;
  - (v) prohibition of charging or accepting compensation for medications except for administrative or minimal dispensing fees;
  - (vi) Dispensing by a pharmacist or a practitioner within the practitioner's scope of practice; and
  - (vii) record keeping, including the source and dispensation of all medication.
- (2) A requirement that the patient receives notification that the medication is being Dispensed by a Repository Program.

## **Section 12. Disposal of Controlled Substances.<sup>152</sup>**

- (a) Any Persons legally authorized to possess controlled substances in the course of their professional practice or the conduct of their business shall dispose of such Drugs by the following procedures and in compliance with federal law:
- (1) The responsible individual shall send the Board of Pharmacy a list of the controlled substances to be disposed of, including the name(s) and quantity of the Drug(s) .
  - (2) The Board shall authorize and instruct the applicant to dispose of the controlled substances in one of the following manners:
    - (i) by Delivery to an agent of the Board of Pharmacy or the Board of Pharmacy office;
    - (ii) by destruction of the Drugs in the presence of a Board of Pharmacy officer, agent, inspector, or other authorized individual; or
    - (iii) by such other means as the Board of Pharmacy may determine to ensure that the Drugs do not become available to unauthorized Persons.

## **Section 13. Prepackaging.**

- (a) A Pharmacy may Prepackage Drugs under the following circumstances:
- (1) written policies and procedures have been developed that address the processes of Prepackaging within the Pharmacy;
  - (2) containers utilized for Prepackaging shall meet, as a minimum requirement, Class B container standards as referenced by USP;
  - (3) the Prepackaging processes are conducted under conditions that ensure the integrity of the Drug and under the direct supervision of a Pharmacist;
  - (4) the Prepackaged Drugs are labeled with the following components:
    - (i) Drug Name;
    - (ii) Drug Strength;
    - (iii) Pharmacy Control and Manufacturer lot number;
    - (iv) Name of the Manufacturer or Distributor of the Drug or the National Drug Code; and
    - (v) Beyond-Use Date, which shall be the Manufacturer's expiration date or one that is required under USP standards,<sup>153</sup> whichever is earlier;
  - (5) Records of all Prepackaging operations are maintained and include the following:
    - (i) the name (nonproprietary and proprietary name), strength, dosage form, quantity per container, and quantity of containers of the Drug being Prepackaged;
    - (ii) the name of the Manufacturer or Distributor of the Drug;
    - (iii) Pharmacy Control and Manufacturer lot number;
    - (iv) expiration date of the Drug according to the original Manufacturer or Distributor container and the Beyond-Use Date;

<sup>152</sup> Boards may give hospitals the authority to dispose of wasted quantities of controlled substances without prior authorization under specified conditions.

<sup>153</sup> See USP General Chapter <7> Labeling.

- (v) the name, initials, or identification codes of the Certified Pharmacy Technician or Certified Pharmacy Technician Candidate that Prepackaged the Drug and the name or initials of the Pharmacist that verified the appropriateness of the Prepackaged Drug; and
  - (vi) the date the Drug is Prepackaged.
- (6) All Drugs Prepackaged are stored at appropriate temperatures and under appropriate conditions in accordance with requirements, if any, in the Labeling of such Drugs, or with requirements in the current edition of an official compendium.
- (b) Pharmacies that store Drugs within an automated counting device or Automated Pharmacy System may, in place of the required Label, maintain records of lot numbers and Beyond-Use Dates that are required on the Label as long as it is fully traceable and is readily retrievable.
- (c) The Repackaging of Drugs shall follow applicable state and federal law.

#### **Section 14. Provision of Pharmacist Care Services Outside of a Licensed Pharmacy.**

- (a) In order for a Pharmacist to provide Pharmacist Care Services outside the premises of a licensed Pharmacy, an applicant shall:
  - (1) register/license with the Board(s);
  - (2) have appropriate security and protections in place to ensure the confidentiality of records or other patient-specific information;
  - (3) maintain such records in readily retrievable form; and
  - (4) follow the patient care process approved by the Board.<sup>154</sup>

#### **Section 15. Approval of Pharmacy Practice Initiatives.**

- (a) Application.<sup>155</sup>

An application for approval of a Pharmacy practice initiative that improves the quality of or access to Pharmacist Care Services, but which falls outside the scope of present regulations, shall be submitted to the Board and shall contain at least the following information:

  - (1) the name, address, telephone number, and the license number of the Pharmacist responsible for overseeing the initiative;
  - (2) the specific location and, if a Pharmacy, the Pharmacy name, address, telephone, and license number where the proposed Pharmacy practice initiative will be conducted; and
  - (3) a detailed summary of the proposed Pharmacy practice initiative, which includes:
    - (i) the goals and/or objectives of the proposed Pharmacy practice initiative;
    - (ii) a full explanation of the initiative and how it will be conducted;
    - (iii) the time frame for the Pharmacy practice initiative, including the proposed start date;
    - (iv) background information or literature review to support the proposal, if applicable;
    - (v) the rule(s) that will have to be waived in order to complete the Pharmacy practice initiative and a request to waive the rule(s); and
    - (vi) procedures to be used during the Pharmacy practice initiative to ensure that the public's health and safety are not compromised as a result of the rule waiver.
- (b) Approval by the Board.
 

The Board shall approve a Pharmacy practice initiative if it determines that:

  - (1) the Pharmacy practice initiative will improve the quality of or access to Pharmacist Care Services;

<sup>154</sup> It is anticipated that Boards use the *Pharmacists' Patient Care Process* approved in May 2014 by the Joint Commission of Pharmacy Practitioners.

<sup>155</sup> Boards may want to develop language addressing the time frame within which they will take action on an application for approval of a Pharmacy practice initiative.

- (2) the Pharmacy practice initiative will not adversely affect, directly or indirectly, the health, safety, or well-being of the public; and
- (3) the alternative measures to be taken, if any, are equivalent or superior to those prescribed in the part for which the rule waiver is requested.

The Board shall deny, Revoke, or refuse to renew an application for a Pharmacy practice initiative if the Board determines that the above requirements have not been met. In issuing an approval for a Pharmacy practice initiative, the Board may impose such terms and conditions it deems appropriate to carry out the purposes of Section 213(a)(14) of this Act and the rules adopted thereunder.

(c) Notification.

The Board shall notify the applicant in writing within 60 days of the Board's decision. If an approval is granted, the notification shall specify the period of time for which the approval and rule waiver will be effective and any conditions to be met by the applicant.

(d) Extension of Approval of Pharmacy Practice Initiatives.

A request for an extension of an approval of a Pharmacy practice initiative shall be submitted in writing at least (\_\_\_\_\_) days prior to the expiration date of the existing approval. Renewal requests shall contain the information specified in subsection (a). An approval of a Pharmacy practice initiative shall be renewed by the Board if the applicant continues to satisfy the Criteria contained in subsection (b) and demonstrates compliance with the alternative measures or conditions imposed at the time the original Pharmacy practice initiative was approved.

## **Section 16. Unprofessional Conduct.**

Unprofessional conduct shall include, but is not limited to, the following acts of a Pharmacist or Pharmacy:

- (a) the publication or circulation of false, misleading, or otherwise deceptive statements concerning the Practice of Pharmacy;
- (b) unreasonably refusing to Compound or Dispense Prescription Drug Orders that may be expected to be Compounded or Dispensed in Pharmacies by Pharmacists;
- (c) attempting to circumvent the Patient Counseling requirements, or discouraging the patient from receiving Patient Counseling concerning their Prescription Drug Orders;
- (d) the illegal use or disclosure of Protected Health Information;
- (e) failure to maintain adequate records, systems, and security to protect against the illegal use or disclosure of Protected Health Information;
- (f) failure to maintain adequate records to account for disclosures of Protected Health Information;
- (g) selling, giving away, or otherwise disposing of accessories, chemicals, or Drugs or Devices found in illegal Drug traffic when the Pharmacist knows or should have known of their intended use in illegal activities;
- (h) engaging in conduct likely to deceive, defraud, or harm the public, or demonstrating a willful or careless disregard for the health, welfare, or safety of a patient, or engaging in conduct which substantially departs from the Standards of Care ordinarily exercised by a Pharmacist, with proof of actual injury not having to be established;
- (i) selling a Drug for which a Prescription Drug Order from a Practitioner is required, without having received a Prescription Drug Order for the Drug;
- (j) willfully and knowingly failing to maintain complete and accurate records of all Drugs received, Dispensed, or disposed of in compliance with the Federal laws and regulations and State laws and rules;
- (k) obtaining any remuneration by fraud, misrepresentation, or deception, including, but not limited to, receiving remuneration for amending or modifying, or attempting to amend or modify, a patient's Pharmacist Care Services, absent a clear benefit to the patient, solely in response to promotion or marketing activities;
- (l) willfully and knowingly completing and submitting inaccurate due diligence questionnaires and attestation documents regarding the purchase or receipt of Drugs from Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors.



# Model Rules for Public Health Emergencies

## Section 1. Purpose and Scope.<sup>156</sup>

By the provision of these rules by the Board, the primary purpose of the section is to enable Pharmacists and Pharmacies to assist in the management and containment of a Public Health Emergency or similar crisis within the confines of a regulatory framework that serves to protect the welfare and health of the public.

## Section 2. Definitions.

- (a) “Declared Disaster Areas” are areas designated by state or federal authorities as those that have been adversely affected by a natural or man-made disaster and require extraordinary measures to provide adequate, safe, and effective health care for the affected population.
- (b) “Emergency Prescription Drug Order” means a standing Prescription Drug Order issued by the State Health Officer for Pharmacists to Dispense designated Prescription Drugs during a Public Health Emergency requiring mass Dispensing to expeditiously treat or provide prophylaxis to large numbers of Patients.<sup>157</sup>

---

<sup>156</sup> States may consider adding the following, more detailed language, which specifically addresses Drug Disposal and reporting requirements in the case of an emergency or disaster, to their emergency rules or guidelines:

### Disposal of Prescription Drugs in Pharmacies Affected by Certain Disasters

- (a) For Pharmacies that sustain flood and/or fire damage in the Prescription department or other damage resulting in an irrevocable loss of the Drug inventory, the entire Drug inventory, including Drugs awaiting pick up by Patients, becomes unfit for Dispensing. In such a case, an accurate record of Prescription Drug losses should be prepared by the Pharmacy.
- (b) For Pharmacies that experience a loss of power for an extended period of time, the Drug inventory must be evaluated for continued Product integrity using USP standards. For example, medications with labeling requiring storage at “controlled room temperature” must be kept at between 68° F and 77° F, with brief deviations of between 56° F and 86° F. Medication inventories found to have been stored outside of USP standards become unfit for Dispensing. In such a case, an accurate record of Prescription Drug losses should be prepared by the Pharmacy. For Pharmacies with questions on USP Product integrity standards, contact USP at 800/227-8772.

### Reporting of Theft or Loss of Controlled Substances During an Emergency or Disaster

- (a) In circumstances of theft by looting, burglary, etc, where evidence or witnesses indicate the medications were taken by someone, the nearest DEA Diversion Field Office must be notified by telephone, facsimile, or brief written message of the circumstances of the theft immediately upon discovery. In addition, the pharmacy must complete DEA Form 106 – Report of Theft or Loss of Controlled Substances, found at [www.deadiversion.usdoj.gov](http://www.deadiversion.usdoj.gov), to formally document the actual circumstances of the theft and the quantity of controlled substances involved, once this information has been conclusively determined.
- (b) In circumstances of damage or where drugs were irrevocably lost to flooding or other circumstance, such information must be reported on DEA Form 41 – Registrants Inventory of Drugs Surrendered, found at [www.deadiversion.usdoj.gov](http://www.deadiversion.usdoj.gov).
- (c) The amount stolen or lost may need to be calculated by taking the most recent controlled substances inventory, adding the amount purchased since that date, then subtracting the amount dispensed and distributed since that date. Absent a calculated amount, a best estimate should be reported.

### Disposal of Prescription Drugs Irrevocably Lost in an Emergency or Disaster

- (a) Controlled Substances  
Reverse Distributors, either individually or in concert with other contractors, are equipped to dispose of controlled substances. Contact your primary distributor for their recommendations for a reverse Distributor or contact a reverse Distributor directly.
- (b) Contaminated Medical Debris  
Non-controlled substance Prescription Drugs and Devices contaminated with flood water or other contaminants should be disposed of using a medical waste transportation, processing, and disposal system vendor. Such vendors must be licensed by the state.
- (c) Hazardous Debris  
Materials are deemed hazardous if they are ignitable, corrosive, toxic, or reactive. Prescription Drugs considered hazardous include, but are not limited to, epinephrine, nicotine, nitroglycerin, physostigmine, reserpine, selenium sulfide, chloral hydrate, and many chemotherapy agents, such as cyclophosphamide, chlorambucil, and daunomycin. Other hazardous items that might be found in a Pharmacy include paints, varnishes and thinners, alcohol, batteries, mercury thermometers, and blood pressure cuffs. It is recommended that Pharmacies handle all contaminated Prescription medications as hazardous debris and dispose of it using a hazardous waste collection and disposal company. These companies must be licensed by the state.
- (d) Commercial Waste  
Over-the-counter Drugs and other store shelf material may be disposed of in the commercial waste stream.

<sup>157</sup> Boards may consider identifying the official who has authority to issue an “Emergency Prescription Drug Order” and reviewing this on a regular basis.

- (c) “Mobile Pharmacy” means a Pharmacy that is self-propelled or movable by another vehicle that is self-propelled.
- (d) “Public Health Emergency” means an imminent threat or occurrence of an illness or health condition caused by terrorism, bioterrorism, epidemic or pandemic disease, novel and highly fatal infectious agent or biological toxin, or natural or man-made disaster, that poses a substantial risk of a significant number of human fatalities or incidents of permanent or long-term disability that is beyond the capacity of local government or nongovernmental organizations to resolve.
- (e) “State of Emergency” means a governmental declaration, usually issued as a result of a Public Health Emergency, that may suspend certain normal functions of government, alert citizens to alter their normal behaviors, and/or direct government agencies to implement emergency preparedness plans.
- (f) “Temporary Pharmacy Facility” means a facility established as a result of a Public Health Emergency or State of Emergency to temporarily provide Pharmacy services within or adjacent to Declared Disaster Areas.

### **Section 3. Emergency Prescription Drug Order.**

- (a) For the duration of a State of Emergency issued due to a Public Health Emergency, a Pharmacist may Dispense a Prescription Drug pursuant to an Emergency Prescription Drug Order if the Pharmacist:
  - (1) performs, to the extent possible, a Prospective Drug Utilization Review (DUR) and Patient Counseling in accordance with these rules;<sup>158</sup>
  - (2) reduces the information to a form that may be maintained for the time required by law or rule, indicates it is an “Emergency Prescription Drug Order,” and files and maintains the record as required by state and federal law.

### **Section 4. Public Health Emergency Refill Dispensing.**

- (a) For the duration of the State of Emergency issued due to a Public Health Emergency in the affected state and in other states engaged in disaster assistance pursuant to a governmental declaration or rule of the Board, a Pharmacist may Dispense a refill of a Prescription Drug, not to exceed a thirty (30)-day supply, without Practitioner authorization if:<sup>159</sup>
  - (1) in the Pharmacist’s professional judgment, the Prescription Drug is essential to the maintenance of the Patient’s life or to the continuation of therapy;
  - (2) the Pharmacist makes a good faith effort to reduce the information to a form that may be maintained for the time required by law or rule, indicates it is an “Emergency Refill Prescription,” and maintains the record as required by state and federal law, as well as state and federal disaster agencies for consideration for possible reimbursement programs implemented to ensure continued provision of care during a disaster or emergency; and
  - (3) the Pharmacist informs the Patient or the Patient’s agent at the time of Dispensing that the Prescription Drug is being provided without the Practitioner’s authorization and that authorization of the Practitioner is required for future refills.
- (b) For the duration of the State of Emergency, in an effort to provide patients with the best possible care in light of limited Drug availability and/or limited information on patients’ current Drug therapy, a Pharmacist may initiate or modify Drug therapy and Dispense an amount of such Drug to accommodate a patient’s health care needs until that patient may be seen by a Practitioner. Pharmacists performing such

<sup>158</sup> Although these services are important, in times of a disaster or emergency, it may not be possible to perform a Prospective Drug Review or provide counseling on Dispensed Drugs.

<sup>159</sup> Boards may consider contacting the US Drug Enforcement Administration ahead of time to ensure these provisions are applicable to controlled substances.

activities must utilize currently accepted Standards of Care when initiating or modifying Drug therapy. These activities may be undertaken if:

- (1) in the Pharmacist's professional judgment, the Prescription Drug is essential to the maintenance of the Patient's life or to the continuation of therapy;
  - (2) the Pharmacist makes a good faith effort to reduce the information to a form that may be maintained for the time required by law or rule, indicates that drug therapy has been initiated or modified due to a disaster or emergency, and maintains the record as required by state and federal law; and<sup>160</sup>
  - (3) the Pharmacist informs the Patient or the Patient's agent at the time of Dispensing that the Prescription Drug is being provided without the Practitioner's authorization and that authorization of the Practitioner is required for future refills.
- (c) The Practitioner and Pharmacist shall not incur any liability as a result of the performance of these activities in good faith pursuant to this section.

## **Section 5. Temporary Recognition of Nonresident Licensure.**

- (a) When a State of Emergency is declared due to a Public Health Emergency:
- (1) a Pharmacist not licensed in this State, but currently licensed in another state, may Dispense Prescription Drugs in areas affected by the Declared Disaster during the time that the State of Emergency exists if:
    - (i) the Board can verify current licensure in good standing of the Pharmacist directly with the state or indirectly via a third-party verification system; and<sup>161</sup>
    - (ii) the Pharmacist is engaged in a legitimate relief effort.
  - (2) a Certified Pharmacy Technician, Certified Pharmacy Technician Candidate, or Pharmacy Intern not licensed in this State, but currently licensed in another state, may assist the Pharmacist in Dispensing Prescription Drugs in affected Disaster Areas during the time that the State of Emergency exists if:
    - (i) the Board can verify current licensure in good standing of the Certified Pharmacy Technician, Certified Pharmacy Technician Candidate, or Pharmacy Intern directly with the state or indirectly via a third-party verification system; and
    - (ii) the Certified Pharmacy Technician, Certified Pharmacy Technician Candidate, or Pharmacy Intern is engaged in a legitimate relief effort.
  - (3) a Wholesale Drug Distributor not licensed in this State, but currently licensed in another state, may Distribute Prescription Drugs in affected Disaster Areas during the time that the State of Emergency exists if:
    - (i) the Board can verify current licensure in good standing of the Wholesale Drug Distributor directly with the state or indirectly via a third-party verification system; and
    - (ii) the Wholesale Drug Distributor is engaged in a legitimate relief effort.
  - (4) the temporary recognition of nonresident licensure or registration shall cease with the termination of the State of Emergency.

---

<sup>160</sup> Boards should be cognizant that state and federal disaster agencies, to ensure continued provision of care during disasters or emergencies, have programs that consider reimbursement requests for medication providers and may request Board assistance in the dispersal of funds. Records of dispensing will likely be needed for possible reimbursement consideration. In addition, records may also be used for post-event evaluation of care.

<sup>161</sup> If the information cannot be verified directly by the state Board of Pharmacy in which the nonresident pharmacist is licensed, the NABP Disciplinary Clearinghouse may be utilized to verify that a nonresident pharmacist has not had disciplinary action taken against his or her license.

## Section 6. Temporary Pharmacy Facilities or Mobile Pharmacies.

- (a) Pharmacies located in Declared Disaster Areas, nonresident Pharmacies, and Pharmacies licensed in another state but not licensed in this State, if necessary to provide Pharmacy services during a State of Emergency, may arrange to temporarily locate or relocate to a Temporary Pharmacy Facility or Mobile Pharmacy if the Temporary Pharmacy Facility or Mobile Pharmacy:<sup>162</sup>
  - (1) is under the control and management of the Pharmacist-in Charge or designated supervising Pharmacist;
  - (2) is located within the Declared Disaster Area or affected areas;
  - (3) notifies the Board of its location;<sup>163</sup>
  - (4) is properly secured to prevent theft and diversion of Drugs;
  - (5) maintains records in accordance with laws and regulations of the state in which the disaster occurred; and
  - (6) ceases the provision of services with the termination of the State of Emergency, unless it is successfully licensed by the Board of Pharmacy in accordance with Article V of this Act.
- (b) The Board, in accordance with Board rules, shall have the authority to approve or disapprove Temporary Pharmacy Facilities and Mobile Pharmacies and shall make arrangements for appropriate monitoring and inspection of the Temporary Pharmacy Facilities and Mobile Pharmacies on a case-by-case basis. Approval of Temporary Pharmacy Facilities and Mobile Pharmacies will be based on the need, type, and scope of Public Health Emergency, as well as the ability of the Temporary Pharmacy Facilities or Mobile Pharmacies to comply with state and federal drug law.
- (c) A Temporary Pharmacy Facility wishing to permanently operate at its temporary site must be licensed by the Board of Pharmacy in accordance with Article V of this Act.
- (d) Mobile Pharmacies, placed in operation during a State of Emergency, may not operate permanently, unless approved by the Board.<sup>164</sup>

---

<sup>162</sup> Boards may consider contacting the US Drug Enforcement Administration ahead of time to ensure that controlled substances may be delivered to and Dispensed from temporary or mobile pharmacy facilities.

<sup>163</sup> Boards may choose to require “approval” of a Temporary Pharmacy Facility or a Mobile Pharmacy, as opposed to requiring only “notification.” “Notification” may imply that the Board of Pharmacy has approved the location of the Temporary Pharmacy Facility or Mobile Pharmacy.

<sup>164</sup> Although many states do not allow the permanent or temporary licensure of Mobile Pharmacies, states that do allow the licensure of Mobile Pharmacies may consider implementing special requirements for permanent licensure; for example, a state may limit Mobile Pharmacies to operation only by nonprofit organizations and only in communities that are medically underserved.

# Model Rules for Nuclear/Radiologic Pharmacy

## Section 1. Purpose and Scope.

The Practice of Nuclear/Radiologic Pharmacy is hereby recognized as a specialty of Pharmacy practice, regulated by State Boards of Pharmacy. As such, the following model rules are included to address those areas specific or unique to this specialty practice. Nuclear/Radiologic Pharmacy Practice refers to a patient-oriented service that embodies the scientific knowledge and professional judgment required to improve and promote health through the assurance of the safe and efficacious use of radiopharmaceuticals and other Drugs.

## Section 2. Definitions.

- (a) "Authentication of Product History" means, but is not limited to, identifying the purchasing source, the ultimate fate, and any intermediate handling of any Component of a radiopharmaceutical.
- (b) "Internal Test Assessment" means, but is not limited to, conducting those tests of quality assurance necessary to ensure the integrity of the test.
- (c) "Nuclear Pharmacy" means a Pharmacy providing radiopharmaceutical services or, as provided in Section 3 of these Rules, an appropriate area of any Institutional Facility.
- (d) "Qualified Licensed Professional" means a non-Pharmacist individual (such as a physician, nurse, or technologist) who possesses a current state license, if applicable, and who has sufficient training and experience to safely handle and Dispense radiopharmaceuticals as defined by the respective requirements of [cite appropriate Nuclear Regulatory Commission (NRC) or Agreement State and State Board of Pharmacy law(s)].
- (e) "Qualified Nuclear Pharmacist" means a currently licensed Pharmacist in the State of practice, who is certified as a Nuclear Pharmacist by the State Board of Pharmacy or by a certification Board recognized by the State Board of Pharmacy, or who meets the following standards:
  - (1) Minimum standards of training for "authorized user status" of radioactive material [cite State Radiation Control Agency or NRC licensure guide].
  - (2) Completed a minimum of 200 contact hours of instruction in nuclear Pharmacy and the safe handling and the use of radioactive materials from a program approved by the State Board of Pharmacy, with emphasis in the following areas:
    - (i) radiation physics and instrumentation;
    - (ii) radiation protection;
    - (iii) mathematics of radioactivity;
    - (iv) radiation biology; and
    - (v) radiopharmaceutical chemistry.
  - (3) Attained a minimum of 500 hours of clinical nuclear Pharmacy training under the supervision of a qualified nuclear Pharmacist.
- (f) "Radiopharmaceutical Quality Assurance" means, but is not limited to, the performance of appropriate chemical, biological, and physical tests on potential radiopharmaceuticals and the interpretation of the resulting data to determine their suitability for use in humans and animals, including internal test assessment, authentication of Product history, and the keeping of proper records.
- (g) "Radiopharmaceutical Service" means, but shall not be limited to, the procurement, storage, handling, preparation, Labeling, quality assurance testing, Dispensing, Delivery, record keeping, and disposal of radiopharmaceuticals and other Drugs.
- (h) "Radiopharmaceuticals" are radioactive Drugs as defined by Food and Drug Administration and the \_\_\_\_\_ State Board of Pharmacy [cite appropriate law(s)].

### Section 3. General Requirements for Pharmacies Providing Radiopharmaceutical Services.

- (a) Nuclear Pharmacy License. A license to operate a Pharmacy providing radiopharmaceutical services shall only be issued to a Qualified Nuclear Pharmacist. All personnel performing tasks in the preparation and Distribution of radioactive Drugs shall be under the direct supervision of a Qualified Nuclear Pharmacist. A Qualified Nuclear Pharmacist shall be responsible for all operations of the Pharmacy and shall be in personal attendance at all times that the Pharmacy is open for business. In emergency situations when a Qualified Nuclear Pharmacist is not present, designated Qualified Licensed Professionals may have access to the licensed area. These individuals may prepare single doses of radiopharmaceuticals for the immediate emergency, and must document such activities.
- (b) Nuclear Pharmacies shall have adequate space and equipment, commensurate with the scope of services required and provided, meeting minimal space requirements established for all pharmacies in the State or as otherwise defined by the \_\_\_\_\_ State Board of Pharmacy.
- (c) The Nuclear Pharmacy area shall be secured from unauthorized personnel.
- (d) Nuclear Pharmacies shall maintain records of acquisition, inventory, and disposition of all radioactive Drugs and other radioactive materials in accordance with [cite appropriate Pharmacy and radiological control agency or NRC Statute(s)].
- (e) All pharmacies handling radiopharmaceuticals shall provide a radioactive storage and Product decay area. Detailed floor plans shall be submitted to the State Board of Pharmacy and the State Radiation Control Agency or NRC before approval of the license.
- (f) Radiopharmaceuticals are to be Dispensed only upon a Prescription Drug Order from a Practitioner authorized to possess, use, and Administer radiopharmaceuticals.
- (g) The permit to operate a Nuclear Pharmacy is conditioned upon an approved State Radiation Control Agency (RCA) or NRC license. Copies of the RCA or NRC inspection reports shall be made available upon request for Board inspection.
- (h) Labeling
  - (1) No radiopharmaceutical may be Dispensed unless a label is affixed to the immediate container bearing the following information:
    - (i) the standard radiation symbol;
    - (ii) the words "Caution – Radioactive Material"; and
    - (iii) the prescription number.
  - (2) No radiopharmaceutical may be Dispensed unless a label is affixed to the outer or Delivery container bearing the following information:
    - (i) the standard radiation symbol;
    - (ii) the words "Caution – Radioactive Material";
    - (iii) the radionuclide and chemical form;
    - (iv) the activity and date and time of assay;
    - (v) the volume, if in liquid form;
    - (vi) the requested activity and the calibrated activity;
    - (vii) the prescription number;
    - (viii) patient name or space for patient name. Where the patient's name is not available at the time of Dispensing, a 72-hour exemption is allowed to obtain the name of the patient. No later than 72 hours after Dispensing the radiopharmaceutical, the patient's name shall become a part of the Prescription Drug Order to be retained for a period of three years;
    - (ix) the name and address of the nuclear Pharmacy;
    - (x) the name of the Practitioner; and
    - (xi) the lot number of the prescription.

**Section 4. Other Requirements.**

All Nuclear/Radiologic Pharmacies shall also adhere to the principles outlined in the Rules for Pharmacist Care Services as these pertain to the practice of Nuclear Pharmacy.

# Model Rules for Compounded or Repackaged Pharmaceuticals

## Section 1. Purpose and Scope.

The purpose of this section is to ensure Compounded Pharmaceuticals are prepared and Dispensed according to practice and quality standards through the provision of: (1) Pharmacist Care Services; and (2) the preparation, Labeling, and Distribution of Compounded or Repackaged Pharmaceuticals by Pharmacies. These standards are intended to apply to all Sterile and nonsterile Compounded Pharmaceuticals, notwithstanding the location of the patient (eg, home, hospital, nursing home, hospice, doctor's office). All facilities and Practitioners engaging in Sterile and nonsterile Compounding or Repackaging shall practice in accordance with Federal law, these Rules, and the current United States Pharmacopeia–National Formulary (USP-NF), including but not limited to General Chapter <797> *Pharmaceutical Compounding – Sterile Preparations*, General Chapter <795> *Pharmaceutical Compounding – Nonsterile Preparations*, General Chapter <800> *Hazardous Drugs – Handling in Healthcare Settings*, and other applicable referenced general chapters. The procedures contained herein are considered to be the minimum current good compounding practices for the Compounding of Drug Products by State-licensed Pharmacies for Dispensing and/or Administration to humans or animals.<sup>165</sup>

## Section 2. Notification.

- (a) On an annual basis, and within 90 days of the beginning of the calendar year, all licensed Persons shall report to the NABP Information Sharing Network the information required by the “MEMORANDUM OF UNDERSTANDING ADDRESSING CERTAIN DISTRIBUTIONS OF COMPOUNDED HUMAN DRUG PRODUCTS BETWEEN THE [insert STATE BOARD OF PHARMACY OR OTHER APPROPRIATE STATE AGENCY] AND THE U.S. FOOD AND DRUG ADMINISTRATION.”
- (b) Upon request from the Board, all licensed Persons shall report to the Board of Pharmacy the number of Compounded Prescription Drug Orders Dispensed in the State where the Pharmacy is located and out of the State where the Pharmacy is located during a specified time period, including the Drugs' Active Ingredients, strength, and dosage form(s).
- (c) The Pharmacist shall notify patients if they may have received a Product found to have a defect or an out-of-specification result and conduct a recall, if the Board deems necessary.

## Section 3. Policy and Procedure Manual.

A policy and procedure manual shall be prepared and maintained for the Compounding, Dispensing, Delivery, Administration, storage, and use of Sterile and nonsterile Compounded Prescription Drugs. The policy and procedure manual shall incorporate all applicable USP requirements and:

- (a) include a quality assurance program for the purpose of monitoring patient care and Pharmacist Care Services outcomes; and
- (b) be current and available for inspection by a Board of Pharmacy-designated agent.

## Section 4. Physical Requirements.

- (a) Any Pharmacy that engages in Compounding shall adhere to physical, equipment, and environmental requirements established by USP.
- (b) Pharmacies shall have sufficient current reference materials applicable to Compounding.

---

<sup>165</sup> The Compounding of Drugs for animals must be done in accordance with the algorithm contained in the Animal Medicinal Drug Use Clarification Act of 1994 and associated FDA Guidance.



## **Section 5. Records and Reports.**

In addition to standard record-keeping and reporting requirements, the following records shall be maintained:

- (a) All Dispensing of sterile Compounded and nonsterile Compounded preparations.
- (b) Any other records required to conform to and demonstrate compliance with USP standards and Federal law.

## **Section 6. Delivery Service.**

The Pharmacist-in-Charge shall ensure the environmental control, stability, and sterility (if applicable) of all preparations shipped. Therefore, any Compounded preparation shall be shipped or Delivered to a patient or patient's agent in appropriate temperature-controlled (as defined by USP Standards) Delivery containers and stored appropriately. Information on appropriate storage shall be provided to the patient or patient's agent.

## **Section 7. Disposal of Hazardous and/or Infectious Wastes.**

The Pharmacist-in-Charge is responsible for ensuring that there is a system for the disposal of hazardous and/or infectious waste in accordance with applicable State and Federal laws and USP requirements.

## **Section 8. Quality Assurance.**

- (a) There shall be a documented, ongoing quality assurance program that monitors personnel performance, Component Verification and usage, Disinfection, sterilization, equipment, and facilities that are appropriate for the Drug being prepared. Quality assurance programs shall at minimum conform to the requirements of USP.
- (b) The Pharmacist has the responsibility and authority to inspect and approve or reject all Components, Drug Product containers, closures, in-process materials, and/or Labeling. The Pharmacist shall have the authority to prepare and review all Compounding records to ensure that no errors have occurred in the Compounding process. If errors have occurred, the Pharmacist is responsible for conducting a full investigation. A written record of the investigation shall be made and shall include conclusions and follow-up. The Pharmacist is also responsible for the proper maintenance, cleanliness, and use of all facilities and equipment used in Compounding.
- (c) All Pharmacists who participate in Compounding, including other Pharmacy personnel who assist the Pharmacist in Compounding, shall be proficient in the science of Compounding and shall acquire the education, training, and/or experience to maintain that proficiency through participation in seminars, studying appropriate literature, and consulting colleagues or by becoming certified by a Compounding certification program approved by the Board.
- (d) Pharmacists and other Compounding Pharmacy personnel (eg, Pharmacy Technicians) shall be trained and proficient in the particular operations that are performed by that individual.
- (e) Training shall be conducted by qualified individuals on a continuing basis and with sufficient frequency to ensure that Compounding Pharmacy personnel remain familiar with applicable operations and policies and procedures.
- (f) Only personnel authorized by the responsible Pharmacist shall be in the immediate vicinity of Compounding operations.
- (g) A Compounded Drug shall be deemed Adulterated if it has been prepared, packed, or held under insanitary conditions whereby it may have been contaminated with filth, or whereby it may have been rendered injurious to health.

## Section 9. Compounded Drug Preparations for Veterinary Use.

- (a) The use of bulk Drug substances for Compounded Drug preparations is prohibited except when:
  - (1) Compounding is pursuant to a patient-specific prescription for a non-food-producing animal or as an antidote to prevent animal suffering or death in food-producing animals;
  - (2) there is no marketed approved, conditionally approved, or indexed (in *The Index of Legally Marketed Unapproved New Animal Drugs for Minor Species*) Drug that can be used as labeled to treat the condition;
  - (3) there is no marketed approved animal or human Drug that can be used to treat the condition through off-label Drug use;
  - (4) the Drug cannot be appropriately Compounded from an approved animal or human Drug;
  - (5) immediate treatment with the Compounded Drug preparation is necessary to avoid animal suffering or death; and
  - (6) FDA has not identified a significant veterinary safety concern with the use of the bulk Drug substance for Compounding.
- (b) It is acceptable for any licensed Pharmacy to Compound veterinary Drug preparations to be used by veterinarians in their offices for Administration to clients' animals.
- (c) Compounded office use Drug preparations may be Dispensed by a veterinarian to clients only in an urgent or emergency situation for use in a single course of treatment, not to exceed a 120-hour supply.
- (d) Prohibition on wholesaling  
The Compounded veterinary Drug preparations will not be Distributed by an entity other than the Pharmacy that Compounded such veterinary Drug preparations. This does not prohibit Administration of a Compounded Drug preparation in a veterinary health care setting or Dispensing of a Compounded Drug preparation pursuant to a Prescription Drug Order executed in accordance with federal and state law.
- (e) Providing samples of Compounded veterinary Drug preparations is prohibited.
- (f) Upon becoming aware of any adverse event or Product defect, the Pharmacy reports the event on the designated FDA form<sup>166</sup> within 15 days and includes the FDA statement about reporting adverse events on the prescription Label.

---

<sup>166</sup> FDA Form 1932a or most current version.

## Model Rules for Outsourcing Facilities

### Section 1. Purpose and Scope.

The purpose of this section is to ensure that Outsourcing Facilities are regulated by this State in a manner consistent with Federal law and to ensure this State has appropriate authority over such facilities.

### Section 2. Registration.

- (a) Any Outsourcing Facility located in this State or that Distributes Compounded Pharmaceuticals to this State must be inspected and registered as an Outsourcing Facility by FDA prior to applying for a license/registration with the Board; and
- (b) The facility must undergo an inspection by the Board or a third party recognized by the Board such as Drug Distributor Accreditation<sup>167</sup> if the facility is registered with FDA but has not received an FDA inspection as an Outsourcing Facility.

### Section 3. Notification.

- (a) All licensed/registered Outsourcing Facilities shall report to the Board the biannual reports they are required to provide to FDA identifying the Drugs Compounded in the previous six (6)-month period, including the Drug's Active Ingredients, strength, and dosage form.

### Section 4. Requirements.

Outsourcing Facilities must:

- (a) Compound Drugs by or under the direct supervision of a licensed Pharmacist;
- (b) Compound Drugs in accordance with current Good Manufacturing Practice (cGMP) as required by Federal law;
- (c) Ensure that Pharmacists conducting or overseeing Compounding at an Outsourcing Facility must be proficient in the art of Compounding and shall acquire the education, training, and/or experience to maintain that proficiency through participation in seminars, studying appropriate literature, and consulting colleagues, and/or by becoming certified by a Compounding certification program approved by the Board.
- (d) Label Compounded Drugs with:
  - (i) required Drug and ingredient information,
  - (ii) facility identification, and
  - (iii) the following or similar statement: "This is a compounded drug. For office use only" or "Not for resale"; and
- (e) Only Compound using bulk Drug substances that meet specified FDA criteria. May also compound Drugs that appear on an FDA shortage list if the bulk drug substances used comply with the aforementioned specified criteria.

---

<sup>167</sup> States may require authentication and tracking of Product, whereby the exchange of information for Compounded Product is traced.

# Model Rules for the Licensure of Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors

## Section 1. Requirements for Licensure.

Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors that provide services within this State, whether located within this State or outside this State, shall be licensed by the Board and shall annually renew their license with the Board using an application provided by the Board. Third-Party Logistics Providers and Wholesale Drug Distributors must report license status to FDA as outlined in Federal law. Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors cannot operate from a place of residence. Where operations are conducted at more than one location, each such location shall be licensed by the Board of Pharmacy.<sup>168</sup>

- (a) Every Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor who engages in the Manufacturing, Repackaging, or Distribution of Prescription Drugs or Devices shall license annually with the Board by application and provide information required by the Board on an application approved by the Board, including but not limited to:
- (1) all trade or business names used by the licensee (includes “is doing business as” and “formerly known as”), which cannot be identical to the name used by another unrelated entity licensed to purchase Prescription Drugs or Devices in the State;
  - (2) name(s) of the owner and operator of the licensee (if not the same person), including:
    - (i) if a Person: the name, business address, Social Security number, and date of birth;
    - (ii) if a partnership: the name, business address, and Social Security number and date of birth of each partner, and the name of the partnership and federal employer identification number;
    - (iii) if a corporation: the name, business address, Social Security number and date of birth, and title of each corporate officer and director, the corporate names, the state of incorporation, federal employer identification number, and the name of the parent company, if any; the name, business address, and Social Security number of each shareholder owning ten percent (10%) or more of the voting stock of the corporation, including over-the-counter (OTC) stock, unless the stock is traded on a major stock exchange and not OTC;
    - (iv) if a sole proprietorship: the full name, business address, Social Security number, and date of birth of the sole proprietor and the name and federal employer identification number of the business entity;
    - (v) if a limited liability company: the name of each member, the name of each manager, their Social Security numbers or unique identifiers and their dates of birth, the name of the limited liability company and federal employer identification number, and the name of the state in which the limited liability company was organized; and
    - (vi) any other relevant information that the Board requires.
  - (3) name(s), business address(es), and telephone number(s) of a person(s) to serve as the Designated Representative(s) for each facility of a Wholesale Distributor that engages in the Wholesale Distribution of Prescription Drugs or Devices and additional information as required in Section 9 (Record Keeping);
  - (4) a list of all State and Federal licenses, registrations, or permits, including the license, registration, or permit numbers issued to the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor by any other State and Federal authority that authorizes the Manufacturer,

---

<sup>168</sup> The application and screening process for licensing entities engaging in the Distribution of Product represents a critical point in efforts to prevent the introduction of Counterfeit and Contraband Products into the medication distribution system. An application that requires detailed information about the applicant and key individuals involved in the operations of the entity is critical.

Repackager, Third-Party Logistics Provider, or Wholesale Distributor to Manufacture, purchase, possess, Repackage, or Distribute Prescription Drugs;

- (5) a list of all disciplinary actions by State and Federal agencies against the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor as well as any such actions against principals, owners, directors, or officers;
- (6) a full description of each facility and warehouse, including all locations utilized for Prescription Drug storage and/or Wholesale Distribution. The description should include the following:
  - (i) square footage;
  - (ii) security and alarm system descriptions;
  - (iii) terms of lease or ownership;
  - (iv) address; and
  - (v) temperature and humidity controls.
- (7) a copy of the deed for the property on which the Manufacturer's, Repackager's, Third-Party Logistics Provider's, or Wholesale Distributor's establishment is located, if the property is owned by the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor; or a copy of the Manufacturer's, Repackager's, Third-Party Logistics Provider's, or Wholesale Distributor's lease for the property on which the establishment is located that has an original term of not less than one (1) calendar year, if the establishment is not owned by the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor;
- (8) information regarding general and Product liability insurance, including copies of relevant policies;
- (9) a description of the Manufacturer's, Repackager's, Third-Party Logistics Provider's, or Wholesale Distributor's Drug import and export activities; and
- (10) a copy of the Manufacturer's, Repackager's, Third-Party Logistics Provider's, or Wholesale Distributor's written policies and procedures as required in Section 11 (Policies and Procedures).
- (11) The information collected pursuant to Section 1(a)(6) and (a)(10) shall be made available only to the Board, a third party recognized by the Board, and to State and Federal law enforcement officials. The Board shall make provisions for protecting the confidentiality of the information collected under this section.

- (b) A "surety" bond of not less than \$100,000, or other equivalent means of security acceptable to the Board or a third party recognized by the Board such as insurance, an irrevocable letter of credit, or funds deposited in a trust account or financial institution, to secure payment of any administrative penalties imposed by the Board and any fees or costs incurred by the Board regarding that licensee when those penalties, fees, or costs are authorized under state law and the licensee fails to pay thirty (30) days after the penalty, fee, or costs becomes final. A separate "surety" bond or other equivalent means of security is not required for each company's separate locations or for affiliated companies/groups when such separate locations or affiliated companies/groups are required to apply for or renew their Third-Party Logistics Provider's or Wholesale Distributor's license with the Board. The Board may make a claim against such bond or other equivalent means of security until one year after the Third-Party Logistics Provider's or Wholesale Distributor's license ceases to be valid or until sixty (60) days after any administrative or legal proceeding before or on behalf of the Board that involves the Third-Party Logistics Provider or Wholesale Distributor is concluded, including any appeal, whichever occurs later. Manufacturers and Repackagers shall be exempt from securing a "surety" bond or other equivalent means of security acceptable to the Board or a third party recognized by the Board. The Board may waive the bond requirement, if the Third-Party Logistics Provider or Wholesale Distributor:<sup>169</sup>

---

<sup>169</sup> Although Wholesale Distributors may be licensed in multiple states, it is not intended for Wholesale Distributors to procure a separate "surety" bond (or other equivalent means) for each state of licensure. States should consider waiving this requirement if the Wholesale Distributor has procured a "surety" bond (or other equivalent means) for the purposes of licensure in another state, or if the wholesaler is a publicly traded company.

- (1) has previously obtained a comparable surety bond or other equivalent means of security for the purpose of licensure in another state, where the Third-Party Logistics Provider or Wholesale Distributor possesses a valid license in good standing; or
- (2) is a publicly held company.
- (c) Every Manufacturer, Repackager, Third Party Logistics Provider, or Wholesale Distributor who engages in Manufacturing, Repackaging, or Wholesale Distribution shall submit a reasonable fee to be determined by the Board.
- (d) Each facility that engages in Distribution must undergo an inspection by the Board or a third party recognized by the Board for the purpose of inspecting the Distribution operations prior to initial licensure and periodically thereafter in accordance with a schedule to be determined by the Board.
- (e) All Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors must publicly display or have readily available all licenses and the most recent inspection report administered by the Board if applicable.
- (f) Changes in any information in this Section shall be submitted to the Board, or to a third party recognized by the Board, within 30 days of such change (unless otherwise noted).
- (g) Information submitted by the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor to the Board or a third party recognized by the Board that is considered trade secret or proprietary information, as defined under this State's privacy and trade secret/proprietary statutes, shall be maintained by the Board or a third party recognized by the Board as private or trade secret/proprietary information and be exempt from public disclosure.<sup>170</sup>
- (h) Per Federal requirements, States shall license Third-Party Logistics Providers (those that provide storage and logistical operations related to Drug Distribution) separately from Wholesale Drug Distributors. Minimum requirements for Wholesale Drug Distributor licensure may also apply to Third-Party Logistics Providers if applicable.<sup>171</sup>
- (i) Per Federal requirements, States shall license Repackagers and Manufacturers separately from Wholesale Drug Distributors. Minimum requirements for Wholesale Drug Distributor licensure may also apply to Repackagers and Manufacturers if applicable.
- (j) Supply chain Trading Partners (Wholesale Drug Distributors and Third-Party Logistics Providers) should report State licensure status and other required information to FDA.

## **Section 2. Minimum Qualifications.**

- (a) The Board will consider the following factors in determining the eligibility for, and renewal of, licensure of Persons who engage in the Wholesale Distribution of Drugs or Devices:
  - (1) any findings by the Board that the applicant has violated or been disciplined by a regulatory agency in any state for violating any Federal, State, or local laws relating to Drug or Device Wholesale Distribution;
  - (2) any criminal convictions of the applicant under Federal, State, or local laws;
  - (3) the applicant's past experience in the Manufacture or Wholesale Distribution of Drugs or Devices;
  - (4) the furnishing by the applicant of false or fraudulent material in any application made in connection with Drug or Device Manufacturing or Wholesale Distribution;
  - (5) Suspension, sanction, or Revocation by federal, State, or local government against any license currently or previously held by the applicant or any of its owners for violations of State or Federal laws regarding Drugs or Devices;
  - (6) compliance with previously granted licenses of any kind;

<sup>170</sup> The Board may designate a third party to conduct inspections and ensure that all requirements for licensure established by the legislature and Board are fulfilled. The NABP Drug Distributor Accreditation program is available to the states.

<sup>171</sup> If a State does not have a licensure category for Third-Party Logistics Providers, facilities that engage in interstate transport of Prescription Drugs must obtain Federal registration.

- (7) compliance with the requirements to maintain and/or make available to the Board licensing authority or to Federal, State, or local law enforcement officials those records required to be maintained by Wholesale Distributors; and
  - (8) any other factors or qualifications the Board considers relevant to and consistent with the public health and safety.
- (b) The Board shall consider the results of a criminal and financial background check of the applicant, including but not limited to, all key personnel involved in the operations of the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor, including the most senior Person responsible for facility operations, purchasing, and inventory control and the Person or Persons they report to; and all company officers, key management, principals, and owners with ten percent (10%) or greater ownership interest in the company (applying to non-publicly held companies only) to determine if an applicant or others associated with the ownership, management, or operations of the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor have committed criminal acts that would constitute grounds for denial of licensure. The background check will be conducted in compliance with any applicable State and Federal laws, at the applicant's expense, and will be sufficient to include all States of residence since the Person has been an adult. Manufacturers shall be exempt from criminal and financial background checks.
- (c) The applicant shall provide, and attest to, a statement providing a complete disclosure of any past criminal convictions and violations of the State and Federal laws regarding Drugs or Devices or an affirmation and attestation that the applicant has not been involved in, or convicted of, any criminal or prohibited acts.

### **Section 3. Personnel.**

Each Person that is issued an initial or renewal license as a Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor, whether in State or out of State, must designate in writing on a form required by the Board, a Person for each facility to serve as the Designated Representative.

- (a) To be certified as a Designated Representative, a Person must:
- (1) submit an application on a form furnished by the Board and provide information that includes, but is not limited to:
    - (i) information required to complete the criminal and financial background checks required under Section 2(b);<sup>172</sup>
    - (ii) date and place of birth;
    - (iii) occupations, positions of employment, and offices held during the past seven (7) years;
    - (iv) principal business and address of any business corporation, or other organization in which each such office of the Person was held or in which each such occupation or position of employment was carried on;
    - (v) whether the Person, during the past seven (7) years, has been enjoined, either temporarily or permanently, by a court of competent jurisdiction from violating any Federal or State law regulating the possession, control, or Wholesale Distribution of Prescription Drugs or Devices, together with details of such events;
    - (vi) description of any involvement by the Person with any business, including any investments, other than the ownership of stock in a publicly traded company or mutual fund, during the past seven (7) years, which Manufactured, Administered, Prescribed, Repackaged, Wholesale Distributed, or stored Prescription Drugs and Devices in which such businesses were named as a party in a lawsuit;

---

<sup>172</sup> Fingerprints represent one of the current means of verifying the identity of the person as well as providing a reliable means to conduct criminal background checks. As technology changes and other means become available to the Board such as retinal scanning or DNA sampling, the Board must stay current with such technologies and amend rules as necessary and appropriate.

- (vii) description of any criminal offense (not including minor traffic violations) of which the Person, as an adult, was found guilty, regardless of whether adjudication of guilt was withheld or whether the Person pled guilty or nolo contendere. If the Person indicates that a criminal conviction is under appeal and submits a copy of the notice of appeal of the criminal offense, the applicant must, within 15 days after the disposition of the appeal, submit to the Board a copy of the final written order of disposition;
  - (viii) photograph of the Person taken within the previous 30 days under procedures as specified by the Board;
  - (ix) name, address, occupation, and date and place of birth for each member of the Person's immediate family, unless the Person is employed by a Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor that is a publicly held company. As used in this subparagraph, the term "member of the immediate family" includes the Person's spouse(s), children, parents, siblings, the spouses of the Person's children, and the spouses of the Person's siblings; and
  - (x) any other information the Board deems relevant.
- (2) have a minimum of two years of verifiable full-time managerial or supervisory experience in a Pharmacy or Wholesale Distribution facility licensed in this State or another state, where the Person's responsibilities included but were not limited to record keeping, storage, and shipment of Prescription Drugs or Devices;
- (3) may serve as the Designated Representative for only one Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor at any one time, except where more than one licensed Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor is co-located in the same facility and such entities are members of an affiliated group, as defined in Section 1504 of the Internal Revenue Code;
- (4) be actively involved in and aware of the actual daily operations of the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor:
  - (i) employed full-time in a managerial position by the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor;
  - (ii) physically present at the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor during normal business hours, except for time periods when absent due to illness, family illness or death, scheduled vacation, or other authorized absence; and
  - (iii) aware of, and knowledgeable about, all policies and procedures pertaining to the operations of the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor.
- (b) The information collected pursuant to Section 3(a) shall be made available only to the Board, a third party recognized by the Board, and to State and Federal law enforcement officials. The Board and a third party recognized by the Board shall make provisions for protecting the confidentiality of the information collected under this section.
- (c) Each licensed Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor located outside of this State that Distributes Prescription Drugs or Devices in this State shall designate a registered agent in this State for service of process. Any licensed Distributor that does not so designate a registered agent shall be deemed to have designated the Secretary of State of this State to be its true and lawful attorney, upon who may be served all legal processes in any action or proceeding against such licensed Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor growing out of or arising from such Manufacturing, Repackaging, or Distribution. A copy of any such service of process shall be mailed to such Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor by the Board by certified mail, return receipt requested, postage prepaid, at the address such licensed entity has designated on its application for licensure in this State. If any such Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor is not licensed in this State, service on the Secretary of State only shall be sufficient service.



- (d) A Designated Representative must complete:<sup>173</sup>
- (1) continuing education programs specified by the Board regarding Federal and State laws in regard to the Wholesale Distribution, handling, and storage of Prescription Drugs or Devices; or
  - (2) if no formal continuing education is specified by the Board, training programs that address applicable Federal and State laws and are provided by qualified in-house specialists, outside counsel, or consulting specialists with capabilities to help ensure compliance.

#### **Section 4. Minimum Requirements for the Storage and Handling of Prescription Drugs and for Establishment and Maintenance of Prescription Drug Records.**

The following are required for the storage, handling, transport, and shipment of Prescription Drugs or Devices, and for the establishment and maintenance of Wholesale Distribution records by Wholesale Distributors, authorized Trading Partners, and their officers, agents, representatives, and employees.

- (a) All facilities at which Prescription Drugs and Devices are received, stored, warehoused, handled, held, offered, marketed, displayed, or transported from shall:
- (1) be of suitable construction to ensure that all Prescription Drugs and Devices in the facilities are maintained in accordance with the Product Labeling of such Prescription Drugs and Devices, or in compliance with official compendium standards such as the United State Pharmacopeia–USP-NF;
  - (2) be of suitable size and construction to facilitate cleaning, maintenance, and proper Wholesale Distribution operations;
  - (3) have adequate storage areas to provide adequate lighting, ventilation, temperature, sanitation, humidity, space, equipment, and security conditions;
  - (4) have a quarantine area for storage of Prescription Drugs and Devices that are outdated, damaged, deteriorated, Misbranded, or Adulterated, Counterfeit, or suspected of being Counterfeit, otherwise unfit for Distribution, or that are in immediate or sealed secondary containers that have been opened;
  - (5) be maintained in a clean and orderly condition;
  - (6) be free from infestation of any kind;
  - (7) be a commercial location and not a personal dwelling or residence;
  - (8) provide for the secure and confidential storage of information with restricted access and policies and procedures to protect the integrity and confidentiality of the information;
  - (9) provide and maintain appropriate inventory controls in order to detect and document any theft, Counterfeiting, or diversion of Prescription Drugs or Devices.
- (b) Wholesale Distributors, Third-Party Logistics Providers, or other Trading Partners involved in the Wholesale Distribution of controlled substances shall be duly registered with Drug Enforcement Administration (DEA) and appropriate state controlled substance agency and in compliance with all applicable laws and rules for the storage, handling, transport, shipment, and Wholesale Distribution of controlled substances.

#### **Section 5. Security.**

- (a) All facilities used for Wholesale Distribution shall be secure from unauthorized entry:
- (1) access from outside the premises shall be kept to a minimum and be well-controlled;
  - (2) the outside perimeter of the premises shall be well-lighted; and
  - (3) entry into areas where Prescription Drugs or Devices are held shall be limited to authorized personnel; all facilities shall be equipped with an alarm system to detect entry after hours.

---

<sup>173</sup> The Board will need to ensure that continuing education programs for the desired content areas are available when considering the implementation of this requirement.

- (b) All facilities shall be equipped with a security system that will provide suitable protection against theft and diversion. When appropriate, the security system shall provide protection against theft or diversion that is facilitated or hidden by tampering with computers or electronic records.
- (c) All facilities shall be equipped with inventory management and control systems that protect against, detect, and document any instances of theft, diversion, or Counterfeiting.
- (d) All common carriers used by a Wholesale Distributor or Third-Party Logistics Provider shall ensure security via one of the following:
  - (1) a verifiable security system; or
  - (2) a Board-approved accreditation or certification program.
- (e) All facilities shall be equipped with security systems to protect the integrity and confidentiality of data and documents and make such data and documents readily available to the Board and other state and federal law enforcement officials.

## **Section 6. Storage.**

All Prescription Drugs and Devices shall be stored at appropriate temperatures and under appropriate conditions in accordance with requirements, if any, in the Product Labeling of such Prescription Drugs and Devices, or with requirements in the current edition of an official compendium such as the USP-NF.

- (a) If no storage requirements are established for a Prescription Drug, the Prescription Drug may be held at “controlled” room temperature, as defined in an official compendium such as USP-NF, to help ensure that its identity, strength, quality, and purity are not adversely affected.
- (b) Appropriate manual, electromechanical, or electronic temperature and humidity recording equipment and/or logs shall be utilized to document proper storage of Prescription Drugs and Devices.
- (c) Packaging of the Prescription Drugs and Devices should be in accordance with an official compendium such as USP-NF and identify any compromise in the integrity of the Prescription Drugs or Devices due to tampering or adverse storage conditions.
- (d) Controlled substance Drugs should be isolated from non-controlled substance Drugs and stored in a secure area in accordance with Drug Enforcement Administration security requirements and standards.
- (e) The record-keeping requirements in Section 9 (Record Keeping) shall be followed for the Wholesale Distribution of all Prescription Drugs and Devices.

## **Section 7. Operations/Reporting Requirements.**

- (a) Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Drug Distributors must comply with all reporting requirements and exchange Transaction History, Transaction Information, and Transaction Statements with authorized Trading Partners as outlined in Federal law.
- (b) Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors shall design and operate a system to identify and report Suspicious Orders of controlled substances and Drugs of Concern to a program approved by the Board.
  - (1) Suspicious Orders shall be submitted electronically to an approved program within five days of the order being identified as suspicious by the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor, and must include, but not be limited to:
    - (i) customer name;
    - (ii) NABP e-Profile ID;
    - (iii) customer address;
    - (iv) customer DEA registration number;
    - (v) state license number(s);
    - (vi) Transaction date;
    - (vii) Drug name;
    - (viii) NDC number;

- (ix) quantity ordered; and
  - (x) indication of whether the Drug was shipped, and if not, the factual basis for the refusal to supply.
- (2) Zero reports shall be submitted if no Suspicious Orders have been identified in a calendar month, and such reports shall be submitted within 15 days of the end of the calendar month.
- (3) Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors may apply to the Board for an exemption from the reporting requirements if they do not distribute controlled substances or Drugs of Concern.
- (c) Except as described in paragraph 7(d), a Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor shall exercise due diligence to identify customers ordering or seeking to order controlled substances or Drugs of Concern and establish the normal and expected transactions conducted by those customers, as well as to identify and prevent the sale of controlled substances or Drugs of Concern that are likely to be diverted from legitimate channels. Such due diligence measures shall include, but are not limited to, the following, which shall be conducted prior to an initial sale and on a regular basis, as necessary:
  - (1) questionnaires and affirmative steps by the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor to confirm the accuracy and validity of the information provided;
  - (2) for a customer who is a prescriber, confirmation of prescriber type, specialty practice area, and if the prescriber personally furnishes controlled substances or Drugs of Concern, the quantity furnished;
  - (3) review of drug utilization reports; and
  - (4) obtaining and conducting a review of the following:
    - (i) methods of payment accepted and in what ratios;
    - (ii) the ratio of controlled versus non-controlled Drug orders and overall sales;
    - (iii) orders for controlled substances or Drugs of Concern from other Manufacturers, Repackagers, Third-Party Logistics Providers, or Wholesale Distributors made available by US DEA's Automation of Reports and Consolidated Orders System (ARCOS); and
    - (iv) the ratio of out-of-state patients served compared to in-state patients.
- (d) A Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor receiving a request for an initial sale of a controlled substance or Drug of Concern may conduct the sale before complying with paragraph 7(c) if all the following apply:
  - (1) the sale is to a new customer;
  - (2) the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor documents that the order is to meet an emergent need;
  - (3) the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor completes the requirements of 7(c) no later than 60 days from the date of sale.
- (e) A Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor receiving a request from an existing customer to purchase a controlled substance or Drug of Concern, the size/quantity of which exceeds the established algorithm limitations or quota restrictions for such customer, may sell the Drug of Concern or controlled substance provided that the customer submits documentation of an emergent need for a specific patient.
- (f) Any customer that is believed to be engaged in potential Diversion Activities, including those to whom a Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor refuses to sell, shall be electronically reported to a program approved by the Board. Such reports shall include:
  - (1) customer name;
  - (2) NABP e-Profile ID;
  - (3) customer address;
  - (4) DEA number;
  - (5) state license number(s); and

(6) a detailed explanation of why the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor identified the customer as a possible diversion risk.

Such reports shall be submitted within 30 days of refusal, cessation, or identification by the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor.

- (g) Within 90 days of the effective date of this rule, a Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor shall provide to a program approved by the Board, information on all customers in the state where the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor has refused to sell or has stopped selling within the past year because the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor has identified the customer(s) as engaging in potential Diversion Activity that may cause reported Drugs to be diverted from legitimate channels.
- (h) All licensed Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors shall submit all reports to a Board-approved program in a DEA ARCOS format.

## **Section 8. Due Diligence.**

- (a) Supply chain Trading Partners (Manufacturers, Repackagers, Wholesale Distributors, and Dispensers) shall exercise due diligence when conducting business to expeditiously identify a Suspect Product and determine whether it is suspect (and after investigation, whether it is illegitimate).
- (b) Supply chain Trading Partners (Manufacturers, Repackagers, Wholesale Distributors, and Dispensers) shall establish a system to:
- (1) Quarantine and investigate Suspect Product to determine if it is illegitimate.
  - (2) Notify FDA, the Board, and immediate Trading Partners if Illegitimate Product is found.
- (c) Manufacturers, Repackagers, Wholesale Distributors, and Dispensers shall establish processes for identifying their Trading Partners and Transactions that require heightened vigilance in preventing the receipt of Suspect Products.
- (d) Heightened vigilance includes the examination of required records (invoices, shipping documents, Transaction History, and Transaction Statement) for suspicious business practices and the physical examination of Products for factors that increase the risk of a Product being suspect, such as:
- (1) a Trading Partner that has been involved in business Transactions where they sold or delivered Illegitimate Product;
  - (2) a Trading Partner that has a history of problematic or potentially false Transaction Histories or pedigrees, such as those that contain misspelled words or incomplete information;
  - (3) a Trading Partner that is reluctant to provide a Transaction History associated with the Product being purchased or does not do so in a timely manner;
  - (4) a Trading Partner that provides Transaction Information, a Transaction Statement, and/or Transaction History that appears to be incomplete or suspicious;
  - (5) the price of a Product is suspicious;
  - (6) the Product has been previously or is currently the subject of a Drug shortage;
  - (7) a Product that is in higher demand because of its potential or perceived relationship to a public health or other emergency;
  - (8) the appearance of the package is suspicious; or
  - (9) the package exhibits unusual or excessive adhesive residue.

## **Section 9. Record Keeping.**

- (a) Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors shall establish and maintain inventories and records of all Transactions regarding the receipt and Distribution or other disposition of Prescription Drugs and Devices as outlined in Federal law. These records shall include:
- (1) dates of receipt and Wholesale Distribution; or

- (2) other disposition of the Prescription Drugs and Devices.
- (b) Such records shall include the Inventories and records and shall be made available for inspection and photocopying by any authorized official of any State, Federal, or local governmental agency for a period of six (6) years following their creation date.
- (c) All records related to the Wholesale Distribution of Prescription Drugs, including, but not limited to, invoices of purchase, packing slips, shipping records, and sales invoices will accurately reflect the name of the Wholesale Distributor as it appears on the facility's license issued by the state in which the Wholesale Distributor is engaged in Wholesale Distribution. Wholesale Distributors to which a license has been issued in the same name and at the same address as another licensee authorized to purchase Prescription Drugs must utilize a method to distinguish purchases and Distributions that are specific to the Wholesale Distributor.
- (d) Records described in this section that are kept at the inspection site or that can be immediately retrieved by computer or other electronic means shall be readily available for authorized inspection during the retention period. Records kept at a central location apart from the inspection site and not electronically retrievable shall be made available for inspection within two (2) working days of a request by an authorized official of any State or Federal governmental agency charged with enforcement of these rules.
- (e) Wholesale Distributors, Third-Party Logistics Providers, Repackagers, and Manufacturers should maintain an ongoing list of Persons with whom they do business.
- (f) All facilities shall establish and maintain procedures for reporting Counterfeit and Contraband or suspected Counterfeit and Contraband Drugs or Devices or Counterfeiting and Contraband or suspected Counterfeiting and Contraband activities to the Board and FDA.
- (g) Wholesale Distributors shall maintain a system for the mandatory reporting of any theft, suspected theft, diversion, or other significant loss of any Prescription Drug or Device to the Board and FDA, and, where applicable, to DEA.<sup>174</sup>

## Section 10. Policies and Procedures.

Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors shall establish, maintain, and adhere to written policies and procedures, which shall be followed for the receipt, security, storage, inventory, transport, and shipping and Wholesale Distribution of Prescription Drugs, including policies and procedures for identifying, recording, and reporting losses or thefts and for correcting all errors and inaccuracies in inventories. Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors shall include in their written policies and procedures the following:<sup>175</sup>

- (a) A procedure to be followed for handling recalls and withdrawals of Prescription Drugs and Devices. Such procedure shall be adequate to deal with recalls and withdrawals due to:
  - (1) Any action initiated at the request of FDA or any other federal, state, or local law enforcement or other government agency, including the Board of Pharmacy; or
  - (2) Any volunteer action by the Manufacturer to remove defective or potentially defective Prescription Drugs or Devices from the market.
- (b) A procedure to ensure that Manufacturers, Repackagers, Third-Party Logistics Providers, and Wholesale Distributors prepare for, protect against, and handle any crisis that affects security or operation of any facility in the event of a strike, fire, flood, or other natural disaster, or other situations of local, State, or national emergency.
- (c) A procedure to ensure that any outdated Prescription Drugs shall be segregated from other Prescription Drugs and either returned to the Manufacturer or destroyed in accordance with Federal and State laws,

<sup>174</sup> This information should be reported to NABP, if serving as a data collection repository, in addition to the other relevant authorities.

<sup>175</sup> In developing policies and procedures for the management and quality improvement of the Wholesale Distribution activities of a Wholesale Distributor, the Board may want to refer to the Healthcare Distribution Management Association and the National Association of Chain Drug Stores.

including all necessary documentation and the appropriate witnessing. This procedure shall provide for written documentation of the disposition of outdated Prescription Drugs.

- (d) A procedure for the destruction of outdated Prescription Drugs in accordance with federal and state laws.
- (e) A procedure for the disposing and destruction of containers, Labels, and packaging to ensure that the containers, Labels, and packaging cannot be used in Counterfeiting activities, including all necessary documentation, maintained for a minimum of three (3) years, and the appropriate witnessing of the destruction of any Labels, packaging, Immediate Containers, or containers in accordance with all applicable Federal and State requirements.
- (f) A procedure for identifying, investigating and reporting significant Prescription Drug inventory discrepancies involving Counterfeit, suspect of being Counterfeit, Contraband, or suspect of being Contraband, in the inventory and reporting of such discrepancies as required to FDA, Board and/or appropriate Federal or State agency upon discovery of such discrepancies.
- (g) A procedure for reporting criminal or suspected criminal activities involving the inventory of Prescription Drug(s) and Device(s) as required to the Board, FDA, and, if applicable, DEA.
- (h) A procedure for verifying security provisions of Common Carriers.
- (i) Procedures addressing:
  - (1) the design and operation of the Suspicious Order monitoring and reporting system;
  - (2) mandatory annual training for staff responsible for identifying and reporting Suspicious Orders and potential Diversion Activities. Such training must include the following:
    - (i) the Manufacturer, Repackager, Third-Party Logistics Provider, or Wholesale Distributor's Suspicious Order monitoring system;
    - (ii) the process to collect all relevant information on customers in accordance with paragraph 7(c);
    - (iii) the requirement and process for submission of Suspicious Orders and information on customers who engage in potential Diversion Activities.
- (j) A procedure for timely responding to customers who submit purchase orders for patients with emergent needs.

## **Section 11. Prohibited Acts.**<sup>176</sup>

It is unlawful for a Person to knowingly and willfully perform or cause the performance of or aid and abet any of the following acts in this State:

- (a) the Manufacture, Repackaging, sale, delivery, or holding or offering for sale any Prescription Drug or Device that is Adulterated, Misbranded, Counterfeit, suspected of being Counterfeit, or has otherwise been rendered unfit for Distribution or Wholesale Distribution;
- (b) the Adulteration, Misbranding, or Counterfeiting of any Prescription Drug or Device;
- (c) the receipt of any Prescription Drug or Device that is Adulterated, Misbranded, stolen, obtained by fraud or deceit, Counterfeit, or suspected of being Counterfeit, or the delivery or proffered delivery of such Prescription Drug or Device for pay or otherwise;
- (d) the Alteration, mutilation, destruction, obliteration, or removal of the whole or any part of the Product Labeling of a Prescription Drug or Device or the commission of any other act with respect to a Prescription Drug or Device that results in the Prescription Drug or Device being Misbranded;
- (e) the forging, Counterfeiting, simulating, or falsely representing of any Prescription Drug or Device without the authority of the Manufacturer, or using any mark, stamp, tag, label, or other identification device without the authorization of the Manufacturer;

---

<sup>176</sup> Boards should be advised that statutory amendments may be necessary in the State practice acts in order to properly integrate the prohibited and criminal acts. It is highly recommended that boards confer with Board Counsel before adopting the Model Rules to make certain that the appropriate statutory language exists in the state pharmacy practice act to support the Model Rules.

- (f) the purchase or receipt of a Prescription Drug or Device from a Person that is not licensed to Distribute Prescription Drugs or Devices to that purchaser or recipient;
- (g) the sale or transfer of a Prescription Drug or Device to a Person who is not legally authorized to receive a Prescription Drug or Device;
- (h) the sale or transfer of a Prescription Drug or Device from Pharmacies to Distributors for resale;<sup>177</sup>
- (i) the failure to maintain or provide records as required by this Act and Rules;
- (j) providing the Board or any of its representatives or any state or federal official with false or fraudulent records or making false or fraudulent statements regarding any matter within the provisions of this Act and Rules;
- (k) the Wholesale Distribution of any Prescription Drug or Device that was:
  - (1) purchased by a public or private hospital or other health care entity;
  - (2) donated or supplied at a reduced price to a charitable organization; or
  - (3) stolen or obtained by fraud or deceit.
- (l) the failure to obtain a license or operating without a valid license when a license is required;
- (m) the Obtaining of or attempting to obtain a Prescription Drug or Device by fraud, deceit, misrepresentation or engaging in misrepresentation or fraud in the Distribution or Wholesale Distribution of a Prescription Drug or Device;
- (n) the Distributing of a Prescription Drug or Device to the patient without a Prescription or Prescription Order from a Practitioner licensed by law to use or prescribe the Prescription Drug or Device;
- (o) the Distributing or Wholesale Distributing of a Prescription Drug or Device that was previously dispensed by a Pharmacy or distributed by a Practitioner; or
- (p) the failure to report any Prohibited Act as listed in these Rules.

## **Section 12. Criminal Acts.**<sup>178</sup>

- (a) A Person who, with intent to defraud or deceive, performs the act of Adulteration, Misbranding, or Counterfeiting of any Prescription Drug or Device commits a felony of the third degree.
- (b) A Person who engages in the Wholesale Distribution of Prescription Drug(s) who, with intent to defraud or deceive, falsely swears or certifies that he or she has authenticated any documents related to the Wholesale Distribution of Prescription Drugs, commits a felony of the third degree.
- (c) A Person who engages in the Wholesale Distribution of Prescription Drug(s) or Device(s) and knowingly purchases or receives Prescription Drug(s) or Device(s) from a Person, not legally authorized to Wholesale Distribute Prescription Drug(s) or Device(s), in Wholesale Distribution commits a felony of the third degree.
- (d) A Person who engages in the Wholesale Distribution of Prescription Drug(s) or Device(s) and knowingly sells, barter, brokers, or transfers Prescription Drug(s) or Device(s) to a Person not legally authorized to purchase Prescription Drug(s) or Device(s), under the jurisdiction in which the Person receives the Prescription Drug(s) or Device(s) in a Wholesale Distribution, commits a felony of the third degree.
- (e) A Person who knowingly possesses, actually or constructively, any amount of a Contraband Drug(s) or Device(s), who knowingly sells or delivers, or who possesses with intent to sell or deliver any amount of a Contraband Drug(s) or Device(s) commits a felony of the third degree.
- (f) A Person who knowingly forges, Counterfeits, or falsely creates any Label for a Prescription Drug(s) or Device(s) or who falsely represents any factual matter contained in any Label of a Prescription Drug(s) or Device(s) commits a felony of the third degree.

---

<sup>177</sup> Returned purchases from Pharmacies to Wholesale Distributors are not considered to be “transfers, Distributions, or sales,” and are not affected by this language.

<sup>178</sup> Boards should be advised that statutory amendments may be necessary in the State practice acts in order to properly integrate the prohibited and criminal acts. It is highly recommended that boards confer with Board Counsel before adopting the Model Rules to make certain that the appropriate statutory language exists in the state pharmacy practice act to support the Model Rules.

- (g) A Person who knowingly Manufactures, purchases, sells, delivers, or brings into the State, or who is knowingly in actual or constructive possession of any amount of Contraband Drug(s) or Device(s), commits a felony of the third degree.
- (h) A Person who knowingly Manufactures, purchases, sells, delivers, or brings into the State, or who is knowingly in actual or constructive possession of any amount of Contraband Drug(s) or Device(s), and whose acts result in the death of a Person, commits a felony in the first degree.
- (i) A Person found guilty of any offense under this section, under the authority of the Court convicting and sentencing the Person, shall be ordered to forfeit to the State any real or Personal property:
  - (1) used or intended to be used to commit, to facilitate, or to promote the commission of such offense; and
  - (2) constituting, derived from, or traceable to the gross proceeds that the defendant obtained directly or indirectly as a result of the offense. Any property or assets subject to forfeiture under this section may be seized pursuant to a warrant obtained in the same manner as a search warrant or as otherwise permitted by law, and held until the case against a defendant is adjudicated. Monies ordered forfeited, or proceeds from the sale of other assets ordered forfeited, shall be equitably divided between the Board and other agencies involved in the investigation and prosecution that led to the conviction. Other property ordered forfeited after conviction of a defendant may, at the discretion of the investigating agencies, be placed into official use by the Board or the agencies involved in the investigation and prosecution that led to the conviction.

### **Section 13. Salvaging and Reprocessing.**

Wholesale Distributors, Third-Party Logistics Providers, and Trading Partners shall be subject to the provisions of any applicable Federal, State, or local laws or rules that relate to Prescription Drug Product salvaging or reprocessing, including Chapter 21, parts 207, 210, and 211k of the Code of Federal Regulations.

### **Section 14. Inspection and Accreditation by a Third Party.**

- (a) The Board shall have the authority to recognize a third party to inspect and accredit Wholesale Distributors.
- (b) The Board may license by reciprocity a Wholesale Distributor and Third-Party Logistics Provider that is licensed under the laws of another state, if:
  - (1) the requirements of that State are deemed by the Board to be substantially equivalent; or
  - (2) the applicant is accredited by a third party recognized by the Board. An applicant that is accredited by a third party recognized and approved by the Board, shall not be subject to duplicative requirements set by the Board. If an applicant is inspected, but not accredited by a third party, that applicant must comply with the requirements set by the Board through regulation.
- (c) Any applicant that is denied accreditation described under paragraph (a), shall have the right of review of the accreditation body's decision, by:
  - (1) the accreditation body; and
  - (2) the Board.
- (d) The Board recognized accreditation body shall ensure that the proprietary information obtained during the accreditation process remains confidential and privileged.
- (e) The Board may waive requirements of this Chapter, by regulation, for Wholesale Distributors that have obtained and maintain a Board-approved accreditation.



# Model Rules for the Licensure of Medical Gas and Medical Gas Related Equipment Wholesale Distributors

## Section 1. Definitions.

- (a) “Adulterated Medical Gas or Medical Gas Related Equipment.” A Medical Gas or Medical Gas Related Equipment shall be deemed to be Adulterated:
- (1) if:
- (i) it consists in whole or in part of any impurities or deleterious substances exceeding normal specifications;
  - (ii) it has been produced, prepared, packed, or held under conditions whereby the medical gas may have been contaminated causing it to be rendered injurious to health; or if the methods used in, or the facilities or controls used for, its manufacture, processing, packing, or holding do not conform to or are not operated or administered in conformity with current good manufacturing practices to ensure that the medical gas meets the requirements of this part as to safety and has the identity and strength, and meets the quality and purity characteristics that it purports or is represented to possess; or
  - (iii) its container interior is contaminated with any poisonous or deleterious substance that may render the contents injurious to health; or
- (2) if it purports to be or is represented as a Medical Gas, the name of which is recognized in the United States Pharmacopeia–National Formulary (USP-NF), and its strength differs from, or its quality or purity falls below, the standard set forth in the USP-NF. Such a determination as to strength, quality, or purity shall be made in accordance with the tests or methods of assay set forth in the USP-NF, or validated equivalent, or in the absence of or inadequacy of these tests or methods of assay, those prescribed under authority of the Federal Act. No medical gas defined in USP-NF shall be deemed to be adulterated under this paragraph because it differs from the standard of strength, quality, or purity therefore set forth in the compendium, if its difference in strength, quality, or purity from that standard is plainly stated on its label; or
- (3) if it is not subject to paragraph (2) and its strength differs from, or its purity or quality falls below, that which it purports or is represented to possess.
- (b) “Authorized Distributor of Record of Medical Gases or Medical Gas Related Equipment” means a distributor with whom a manufacturer has established an ongoing relationship to distribute the Manufacturer’s Products. An ongoing relationship is deemed to exist between such Wholesale Distributor and a Manufacturer when the Wholesale Distributor, including any affiliated group of the Wholesale Distributor, as defined in Section 1504 of the Internal Revenue Code, complies with the following:
- (1) the Wholesale Distributor has a written agreement currently in effect with the Manufacturer evidencing such ongoing relationship; and
  - (2) the Wholesale Distributor is listed on the Manufacturer’s current list of authorized distributors of record, which must be updated by the Manufacturer when changes are made.
- (c) “Common Carrier of Medical Gases or Medical Gas Related Equipment” means any person or entity who undertakes, whether directly or by any other arrangement, to transport, load, or offload property including Medical Gas or Medical Gas Related Equipment for compensation.<sup>179</sup>
- (d) “Designated Representative of Medical Gas or Medical Gas Related Equipment Wholesale Distributors” means any and all individuals designated by the Wholesale Distributor of Medical Gases or Medical Gas

<sup>179</sup> Common carriers frequently use the terms “to load,” which means placing property from the shipping location onto the transport vehicle, and “to offload,” which means removing property from the transport vehicle at the delivery location.

Related Equipment who will serve as a responsible individual of such Wholesale Distributor with the Board who is actively involved in and aware of the actual daily operation of such Wholesale Distributor.

- (e) “Distribute Medical Gas or Medical Gas Related Equipment” or “Distribution of Medical Gas or Medical Gas Related Equipment” means to sell, offer to sell, deliver, offer to deliver, broker, give away, or transfer a Medical Gas or Medical Gas Related Equipment, whether by passage of title, physical movement, or both. The term does not include:
- (1) to Dispense or Administer; or
  - (2) delivering or offering to deliver a Medical Gas or Medical Gas Related Equipment by a common carrier in the usual course of business as a common carrier.
- (f) “Emergency Medical Reasons for the Distribution of Medical Gases or Medical Gas Related Equipment” include, but are not limited to, transfers of a Medical Gas or Medical Gas Related Equipment between a Wholesale Distributor of Medical Gases or Medical Gas Related Equipment or Pharmacy to alleviate a temporary shortage of a Medical Gas or Medical Gas Related Equipment arising from delays in or interruption of regular distribution schedules; sales to nearby emergency medical services, ie, ambulance companies and firefighting organizations in the same state or same marketing or service area, or nearby licensed Practitioners allowed to dispense Medical Gases or Medical Gas Related Equipment for use in the treatment of acutely ill or injured Persons; provision of minimal emergency supplies of Medical Gases or Medical Gas Related Equipment to nearby nursing homes for use in emergencies or during hours of the day when necessary Medical Gases or Medical Gas Related Equipment cannot be obtained; and transfers of Medical Gases or Medical Gas Related Equipment by a retail Pharmacy to another retail Pharmacy to alleviate a temporary shortage.
- (g) “Emergency Use Oxygen” means Oxygen USP administered in emergency situations without a prescription. The container must be labeled in accordance with federal FDA requirements: “For emergency use only when administered by properly trained personnel for oxygen deficiency and resuscitation. For all other medical applications, Rx Only.”
- (h) “FDA” means the Food and Drug Administration, a federal agency within the United States Department of Health and Human Services, established to set safety and quality standards for Drugs, food, cosmetics, and other consumer Products.
- (i) “Federal Act” means the Federal Food, Drug, and Cosmetic Act.
- (j) “Health Care Entity” means any Person that provides diagnostic, medical, surgical, dental treatment, or rehabilitative care, including home respiratory care providers and (in the case of Oxygen USP) to an authorized administrator of “Emergency Use Oxygen,” but does not include any retail Pharmacy or Wholesale Distributor.
- (k) “Immediate Container for Medical Gases” means compressed gas cylinders and liquid containers containing a Medical Gas, but does not include large bulk liquid or high pressure containers such as storage tanks, vehicle mounted vessels, trailers, and/or railcars.
- (l) “Intracompany Transaction” means any transaction between a division, subsidiary, parent, and/or affiliated or related company under the common ownership and control of a corporate entity.
- (m) “Label for Medical Gases” means a display of written, printed, or graphic matter upon the immediate container of any Medical Gas.
- (n) “Label for Medical Gas Related Equipment” means a display of written, printed, or graphic matter upon the immediate container of any Medical Gas Related Equipment.
- (o) “Legally Authorized to Receive” means persons that are licensed Manufacturers of Medical Gases or Medical Gas Related Equipment, Wholesale Distributors of Medical Gases or Medical Gas Related Equipment, home respiratory care companies, and Pharmacies. Also includes Health Care Entities, persons authorized to receive Emergency Use Oxygen without a prescription, and companies that require the use of a Medical Gas in the installation and refurbishment of piping and equipment, including Medical Gas Related Equipment that will be used to distribute or contain a Medical Gas.

- (p) “Medical Gas” means gases (including liquefied gases) classified by FDA as drugs or devices that are used for medical applications and which may be stored and administered through the use of Medical Gas Related Equipment, which may or may not be required under Federal or State law for the immediate container to bear the label, “Rx only” or “Caution: Federal or State law prohibits dispensing without a prescription.”
- (q) “Manufacturer of Medical Gases” means persons manufacturing bulk medical gases or persons transferring gas or liquefied gas product from one container to another (eg, liquid to gas, gas to gas, liquid to liquid).
- (r) “Medical Gas Related Equipment” means a device used as a component part or accessory used to contain or control the flow, delivery, and/or pressure during the Administration of a medical gas (eg, liquid oxygen base and portable units, pressure regulators and flow meters, oxygen concentrators).
- (s) “Misbranded Medical Gas or Medical Gas Related Equipment” means a Medical Gas or Medical Gas Related Equipment shall be deemed to be misbranded if the label is false or misleading in any particular; or the label does not bear the name and address of the Manufacturer, packer, or Distributor and does not have an accurate statement of the quantities of the active ingredients in the case of a Medical Gas; or the label does not show an accurate monograph for the Medical Gas.
- (t) “Prescription Medical Gas” means a Medical Gas which is required under law to be labeled with the following statement: “Rx Only.”
- (u) “Product Labeling” means all labels and other written, printed, or graphic matter upon any article or any of its containers or wrappers, or accompanying such article.
- (v) “USP Standards” means standards published in the current official United States Pharmacopeia or National Formulary.
- (w) “Wholesale Distribution of Medical Gases or Medical Gas Related Equipment” means the Distribution of Medical Gas or Medical Gas Related Equipment, by Wholesale Distributors of Medical Gases or Medical Gas Related Equipment to Persons other than consumers or patients. To the extent permitted by the Prescription Drug Marketing Act, Wholesale Distribution of Medical Gases, or Medical Gas Related Equipment does not include:
  - (1) the sale, purchase, or trade of a Medical Gas or Medical Gas Related Equipment, an offer to sell, purchase, or trade a Prescription Drug or Device, or the Dispensing of a Medical Gas or Medical Gas Related Equipment pursuant to a Prescription;
  - (2) the sale, purchase, or trade of a Medical Gas or Medical Gas Related Equipment or an offer to sell, purchase, or trade a Medical Gas or Medical Gas Related Equipment for Emergency Medical Reasons;
  - (3) intracompany Transactions, unless in violation of own use provisions;
  - (4) the sale, purchase, or trade of a Medical Gas or Medical Gas Related Equipment or an offer to sell, purchase, or trade a Medical Gas or Medical Gas Related Equipment among hospitals, Pharmacies, or other health care entities that are under common control;
  - (5) the sale, purchase, or trade of a Medical Gas or Medical Gas Related Equipment or the offer to sell, purchase, or trade a Medical Gas or Medical Gas Related Equipment by a charitable organization described in 503(c)(3) of the Internal Revenue Code of 1954 to a nonprofit affiliate of the organization to the extent otherwise permitted by law;
  - (6) the purchase or other acquisition by a hospital or other similar health care entity that is a member of a group purchasing organization of a Medical Gas or Medical Gas Related Equipment for its own use from the group purchasing organization or from other hospitals or similar health care entities that are members of these organizations;
  - (7) the Return of residual Medical Gas that may be reprocessed in accordance with Manufacturer’s procedures, or the Return of recalled, expired, damaged, or otherwise non-salable Medical Gas or Medical Gas Related Equipment, when conducted by a hospital, health care entity, Pharmacy, or charitable institution in accordance with the Board’s regulations; or
  - (8) other Transactions excluded from the definition of “wholesale distribution” under 21 CFR 203.3(CC), including any amendments thereto.

- (x) “Wholesale Distributor of Medical Gases or Medical Gas Related Equipment” means any Person engaged in Wholesale Distribution of Medical Gas or Medical Gas Related Equipment in or into the State, including but not limited to Manufacturers, own-label distributors, private-label distributors, warehouses, including Manufacturers’ and Distributors’ warehouses, and Wholesale Medical Gas or Medical Gas Related Equipment warehouses.

## **Section 2. Requirements for Licensure.**

Wholesale Distributors of Medical Gases or Medical Gas Related Equipment that reside in this state and provide services within this state or other states shall be licensed by the Board and shall periodically renew their license with the Board using an application provided by the Board. Wholesale Distributors of Medical Gases or Medical Gas Related Equipment that provide services within this state though are not residents of this state shall maintain a valid license with the state Board in which they reside and in all states in which they distribute, if required. Wholesale Distributors cannot operate from a place of residence, except when that place of residence is used for “on call” delivery of homecare oxygen and oxygen related equipment by a home respiratory care technician. Where Wholesale Distribution operations are conducted at more than one location within this state, each such location shall be licensed by the Board of Pharmacy.

- (a) Subject to the Federal Act and all applicable federal law and regulations, an FDA-registered Medical Gas or Medical Gas Related Equipment manufacturer, including its affiliates, subsidiaries, agents, and other entities under common ownership and control of the Manufacturer, that exclusively distributes its own Medical Gas or Medical Gas Related Equipment, may be exempted from the requirements for licensure.
- (b) Every Wholesale Distributor who engages in the Wholesale Distribution of Medical Gases or Medical Gas Related Equipment shall license with the Board by application and provide information required by the Board on an application approved by the Board, including but not limited to:
- (1) all trade or business names used by the licensee (includes “doing business as (dba)” and “formerly known as”), which cannot be identical to the name used by another unrelated Wholesale Distributor licensed to purchase Medical Gas or Medical Gas Related Equipment in the State;
  - (2) name(s) of the owner and operator of the licensee (if not the same person), including:<sup>180</sup>
    - (i) if a Person: the name, business address, Social Security number, and date of birth;
    - (ii) if a partnership: the name, business address, and Social Security number, and date of birth of each partner, the name of the partnership, and federal employer identification number;
    - (iii) if a corporation: the name, business address, and title of each corporate officer and director, the corporate names, the state of incorporation, federal employer identification number, and the name and business address of the parent company, if any;
    - (iv) if a sole proprietorship: the full name and business address of the sole proprietor and the name and federal employer identification number of the business entity;
    - (v) if a limited liability company: the name, business address, and title of each company officer, the name of the limited liability company and federal employer identification number, and the name of the state in which the limited liability company was organized; and
    - (vi) any other relevant information that the Board requires.
  - (3) name(s), business address(es), and telephone number(s) of a person(s) to serve as the Designated Representative(s) for each facility of the Wholesale Distributor that engages in the Wholesale Distribution of Medical Gas /or Medical Gas Related Equipment and additional information as required in Section 10 (Record Keeping);

---

<sup>180</sup> The risk of diversion and adulteration are not concerns for medical gases. With this in mind, the depth of personal identification information required for licensure of Wholesale Distributors of Medical Gases or Medical Gas Related Equipment is less than that of Wholesale Distributors of Prescription Drugs. In addition, the provision of facility details such as square footage, lease details, and temperature and humidity controls is not required as it is for Wholesale Distributors of Prescription Drugs.

- (4) a list of all State and Federal licenses, registrations, or permits, including the license, registration, or permit numbers issued to the Wholesale Distributor by any other state and federal authority that authorizes the Wholesale Distributor to purchase, possess, and Wholesale Distributes Medical Gas or Medical Gas Related Equipment in this state;
  - (5) a list of all disciplinary actions pertinent to Wholesale Distributors of Medical Gases or Medical Gas Related Equipment by any State and Federal agencies against the Wholesale Distributor distributing Medical Gas or Medical Gas Related Equipment into the state as well as any such actions against principals, owners, directors, or officers;
  - (6) an address and description of each facility and warehouse, including all locations utilized for Medical Gas or Medical Gas Related Equipment storage or Wholesale Distribution including a description of the security system;
  - (7) information regarding general and Product liability insurance, including copies of relevant policies;
  - (8) a description of import and export activities;
  - (9) a copy of the Wholesale Distributor's written policies and procedures as required in Section 11 (Policies and Procedures); and
  - (10) the information collected by the Board pursuant to Section 1(a)(6) shall be made available only to the Board, a third party recognized by the Board, and to State and Federal law enforcement officials. The Board shall make provisions for protecting the confidentiality of the information collected under this section.
- (c) A "surety" bond of not less than \$100,000, or other equivalent means of security acceptable to the Board or a third party recognized by the Board such as insurance, an irrevocable letter of credit, or funds deposited in a trust account or financial institution, to secure payment of any administrative penalties imposed by the Board and any fees or costs incurred by the Board regarding that licensee when those penalties, fees, or costs are authorized under state law and the licensee fails to pay thirty (30) days after the penalty, fee, or costs becomes final. A separate "surety" bond or other equivalent means of security is not required for each company's separate locations or for affiliated companies/groups when such separate locations or affiliated companies/groups are required to apply for or renew their Wholesale Distributor of Medical Gases or Medical Gas Related Equipment license with the Board. The Board may make a claim against such bond or other equivalent means of security until one year after the Wholesale Distributor's license ceases to be valid or until sixty (60) days after any administrative or legal proceeding before or on behalf of the Board that involves the Wholesale Distributor is concluded, including any appeal, whichever occurs later. Manufacturers of Medical Gases shall be exempt from securing a "surety" bond or other equivalent means of security acceptable to the Board. The Board may waive the bond requirement, if the Wholesale Distributor of Medical Gases or Medical Gas Related Equipment:
- (1) has previously obtained a comparable surety bond or other equivalent means of security for the purpose of licensure in another state, where the Wholesale Distributor possesses a valid license in good standing; or
  - (2) is a publicly held company.
- (d) Every Wholesale Distributor of Medical Gases or Medical Gas Related Equipment who engages in Wholesale Distribution of Medical Gases or Medical Gas Related Equipment shall submit a reasonable fee to be determined by the Board.
- (e) Manufacturing facilities of Medical Gases are exempt from inspection by the Board, if the Manufacturing facilities:
- (1) are currently registered with FDA in accordance with Section 510 of the Federal Act and can provide proof of such registration, such as a copy of the online verification page; and
  - (2) can provide proof of inspection by the FDA, or other regulatory body within the past three (3) years.
- (f) The Board may require each facility that engages in Wholesale Distribution of Medical Gases or Medical Gas Related Equipment to undergo an inspection in accordance with Section 15 of this rule and in accordance

with a schedule to be determined by the Board. Wholesale Distributors of Medical Gas or Medical Gas Related Equipment do not qualify for the Drug Distributor Accreditation program.<sup>181</sup>

- (g) All Wholesale Distributors of Medical Gases or Medical Gas Related Equipment must publicly display or have readily available all state licenses and the most recent inspection report administered by the Board.
- (h) Changes in any information in this Section shall be submitted to the Board, or to a third party recognized by the Board, within 30 days of such change (unless otherwise noted).
- (i) Information submitted by the Wholesale Distributor of Medical Gases or Medical Gas Related Equipment to the Board or a third party recognized by the Board that is considered trade secret or proprietary information, as defined under this State's privacy and trade secret/proprietary statutes, shall be maintained by the Board or a third party recognized by the Board as private or trade secret/proprietary information and be exempt from public disclosure.

### **Section 3. Minimum Qualifications.**

- (a) The Board will consider the following factors in determining the eligibility for, and renewal of, licensure of Persons who engage in the Wholesale Distribution of Medical Gas or Medical Gas Related Equipment:
  - (1) any findings by the Board that the applicant has violated or been disciplined by a regulatory agency in any state for violating any Federal, State, or local laws relating to or the Wholesale Distribution of Medical Gases or Medical Gas Related Equipment;
  - (2) any criminal convictions of the applicant under Federal, State, or local laws;
  - (3) the applicant's past experience in the Manufacture or Wholesale Distribution of Medical Gases or Medical Gas Related Equipment;
  - (4) the furnishing by the applicant of false or fraudulent material in any application made in connection with the or Manufacturing or Wholesale Distribution of Medical Gases or Medical Gas Related Equipment;
  - (5) Suspension, sanction, or Revocation by federal, State, or local government against any license currently or previously held by the applicant or any of its owners for violations of State or Federal laws regarding Medical Gas or Medical Gas Related Equipment;
  - (6) compliance with previously granted licenses of any kind;
  - (7) compliance with the requirements to maintain and/or make available to the Board licensing authority or to Federal, State, or local law enforcement officials those records required to be maintained by Wholesale Distributors of Medical Gases or Medical Gas Related Equipment; and
  - (8) any other factors or qualifications the Board considers relevant to and consistent with the public health and safety.
- (b) The Board shall consider the results of a criminal and financial background check of the applicant, including but not limited to, all key personnel involved in the operations of the Wholesale Distributor of Medical Gases or Medical Gas Related Equipment, including the most senior Person responsible for facility operations, purchasing, and inventory control and the Person or Persons they report to; and all company officers, key management, principals, and owners with ten percent (10%) or greater ownership interest in the company (applying to non-publicly held companies only) to determine if an applicant or others associated with the ownership, management, or operations of the Wholesale Distributor of Medical Gases or Medical Gas Related Equipment have committed criminal acts that would constitute grounds for denial of licensure. The background check will be conducted in compliance with any applicable state and Federal laws, at the applicant's expense, and will be sufficient to include all states of residence since the Person has been an adult. Manufacturers of Medical Gases or Medical Gas Related Equipment shall be exempt from criminal and financial background checks.

---

<sup>181</sup> Although a Board may allow a firm to be third-party accredited, Wholesale Distributors of Medical Gases or Medical Gas Related Equipment do not qualify for the NABP Drug Distributor Accreditation program as the inspection criteria is not applicable to Medical Gas or Medical Gas Equipment Related operations.

- (c) The applicant shall provide, and attest to, a statement providing a complete disclosure of any past criminal convictions and violations of the State and Federal laws regarding Medical Gases or Medical Gas Related Equipment or an affirmation and attestation that the applicant has not been involved in, or convicted of, any criminal or prohibited acts.

#### **Section 4. Personnel.**

Each Person that is issued an initial or renewal license as a Wholesale Distributor of Medical Gases or Medical Gas Related Equipment, whether in state or out of state, must designate in writing, Person(s) for each facility to serve as Designated Representatives of such Wholesale Distributor. The members of the quality control unit, per 21 CFR 211.22, shall act as the Designated Representatives for the Wholesale Distributer.

- (a) To be certified as a Designated Representative for a Wholesale Distributor of Medical Gases or Medical Gas Related Equipment, a Person:
  - (1) must have the appropriate amount of education, training and experience or any combination thereof to perform the functions required to serve as the Designated Representative of such Wholesale Distributor; and
  - (2) must be actively involved in and aware of the daily operations of the Wholesale Distributor location(s) including all policies and procedures pertaining to those operations and may cover multiple locations. The Designated Representative is therefore not required to be present at each site during normal business hours.
- (b) The information collected pursuant to Section 3(a) shall be made available only to the Board, a third party recognized by the Board, and to State and Federal law enforcement officials. The Board and a third party recognized by the Board shall make provisions for protecting the confidentiality of the information collected under this section.
- (c) Each licensed Wholesale Distributor of Medical Gases or Medical Gas Related Equipment located outside of this State that Wholesale Distributes Medical Gases or Medical Gas Related Equipment in this State shall designate a registered agent in this State for service of process. Any licensed Wholesale Distributor of Medical Gases or Medical Gas Related Equipment that does not so designate a registered agent shall be deemed to have designated the Secretary of State of this State to be its true and lawful attorney, upon who may be served all legal processes in any action or proceeding against such licensed Wholesale Distributor growing out of or arising from such Wholesale Distribution. A copy of any such service of process shall be mailed to such Wholesale Distributor by the Board by certified mail, return receipt requested, postage prepaid, at the address such licensed Wholesale Distributor has designated on its application for licensure in this State. If any such Wholesale Distributor is not licensed in this State, service on the Secretary of State only shall be sufficient service.
- (d) A Designated Representative must complete either:
  - (1) continuing education programs specified by the Board regarding Federal and State laws in regard to the Wholesale Distribution, handling, and storage of Medical Gases or Medical Gas Related Equipment; or
  - (2) training programs that address applicable Federal and State laws and are provided by qualified in-house specialists, outside counsel, or consulting specialists with capabilities to help ensure compliance.

#### **Section 5. Minimum Requirements for the Storage and Handling of Medical Gases or Medical Gas Related Equipment and for Establishment and Maintenance of Medical Gas or Medical Gas Related Equipment Records.**

The following are required for the storage, handling, transport, and shipment of Medical Gases or Medical Gas Related Equipment and for the establishment and maintenance of Wholesale Distribution records by Wholesale

Distributors of Medical Gases and Medical Gas Related Equipment and their officers, agents, representatives, and employees.

- (a) All facilities at which a Medical Gas or Medical Gas Related Equipment is received, stored, warehoused, handled, held, offered, marketed, displayed, or transported from shall:
  - (1) be of suitable construction to ensure that all Medical Gases or Medical Gas Related Equipment in the facilities are maintained in accordance with the Product Labeling of such Medical Gas or Medical Gas Related Equipment, or in compliance with official compendium standards such as the USP-NF;
  - (2) be of suitable size and construction to facilitate cleaning, maintenance, and proper Wholesale Distribution operations;
  - (3) have adequate storage areas with appropriate lighting, ventilation, sanitation, space, equipment, and security conditions;
  - (4) have a quarantine area for storage of Medical Gas or Medical Gas Related Equipment that are suspected of being outdated, Misbranded, or Adulterated, or otherwise unfit for Distribution or Wholesale Distribution;
  - (5) be maintained in a clean and orderly condition;
  - (6) be free from infestation that may impact the identity, strength, quality, or purity of the Medical Gas;
  - (7) be a commercial location and not a personal dwelling or residence, except when that personal dwelling is used for “on call” delivery of Oxygen USP and oxygen related equipment for homecare use;<sup>182</sup>
  - (8) provide for the secure and confidential storage of information with restricted access and policies and procedures to protect the integrity and confidentiality of the information; and
  - (9) provide and maintain appropriate inventory controls in order to detect and document any theft of nitrous oxide.

## **Section 6. Security.**

- (a) All facilities used for Wholesale Distribution of Medical Gases or Medical Gas Related Equipment shall be secure from unauthorized entry:
  - (1) access from outside the premises shall be kept to a minimum and be well-controlled;
  - (2) the outside perimeter of the premises shall be well-lighted; and
  - (3) entry into areas where Medical Gas or Medical Gas Related Equipment are held shall be limited to authorized personnel; all facilities shall be equipped with a system to detect or deter entry after hours.
- (b) All facilities shall be equipped with a system that will provide suitable protection against theft. When appropriate, the system shall provide protection against theft that is facilitated or hidden by tampering with computers or electronic records.
- (c) All facilities shall be equipped with inventory management and control systems that protect against, detect, and document any instances of theft of nitrous oxide.
- (d) Where Wholesale Distributors of Medical Gases or Medical Gas Related Equipment use electronic distribution records, they shall employ, train, and document the training of personnel in the proper use of such technology and equipment.
- (e) All facilities shall be equipped with security systems to protect the integrity and confidentiality of data and documents and make such data and documents readily available to the Board and other state and federal law enforcement officials.
- (f) Vehicles utilized for on-call delivery of Oxygen USP and oxygen related equipment for home care use by home care providers may be parked at a place of residence and shall be locked and equipped with an audible alarm while not attended.

---

<sup>182</sup> Some home respiratory care providers provide “on call” services to patients. This requires home respiratory care technicians to keep parked at their personal dwelling the company vehicle stocked with Medical Gases or Medical Gas Related Equipment.



- (g) All Wholesale Distributors of Medical Gases or Medical Gas Related Equipment shall maintain records documenting from whom Medical Gases or Medical Gas Related Equipment are received and to whom Medical Gases and/or Medical Gas Related Equipment are distributed with information sufficient to perform a recall of Medical Gases or Medical Gas Related Equipment received and distributed in compliance with 21 CFR 150b, 21 CFR 211.196, and 21 CFR 820.160b.

## **Section 7. Storage.**

All Medical Gases or Medical Gas Related Equipment shall be stored under appropriate conditions in accordance with regulations or, in the absence of regulations, in accordance with applicable industry standards, and the manufacturers' recommendations on the Product labeling.

- (a) Packaging of the Medical Gas or Medical Gas Related Equipment should be in accordance with an official compendium such as USP-NF, if applicable.
- (b) The record-keeping requirements in Section 10 (Record Keeping) shall be followed for the Wholesale Distribution of all Medical Gases or Medical Gas Related Equipment.

## **Section 8. Examination of Materials.**

- (a) Upon receipt, each Medical Gas container and related equipment shall be visually examined for identity and to determine if it is damaged or otherwise unfit for Wholesale Distribution. This examination shall be adequate to reveal container damage that would suggest possible Adulteration or Misbranding.
- (b) The Medical Gas or Medical Gas Related Equipment found to be unacceptable under paragraph (a) should be quarantined from the rest of stock until the examination and determination that the Medical Gas or Medical Gas Related Equipment are not Misbranded or Adulterated.
- (c) Each outgoing shipment shall be carefully inspected for identity of the Medical Gas or Medical Gas Related Equipment and to ensure that there is no Delivery of Medical Gas or Medical Gas Related Equipment that have been damaged in storage or held under improper conditions.
- (d) Upon receipt, a Wholesale Distributor of Medical Gases or Medical Gas Related Equipment must review records for the acquisition of Medical Gases or Medical Gas Related Equipment for accuracy and completeness.
- (e) The record-keeping requirements in Section 10 (Record Keeping) shall be followed for all incoming and outgoing Medical Gases or Medical Gas Related Equipment.

## **Section 9. Returned, Damaged, and Outdated Medical Gases or Medical Gas Related Equipment.**

- (a) Medical Gas that has left the control of the Wholesale Distributor may be returned to the Wholesale Distributor or Manufacturer from which it was acquired but may not be resold as a Medical Gas even if the integrity of the Product is maintained, unless it is reprocessed by the Manufacturer employing proper and adequate controls to ensure the identity, strength, quality, and purity of the reprocessed Medical Gas.
- (b) Reusable Medical Gas Related Equipment that has left the control of the Wholesale Distributor may be returned to Wholesale Distributor or Manufacturer for inspection. The Medical Gas Related Equipment may be repaired and or refurbished, if necessary, provided the Manufacturer or Wholesale Distributor employs proper and adequate controls to return the Medical Gas Related Equipment to proper condition.
- (c) Any Medical Gas, including its container, that is damaged, Misbranded, or Adulterated shall be quarantined and physically separated from other Medical Gases until it is destroyed or returned to either the Manufacturer or Wholesale Distributor from which it was acquired. External contamination to Medical Gas containers or closure system, not impacting the integrity of the Medical Gas, is not considered damage or Adulteration for purposes of this paragraph. When Medical Gas or Medical Gas Related Equipment are Adulterated, Misbranded, or suspected of being Adulterated, or Misbranded, notice of the Adulteration, Misbranding, or suspected Adulteration, or Misbranding shall be provided to the manufacturer or wholesale distributor from which they were acquired and also the appropriate boards and federal regulatory bodies.

- (d) Any Medical Gas container that has been opened or used, but is not Adulterated or Misbranded, shall be considered empty, quarantined and physically separated from non-empty Medical Gas containers and returned to the Manufacturer for destruction or reprocessing.
- (e) Any Medical Gas, its container, or Medical Gas Related Equipment including its associated documentation or labeling, suspected of being involved in a criminal activity shall be retained and not destroyed until its disposition is authorized by the Board, or applicable law enforcement agency.
- (f) The record-keeping requirements in Section 10 (Record Keeping) of this rule shall be followed for all Misbranded or Adulterated Medical Gases.

## **Section 10. Due Diligence.**

A Wholesale Distributor of Medical Gases or Medical Gas Related Equipment licensed in accordance with these Rules shall comply with the following Due Diligence requirements:

- (a) Prior to the initial Wholesale Distribution or acquisition of a Medical Gases or Medical Gas Related Equipment to or from any Wholesale Distributor (or prior to any Wholesale Distribution to a Wholesale Distributor by a Manufacturer), the Distributing Wholesale Distributor (or Manufacturer) shall provide the following information to the acquiring Wholesale Distributor:
  - (1) If a Manufacturer is distributing to a Wholesale Distributor, evidence that the Manufacturer is registered, and the Medical Gas or Medical Gas Related Equipment is listed with FDA;
  - (2) If a Wholesale Distributor is distributing to a Wholesale Distributor, evidence that the Wholesale Distributor supplying the Medical Gas or Medical Gas Related Equipment is licensed to provide Product into the State, if required by the State;
  - (3) the name(s) of the responsible facility contact person(s) at the supplying Manufacturer or Wholesale Distributor; and
  - (4) a certification that the Manufacturer or Wholesale Distributor's policies and procedures comply with this Act.
- (b) A Manufacturer or Wholesale Distributor that Wholesale Distributes or acquires Medical Gases or Medical Gas Related Equipment to or from another Wholesale Distributor of Medical Gases or Medical Gas Related Equipment shall provide to or obtain from the distributing or acquiring entities as applicable the information set forth in Section 10 (Record Keeping).
- (c) Wholesale Distributors of Medical Gases or Medical Gas Related Equipment are exempt from inspecting and obtaining the information from Manufacturers of Medical Gases or Medical Gas Related Equipment as required in Section 9 (Due Diligence) when the Manufacturer is registered with FDA in accordance with Section 510 of the Federal Act and can:<sup>183</sup>
  - (1) provide proof of such registration; and
  - (2) either:
    - (i) can provide proof of inspection by the FDA, or other regulatory body within the past three (3) years; or
    - (ii) in the event that no regulatory body has inspected within the past three (3) years, conformance with industry standards or guidelines, as identified by the Board.

## **Section 11. Record Keeping.**

- (a) Wholesale Distributors of Medical Gases or Medical Gas Related Equipment shall establish and maintain records of all Transactions regarding the receipt and Wholesale Distribution or other disposition of Medical Gases or Medical Gas Related Equipment. These records shall include:
  - (1) dates of receipt and Wholesale Distribution or other disposition of the Medical Gas or Medical Gas Related Equipment; and

---

<sup>183</sup> The Board may refer to the following industry guideline: CGA M-7, *Guideline for Qualifying Suppliers Used by Medical Gas Manufacturers and Distributors*.

- (2) Information sufficient to perform a recall of Medical Gases or Medical Gas Related Equipment received and distributed.
- (b) Such records shall be made available for inspection and photocopying by any authorized official of any State, Federal, or local governmental agency for a period of:<sup>184</sup>
  - (1) three (3) years following their creation date for high pressure Medical Gases;
  - (2) one (1) year following their creation date for cryogenic or refrigerated liquid Medical Gases; and
  - (3) three (3) years following their creation date for Medical Gas Related Equipment.
- (c) Records described in this section that are kept at the inspection site or that can be immediately retrieved by computer or other electronic means shall be readily available for authorized inspection during the retention period. Records kept at a central location apart from the inspection site and not electronically retrievable shall be made available for inspection within two (2) working days of a request by an authorized official of any State or Federal governmental agency charged with enforcement of these rules.
- (d) Wholesale Distributors and Manufacturers of Medical Gases or Medical Gas Related Equipment should maintain an ongoing list of Persons from whom they receive or to whom they distribute Medical Gases or Medical Gas Related Equipment.
- (e) Wholesale Distributors of Medical Gases or Medical Gas Related Equipment shall maintain a system for the mandatory reporting of any theft, suspected theft, or other significant loss of Nitrous Oxide to the Board and other appropriate law enforcement agencies.

## **Section 12. Policies and Procedures.**

Wholesale Distributors of Medical Gases or Medical Gas Related Equipment shall establish, maintain, and adhere to written policies and procedures, which shall be followed for the receipt, security, storage, transport, and shipping and Wholesale Distribution of Medical Gases or Medical Gas Related Equipment, including policies and procedures for maintaining inventories, identifying, recording, and reporting losses or thefts and for correcting all errors and inaccuracies in inventories associated with nitrous oxide. Wholesale Distributors of Medical Gases or Medical Gas Related Equipment shall include in their written policies and procedures the following:

- (a) A procedure to be followed for handling recalls and withdrawals of Medical Gases or Medical Gas Related Equipment. Such procedure shall be adequate to deal with recalls and withdrawals due to:
  - (1) Any action initiated at the request of FDA or any other federal, state, or local law enforcement or other government agency, including the Board of Pharmacy; or
  - (2) Any volunteer action by the Manufacturer of Medical Gases or Medical Gas Related Equipment to remove defective or potentially defective Medical Gases or Medical Gas Related Equipment from the market.
- (b) A procedure to ensure that Wholesale Distributors of Medical Gases or Medical Gas Related Equipment prepare for, protect against, and handle any crisis that affects security or operation of any facility in the event of a strike, fire, flood, or other natural disaster, or other situations of local, State, or national emergency.
- (c) A procedure for reporting criminal or suspected criminal activities involving the inventory of nitrous oxide to the Board, and applicable law enforcement agencies, within three (3) business days of becoming aware of the criminal or suspect criminal activity.
- (d) A procedure for verifying security provisions of Common Carriers.

---

<sup>184</sup> Record retention requirements are determined based on cryogenic and liquefied gas product profiles.

### **Section 13. Prohibited Acts.**

It is unlawful for a Person to knowingly and willfully perform or cause the performance of or aid and abet any of the following acts in this State:

- (a) the Manufacture, Repackaging, sale, delivery, or holding or offering for sale any Medical Gas or Medical Gas Related Equipment that is Adulterated, Misbranded, or has otherwise been rendered unfit for Distribution or Wholesale Distribution;
- (b) the Adulteration, or Misbranding of any Medical Gas or Medical Gas Related Equipment;
- (c) the receipt of any Medical Gas or Medical Gas Related Equipment that is Adulterated, Misbranded, stolen, obtained by fraud or deceit, or the delivery or proffered delivery of such Medical Gas or Medical Gas Related Equipment for pay or otherwise;
- (d) the Alteration, mutilation, destruction, obliteration, or removal of the whole or any part of the Product Labeling of a Medical Gas or Medical Gas Related Equipment or the willful commission of any other act with respect to a Medical Gas or Medical Gas Related Equipment that results in the Medical Gas or Medical Gas Related Equipment being Misbranded;
- (e) the purchase or receipt of a Medical Gas or Medical Gas Related Equipment from a Person that is not licensed to Wholesale Distribute Medical Gas or Medical Gas Related Equipment to that purchaser or recipient;
- (f) the sale or transfer of a Medical Gas or Medical Gas Related Equipment to a Person who is not legally authorized to receive a Medical Gas or Medical Gas Related Equipment;
- (g) the failure to maintain or provide records as required by this Act and Rules;
- (h) providing the Board or any of its representatives or any state or federal official with false or fraudulent records or making false or fraudulent statements regarding any matter within the provisions of this Act and Rules;
- (i) the Wholesale Distribution of any Medical Gas or Medical Gas Related Equipment that was:
  - (1) purchased by a public or private hospital or other health care entity;
  - (2) donated or supplied at a reduced price to a charitable organization; or
  - (3) stolen or obtained by fraud or deceit.
- (j) the failure to obtain a license or operating without a valid license when a license is required;
- (k) the Obtaining of or attempting to obtain a Medical Gas or Medical Gas Related Equipment by fraud, deceit, misrepresentation, or engaging in misrepresentation or fraud in the Distribution or Wholesale Distribution of a Medical Gas/or Medical Gas Related Equipment;
- (l) the Distributing of a Medical Gas or Medical Gas Related Equipment to a patient without a Prescription or Prescription Order from a Practitioner licensed by law to use or prescribe the Medical Gas or Medical Gas Related Equipment;
- (m) the Distributing or Wholesale Distributing of a Medical Gas or Medical Gas Related Equipment that was previously dispensed by a Pharmacy or distributed by a Practitioner;
- (n) the Distributing of a Medical Gas or Medical Gas Related Equipment to a patient without providing appropriate information and counseling on use, storage, and disposal;
- (o) the failure to report any Prohibited Act as listed in these Rules; or
- (p) the failure to exercise Due Diligence as provided in Section 9 (Due Diligence) of these regulations.

### **Section 14. Criminal Acts.**

- (a) A Person who, with intent to defraud or deceive, performs the act of Adulteration or Misbranding of any Medical Gas or Medical Gas Related Equipment commits a felony of the third degree.
- (b) A Person who engages in the Wholesale Distribution and knowingly purchases or receives Medical Gas or Medical Gas Related Equipment from a Person, not legally authorized to Wholesale Distribute Medical Gas or Medical Gas Related Equipment, in Wholesale Distribution commits a felony of the third degree.

- (c) A Person who engages in the Wholesale Distribution and knowingly sells, barter, brokers, or transfers Medical Gases or Medical Gas Related Equipment to a Person not legally authorized to purchase Medical Gases or Medical Gas Related Equipment, under the jurisdiction in which the Person receives the Medical Gas or Medical Gas Related Equipment in Wholesale Distribution, commits a felony of the third degree.
- (d) A Person who knowingly falsely creates any Label for a Medical Gas or Medical Gas Related Equipment or who falsely represents any factual matter contained in any Label of a Medical Gas or Medical Gas Related Equipment commits a felony of the third degree.
- (e) A Person found guilty of any offense under this section, under the authority of the Court convicting and sentencing the Person, shall be ordered to forfeit to the State any real or Personal property:
  - (1) used or intended to be used to commit, to facilitate, or to promote the commission of such offense; and
  - (2) constituting, derived from, or traceable to the gross proceeds that the defendant obtained directly or indirectly as a result of the offense. Any property or assets subject to forfeiture under this section may be seized pursuant to a warrant obtained in the same manner as a search warrant or as otherwise permitted by law, and held until the case against a defendant is adjudicated. Monies ordered forfeited, or proceeds from the sale of other assets ordered forfeited, shall be equitably divided between the Board and other agencies involved in the investigation and prosecution that led to the conviction. Other property ordered forfeited after conviction of a defendant may, at the discretion of the investigating agencies, be placed into official use by the Board or the agencies involved in the investigation and prosecution that led to the conviction.

#### **Section 15. Salvaging and Reprocessing.**

- (a) Medical Gas or Medical Gas Related Equipment that has been subjected to improper conditions such as a fire, accident or natural disaster, shall not be Salvaged or Reprocessed;
- (b) Medical Gas product in a Medical Gas container that has left the control of the Wholesale Distributor may be returned to the Manufacturer and reprocessed provided the Manufacturer employs proper and adequate controls to assure the identity, strength, quality, and purity of the reprocessed Medical Gas; and
- (c) Reusable Medical Gas Related Equipment that has left the control of the Wholesale Distributor may be returned to the Wholesale Distributor or Manufacturer for inspection. The Medical Gas Related Equipment may be repaired and or refurbished (servicing), if necessary, provided the Manufacturer or Wholesale Distributor employs proper and adequate controls to ensure the Medical Gas Related Equipment complies with the manufacturers' design and performance specifications following completion of servicing.

#### **Section 16. Inspection.**

- (a) The Board shall have the authority to recognize a third party to inspect Wholesale Distributors of Medical Gases or Medical Gas Related Equipment in that State or in other State(s).
- (b) The Board shall have the authority to recognize other State(s) inspections of Wholesale Distributors of Medical Gases or Medical Gas Related Equipment operations in other State(s), if such state's laws are deemed to be substantially equivalent.
- (c) The Board may license by reciprocity, a Wholesale Distributor of Medical Gases or Medical Gas Related Equipment that is licensed under the laws of another state, if the requirements of that State are deemed by the Board to be substantially equivalent.
- (d) Any applicant that is denied a license due to an inspection shall have the right of review of the Board's decision.
- (e) The Board shall ensure that the proprietary information obtained during the inspection process remains confidential and privileged.
- (f) The Board may waive requirements of this Chapter.

## **Appendix A**

### **Multistate Pharmacy Inspection Blueprint**

#### **1.1.0 Practice**

- 1.1.1 Classification
  - 1.1.1.1 Traditional Community
  - 1.1.1.2 Mail Order
  - 1.1.1.3 Manufacturer
  - 1.1.1.4 Compounding
    - 1.1.1.4.1 Nonsterile Compounding
    - 1.1.1.4.2 Sterile Compounding
    - 1.1.1.4.3 Nonsterile HD Compounding
    - 1.1.1.4.4 Sterile HD Compounding
- 1.1.2 Institutional
- 1.1.3 Closed Door
  - 1.1.3.1 Long-Term Care
- 1.1.4 HMO/PBM only
  - 1.1.4.1 Internet Pharmacy
  - 1.1.4.2 Telepharmacy
- 1.1.5 Central Fill/Processing/Workload Balancing/Shared Services
- 1.1.6 Outsourcing Facility
- 1.1.7 Wholesale Distributor
- 1.1.8 Clinical Trials/Research
  - 1.1.8.1 Medical Marijuana
- 1.1.9 Nuclear
- 1.1.10 Veterinary
- 1.1.11 Specialty
- 1.1.12 Specify for any of the classifications above
  - 1.1.12.1 Mail/Deliver (in-state)
  - 1.1.12.2 Mail/Deliver (out of state, please list below)

#### **1.2.0 Other Practices to note**

- 1.2.1 Provide products for “Office Use”
  - 1.2.1.1 Patient specific for administration
  - 1.2.1.2 Any non-patient specific for administration
- 1.2.2 Wholesale distribution (less than 5%)

#### **1.3.0 Business Information**

- 1.3.1 Business Name
  - 1.3.1.1 DBA
  - 1.3.1.2 Street Address
  - 1.3.1.3 City, State, Zip
  - 1.3.1.4 Telephone Number

- 1.3.1.5 Toll-free Number
- 1.3.1.6 Fax Number
- 1.3.1.7 Email Address

#### **1.4.0 Website\***

- 1.4.1 Operational Website
- 1.4.2 Affiliated Websites

#### **1.5.0 Personnel**

- 1.5.1 Pharmacists
  - 1.5.1.1 PIC/Supervising Pharmacist
  - 1.5.1.2 PIC email
  - 1.5.1.3 Nonresident PIC (if applicable)
  - 1.5.1.4 Compounding Pharmacists
  - 1.5.1.5 Total Pharmacists
  - 1.5.1.6 Total Pharmacist Hours Per Week
- 1.5.2 Technicians
  - 1.5.2.1 Technicians in Training
  - 1.5.2.2 Registered/Licensed Technicians
  - 1.5.2.3 Nationally Certified Technicians
  - 1.5.2.4 Compounding Technicians
  - 1.5.2.5 Total Technicians
  - 1.5.2.6 Total Technician Hours Per Week
  - 1.5.2.7 Ratio at time of inspection
- 1.5.3 Interns/Students
  - 1.5.3.1 Total Student Interns
  - 1.5.3.2 Total Graduate Interns
- 1.5.4 Other Personnel
  - 1.5.4.1 Nonsterile Compounding Manager
  - 1.5.4.2 Sterile Compounding Manager
  - 1.5.4.3 Hazardous Compounding Supervisor
  - 1.5.4.4 Total Other Personnel

### ***Section 2.0.0 State and Federal Licensure/Registration Information of State of Residence***

#### **2.1.0 Types of Licensure and/or Registration**

- 2.1.1 State permits, registrations, and licenses
  - 2.1.1.1 Business Name on License/Registration
  - 2.1.1.2 License/Registration Agency
  - 2.1.1.3 License/Registration Number
  - 2.1.1.4 Expiration Date
  - 2.1.1.5 Frequency of renewal (annual/bi/tri)
- 2.1.2 Federal permits, registration, and licenses
  - 2.1.2.1 DEA

- 2.1.2.1.1 Business Name on License/Registration
- 2.1.2.1.2 License/Registration Agency
- 2.1.2.1.3 License/Registration Number
- 2.1.2.1.4 Expiration Date
- 2.1.2.1.5 Restrictions/limitations/waivers
- 2.1.2.2 FDA
  - 2.1.2.2.1 Business Name on License/Registration
  - 2.1.2.2.2 License/Registration Agency
  - 2.1.2.2.3 License/Registration Number
  - 2.1.2.2.4 Expiration Date
- 2.1.2.3 Other businesses at that address
  - 2.1.2.3.1 Type of business

## **2.2.0**

### ***Inspections***

- 2.2.1 State
  - 2.2.1.1 Inspection/Response (dates)
  - 2.2.1.2 Frequency of inspection
  - 2.2.1.3 Warning Letters/Response
  - 2.2.1.4 Consent/Response
- 2.2.2 Federal
  - 2.2.2.1 DEA
    - 2.2.2.1.1 Inspection/Response (dates)
    - 2.2.2.1.2 Frequency
    - 2.2.2.1.3 Warning Letters / Response
    - 2.2.2.1.4 Consent/Response
  - 2.2.2.2 FDA
    - 2.2.2.2.1 483 Inspection/Response (dates)
    - 2.2.2.2.2 Frequency
    - 2.2.2.2.3 Warning Letters/Response
    - 2.2.2.2.4 Consent/Response
- 2.2.3 Other
  - 2.2.3.1 Inspection/Response
  - 2.2.3.2 Warning Letters/Response
  - 2.2.3.3 Consent/Response

## **2.3.0**

### ***Accreditations/Certifications\****

- 2.3.1 Pharmacy Accreditations
- 2.3.2 Pharmacy Accreditations or Certifications that have been rescinded or suspended



### ***Section 3.0.0 Physical Description of Facility***

Each Pharmacy shall be of sufficient size, as determined by the Board, to allow for the safe and proper storage of Prescription Drugs and for the safe and proper Compounding and/or preparation of Prescription Drug Orders.

#### **3.1.0        *Policies and Procedures***

- 3.1.1    Confirmation of P&Ps in place and readily available

#### **3.2.0        *Size***

- 3.2.1    Pharmacy Size
  - 3.2.1.1 Size of dedicated space within the Pharmacy where prescription processing activities occur

#### **3.3.0        *Appearance***

- 3.3.1    Cleanliness

#### **3.4.0        *Records Storage***

- 3.4.1    On-site storage of required paper records
- 3.4.2    Off-site storage, if allowed of records
- 3.4.3    Ready retrieval of electronic records
- 3.4.4    Record retention length of time

#### **3.5.0        *Equipment***

- 3.5.1    Equipment to maintain and monitor temperature and humidity control of environment
- 3.5.2    Drug storage equipment (eg, refrigerators and freezers and appropriate equipment to monitor storage temperatures)
- 3.5.3    Necessary reference materials in accordance with scope of practice at pharmacy

#### **3.6.0        *Security***

- 3.6.1    Physical Security
  - 3.6.1.1 Entry detection and access
  - 3.6.1.2 Alarm system(s)
  - 3.6.1.3 Controlled substances secured
- 3.6.2    Pharmacist Responsibility
  - 3.6.2.1 Effective control against theft or diversion of Drugs and/or Devices
  - 3.6.2.2 Resignation or termination of staff for cause
  - 3.6.2.3 Proper reporting to law enforcement for diversion/termination for cause
- 3.6.3    Patient Confidentiality
  - 3.6.3.1 System security
  - 3.6.3.2 Personnel access, monitoring, and revocation
  - 3.6.3.3 Disposal
    - 3.6.3.3.1 Prescription information (PHI)
    - 3.6.3.3.2 Prescription packaging (vials, etc)
    - 3.6.3.3.3 Hazardous waste
    - 3.6.3.3.4 Drugs
- 3.6.4    Systems Backup
  - 3.6.4.1 Facility
  - 3.6.4.2 Prescription processing

- 3.6.5 Equipment
  - 3.6.5.1 General
  - 3.6.5.2 Automated
    - 3.6.5.2.1 Dispensing
    - 3.6.5.2.2 Packaging
  - 3.6.5.3 System Backup

### **3.7.0      *Compounding Area***

- 3.7.1 General Condition
  - 3.7.1.1 Cleanliness
  - 3.7.1.2 Risk level
- 3.7.2 Nonsterile
  - 3.7.2.1 Compounding area size
  - 3.7.2.2 Compounding powder hoods number
- 3.7.3 Sterile Compounding
  - 3.7.3.1 Ante Room size
  - 3.7.3.2 Clean/Buffer Room size
  - 3.7.3.3 LAFW hoods/areas
  - 3.7.3.4 Number BSC hoods
  - 3.7.3.5 Number CAI/CACI hoods
- 3.7.4 Nonsterile HD Compounding Area size
  - 3.7.4.1 Designated HD Compounding hoods number (in addition to 3.7.2.2)
- 3.7.5 Sterile HD Compounding
  - 3.7.5.1 Negative Pressure
  - 3.7.5.2 Sterile HD Room size
  - 3.7.5.3 Number of BSC hoods
  - 3.7.5.4 Number of CACI hoods

### **3.8.0      *Hours of Operation***

### **3.9.0      *Drive-through window***

## **Section 4.0.0   *Description of Dispensing and/or Distribution Information***

### **4.1.0      *Policies and Procedures***

- 4.1.1 Confirmation of P&Ps in place and readily available

### **4.2.0      *Prescription Volume Dispensed (DISPENSE means to provide a prescription product or compound pursuant to a patient-specific prescription.)***

- 4.2.1 Average Prescriptions Dispensed/day/week/month
- 4.2.2 State-specific breakdown

**4.3.0**      ***Volume Distributed (DISTRIBUTE means to provide a prescription product or compound to a prescriber or health care entity for office use or stock and is NOT patient specific – is not labeled with the patient name at the pharmacy.)***

- 4.3.1    Average Orders Distributed/day/week/month
- 4.3.2    State-specific breakdown

**Section 5.0.0**   ***Prescription Processing***

**5.1.0**      ***Policies and Procedures***

- 5.1.1    Confirmation of P&Ps in place and readily available

**5.2.0**      ***Processing at Pharmacy***

- 5.2.1    Processes in place to assure the integrity, legitimacy, and authenticity of prescription orders
- 5.2.2    Staff Training
- 5.2.3    Red Flags
- 5.2.4    Procedure to follow when a prescription is suspected of (or actually is) fraudulent
  - 5.2.4.1    Action Steps
  - 5.2.4.2    Reporting
- 5.2.5    Electronic prescription capability and Processing
  - 5.2.5.1    Non-controlled substances
  - 5.2.5.2    Controlled substances
- 5.2.6    Processes in place for ensuring that prescriptions are verified for accuracy prior to dispensing
- 5.2.7    Processes in place to ensure integrity, stability, and purity of prescription medications in transport, if applicable
- 5.2.8    Brand of prescription processing system
- 5.2.9    Veterinary prescriptions
  - 5.2.9.1    Records and labeling include species and name of the animal/owner as required by resident state law.
  - 5.2.9.2    Identification as a veterinary prescription in pharmacy records/system?
- 5.2.10    Labeling of prescriptions
- 5.2.11    Records of dispensing
  - 5.2.11.1        Written and verbal (reduced to writing) prescriptions
  - 5.2.11.2        Electronic prescriptions
  - 5.2.11.3        Dispensing records
  - 5.2.11.4        On-site or readily retrievable for required retention time

**5.3.0**      ***Shared Services /Central Fill/Central Processing***

- 5.3.1    P&Ps for shared services
- 5.3.2    Appropriate records showing identification of person performing and accountability for each separate process, and appropriate labeling
- 5.3.3    Portions of the prescription processing performed at a different location
  - 5.3.3.1    Prescriptions received by another location (including written, telephone,

- fax, electronic)
  - 5.3.3.2 Patient information (demographics and contact information) and profile information (allergies, disease states, etc) entered into the computer at another location
  - 5.3.3.3 Prescription information entered into the computer system at another location
  - 5.3.3.4 Accuracy of the prescription information entered into the computer verified at another location
  - 5.3.3.5 Prescriptions dispensed or sold pursuant to shared services agreement
  - 5.3.3.6 DUR process (including assessing and acting on DUR alerts and warnings) performed at another location (prospective review)
- 5.3.4 Ownership/Organization
  - 5.3.4.1 Common ownership
  - 5.3.4.2 Central fill
  - 5.3.4.3 Other agreement

## **5.4.0      *Drug Utilization Review***

- 5.4.1 Prospective DUR prior to the dispensing of a medication or product
  - 5.4.1.1 Drug-drug interaction (prescription and OTC)
  - 5.4.1.2 Drug-allergy interaction
  - 5.4.1.3 Therapeutic duplication
  - 5.4.1.4 Under- or over-utilization (including clinical abuse/misuse)
  - 5.4.1.5 Disease state or condition contraindication
  - 5.4.1.6 Incorrect dosage or duration of therapy
  - 5.4.1.7 Gender or age-related contraindications
  - 5.4.1.8 Additional information in the DUR process
  - 5.4.1.9 DUR overrides/bypasses documented (documented via a password/biometric override or by computer logs)
  - 5.4.1.10 Manual or electronic process
    - 5.4.1.10.1 Name of vendor and system

## **5.5.0      *Prescription Monitoring Program***

- 5.5.1 Access by pharmacists to the state PMP/PDMP program data for specific patients?
- 5.5.2 Verification of a policy regarding access and follow-up or reporting
- 5.5.3 Pharmacist utilization of the PMP data
- 5.5.4 Required reporting to PMPs by the pharmacy.

## **5.6.0      *Off-site***

- 5.6.1 Emergency kits in nursing homes, long-term care facilities, or other entities (such as hospice, EMTs, ambulances)
- 5.6.2 Maintain any automated prescription dispensing devices outside the pharmacy, such as Pyxis in a nursing home, secure mailbox device patients access after hours, etc
  - 5.6.2.1 Licenses
- 5.6.3 Do emergency kits and automated prescription dispensing devices contain any compounded sterile products?

## **Section 6.0.0 Patient Care**

### **6.1.0 Policies and Procedures**

- 6.1.1 Confirmation of P&Ps in place and readily available

### **6.2.0 Patient Records and Profile Data**

- 6.2.1 Organized and readily accessible to facilitate consultation with the prescriber, patient, or caregiver.

### **6.3.0 Patient Counseling and Communication**

- 6.3.1 Pharmacists provide counseling for all new prescriptions/changes in therapy, prescriptions picked up at the pharmacy (proactively, no “offer”)
  - 6.3.1.1 Documentation
- 6.3.2 An “offer” to counsel is made for all new prescriptions/changes in therapy, prescriptions picked up at the pharmacy
  - 6.3.2.1 Personnel making offer
  - 6.3.2.2 Documentation of offer for counseling
- 6.3.3 Patient counseling provided for delivered prescriptions
- 6.3.4 Patient counseling provided for mailed prescriptions
- 6.3.5 Patient Information
  - 6.3.5.1 PPIs provided with every fill and refill of medications for which they are required (such as hormone products, inhalers, etc)
  - 6.3.5.2 MedGuides provided with every fill and refill of medications for which they are required (such as NSAIDS, antidepressants, etc)
- 6.3.6 REMS implementation programs performed
  - 6.3.6.1 Programs (such as iPledge for isotretinoin, or Tikosyn)
- 6.3.7 Patient counseling documented
  - 6.3.7.1 Pharmacist notes in the profile of the patient during counseling
  - 6.3.7.2 Refusal of counseling documented
- 6.3.8 Patients have 24-hour access to a pharmacist
- 6.3.9 Process for drug recalls
- 6.3.10 Drug take-back program
  - 6.3.10.1 DEA registration modification for controlled substances take back
- 6.3.11 Private area for patient counseling and providing patient counseling service
- 6.3.12 Face-to-face/remote

### **6.4.0 Other Patient Services**

- 6.4.1 Collaborative Drug Therapy Management, Medication Therapy Management
- 6.4.2 Immunizations
- 6.4.3 Patient lab testing such as blood glucose tests, cholesterol tests, etc?
  - 6.4.3.1 CLIA waiver expiration date
  - 6.4.3.2 Name of lab director listed. Verify that the lab director is current (usually the PIC is the lab director named)
- 6.4.4 Training for additional services
- 6.4.5 P&Ps for other patient services

## ***Section 7.0.0 Product Receipt and Inventory***

### ***7.1.0 Policies and Procedures***

- 7.1.1 Confirmation of P&Ps in place and readily available

### ***7.2.0 Pharmacy drug ordering***

- 7.2.1 Received product from authorized trading partners

#### ***7.2.1.1 List trading partners***

- 7.2.2 Receive transaction data (transaction history, transaction information, transaction statement)
- 7.2.3 Verification-suspect and illegitimate product

### ***7.3.0 Controlled Substances (all C-I through C-V)***

- 7.3.1 Records of receipt/invoices
- 7.3.2 Procurement of C-II controlled substances
  - 7.3.2.1 DEA-222 forms to procure C-II substances
  - 7.3.2.2 CSOS utilization (electronic procurement) to order and receive C-II controlled substances
  - 7.3.2.3 Power of Attorney
- 7.3.3 Inventories
  - 7.3.3.1 Biennial (or more frequent if required by state)
  - 7.3.3.2 Other required inventories (eg, change of PIC, theft/loss)
  - 7.3.3.3 Perpetual inventory log of all C-II controlled substances (including APIs, if applicable)
    - 7.3.3.3.1 Perpetual inventory log reconciliation

### ***7.4.0 Pseudoephedrine***

- 7.4.1 Record of sales

### ***7.5.0 Other restricted products (such as exempt C-V controlled substances, paraphernalia, dextromethorphan, Plan B, etc)***

- 7.5.1 Record of sales

### ***7.6.0 Outdated, damaged, or recalled products segregated***

- 7.6.1 P&Ps on reverse distribution

### ***7.7.0 Repackage prescription medications***

- 7.7.1 Pre-pack bulk containers of prescription medications
  - 7.3.3.4 Unit-of-use quantities
  - 7.3.3.5 Blister packaging for subsequent dispensing; eg, long-term care system
  - 7.3.3.6 Patient compliance packaging
  - 7.3.3.7 Other
- 7.7.2 Repackaging record/log
- 7.7.3 Appropriate labeling of repackaged products
- 7.7.4 Repackaging and shipping interstate or intrastate
  - 7.3.3.8 Registered as a repackager with FDA, if applicable
- 7.7.5 Return to Stock P&Ps for dispensed prescriptions not picked up

## ***Section 8.0.0 Compounding***

### **8.1.0        *Nonsterile Compounding***

- 8.1.1 Policies and Procedures
  - 8.1.1.1 Confirmation of P&Ps in place and readily available
- 8.1.2 USP Chapter <795> and other referenced chapters
- 8.1.3 Veterinary Compounding

### **8.2.0        *Sterile Compounding***

- 8.2.1 Policies and Procedures
  - 8.2.1.1 Confirmation of P&Ps in place and readily available
- 8.2.2 USP Chapter <797> and other referenced chapters
- 8.2.3 Veterinary Compounding

### **8.3.0        *Nuclear Compounding***

- 8.3.1 Policies and Procedures
  - 8.3.1.1 Confirmation of P&Ps in place and readily available
- 8.3.2 USP Chapters <795> and <797> and other referenced chapters (with nuclear exemptions noted in the chapter)
- 8.3.3 US Nuclear Regulatory Commission portions of CFR Title 10

## ***Section 9.0.0 Quality Assurance/Quality Improvement Program***

### **9.1.0        *Documented formalized QA/QI program***

- 9.1.1 Oversight of the program
- 9.1.2 Formal performance program and written P&Ps
- 9.1.3 QA data readily retrievable

### **9.2.0        *QRE defined***

- 9.2.1 Recording
- 9.2.2 Reporting
  - 9.2.2.1 Internal
  - 9.2.2.2 Outside peer review committee or patient safety organization

### **9.3.0        *Errors***

- 9.3.1 External errors
  - 9.3.1.1 Documented and tracked
  - 9.3.1.2 Reached the patient
- 9.3.2 Internal errors
  - 9.3.2.1 Documented and tracked
  - 9.3.2.2 Reporting
- 9.3.3 Root cause analysis implemented

### **9.4.0        *Reporting adverse events to appropriate entities (eg, board of pharmacy, FDA MedWatch, VAERS)***

- 9.4.1 Prescription medications
- 9.4.2 Compounded products
- 9.4.3 Vaccinations

**9.5.0**      ***Incidents involving malfunctioning or defective medical equipment or devices (glucose meters, DME, injection devices, etc) documented and reported to the manufacturer or distributor***

**9.6.0**      ***Patient complaints***

9.6.1      Documented and tracked

**9.7.0**      ***Data***

9.7.1      Evaluated

9.7.2      Summary QA/QI

9.7.2.1      Report

9.7.2.2      Shared with staff

9.7.3      Process or policy changes or improvements made based upon other data collected in the QA/QI program

9.7.4      Improvements or changes evaluated for performance as a way to measure the effectiveness of the QA/QI program



## ***Pharmacy Interstate Inspection Blueprint Key***

<b>API:</b>	<b>Active pharmaceutical ingredient</b>
<b>BSC:</b>	<b>Biological safety cabinet</b>
<b>CAI:</b>	<b>Compounding Aseptic Isolator</b>
<b>CACI:</b>	<b>Compounding Aseptic Containment Isolator</b>
<b>CLIA:</b>	<b>Clinical Laboratory Improvement Amendment</b>
<b>CSOS:</b>	<b>Controlled Substances Ordering System</b>
<b>DBA:</b>	<b>Doing business as</b>
<b>DEA:</b>	<b>Drug Enforcement Administration</b>
<b>DME:</b>	<b>Durable medical equipment</b>
<b>DMEPOS:</b>	<b>Durable medical equipment, prosthetics, orthotics, and supplies</b>
<b>DUR:</b>	<b>Drug Use Review</b>
<b>EMT:</b>	<b>Emergency medical technician</b>
<b>FDA:</b>	<b>Food and Drug Administration</b>
<b>HD:</b>	<b>Hazardous drug</b>
<b>HMO:</b>	<b>Health maintenance organization</b>
<b>LAFW:</b>	<b>Laminar airflow workbench</b>
<b>NSAID:</b>	<b>Non-steroidal anti-inflammatory drug</b>
<b>OTC:</b>	<b>Over-the-counter</b>
<b>P&amp;P:</b>	<b>Policy and procedure</b>
<b>PBM:</b>	<b>Pharmacy benefits manager</b>

<b>PDMP:</b>	<b>Prescription drug monitoring program</b>
<b>PHI:</b>	<b>Protected health information</b>
<b>PIC:</b>	<b>Pharmacist-in-charge</b>
<b>PMP:</b>	<b>Prescription monitoring program</b>
<b>PPI:</b>	<b>Patient package insert</b>
<b>QA:</b>	<b>Quality assurance</b>
<b>QI:</b>	<b>Quality Improvement</b>
<b>QRE:</b>	<b>Quality-related event</b>
<b>REMS:</b>	<b>Risk evaluation mitigation strategy</b>
<b>USP:</b>	<b>United States Pharmacopeia</b>
<b>VAERS:</b>	<b>Vaccine Adverse Event Reporting System</b>

**Please contact [VPP@nabp.pharmacy](mailto:VPP@nabp.pharmacy) for additional inquiries**

## **Appendix B**

### **Sample Pharmacy Automation Policy and Procedure Outline**

- I. Access
  - A. System Entry
  - B. Access Codes
  - C. System Access Privileges
  - D. Changing Access Privileges
  - E. Termination of User
  - F. Temporary Access Codes
  - G. Password Assignment
- II. Controlled Substances
  - A. Chain of Custody
  - B. Discrepancy Resolution
- III. Data
  - A. Archiving
  - B. Stored/Uploading to Database
  - C. Backup
- IV. Definitions
- V. Dispensing/Distribution
  - A. Removal of Medications and/or Pharmaceutical Supplies
  - B. Medication Access
- VI. Downtime Procedures (see Malfunction)
- VII. Emergency Procedures
- VIII. Information Security/Confidentiality
  - A. Patient Information
  - B. Medication Information
  - C. Transaction Files
  - D. Information Update Plan
  - E. Patient Update Plan
  - F. Information Access
- IX. Inspection
- X. Installation Requirements
- XI. Maintenance
  - A. Service and Repair Protocols

- XII. Medication Administration
  - A. Medication and Patient Validation
  - B. Administration Verification
- XIII. Medication Security
  - A. Security Management and Control
  - B. Medication Loading and Storage
  - C. Medication Loading Records
  - D. Medication Containers
  - E. Cross Contamination
  - F. Lot Number Control
  - G. Inventory
  - H. Utilization Review
  - I. Research
- XIV. Malfunction
  - A. Troubleshooting
  - B. Power Failure
- XV. Quality Assurance/Quality Improvement
  - A. Documentation and Verification of Proper Loading and Refilling of Device
  - B. Proof of Delivery
  - C. Removal of Drugs for Administration, Return, or Waste
  - D. Chain of Custody of Controlled Substances (Institutions)
  - E. Recording, Resolving, and Reporting of Discrepancies
  - F. Periodic Audits to Assure Compliance with Policies and Procedures
- XVI. Reports
  - A. System Maintenance
  - B. Administrative Functions
  - C. Inventory
  - D. Error
  - E. Discrepancies
  - F. Activity
  - G. Problem
- XVII. Medication Inventory
  - A. Management
- XVIII. Staff Education and Training
- XIX. System Setup

## **Appendix C**

### **Guidelines for Disciplinary Sanctions**

#### **Improperly Obtaining or Attempting to Obtain a License**

1. Fraud or Misrepresentation in applying for or procuring a pharmaceutical license or in connection with applying for or procuring periodic reregistration of a pharmaceutical license.  
Range of action: from Fine to Revocation or denial
2. Cheating on or attempting to subvert the Pharmacist licensure examination(s).  
Range of action: Revocation or denial

#### **Misdemeanors/Felonies**

3. The commission or conviction of a gross misdemeanor or a felony, whether or not related to the Practice of Pharmacy, or the entry of a guilty or nolo contendere plea to a gross misdemeanor or a felony charge.  
Range of action: from Probation to Revocation

#### **Deception/Fraud/Misrepresentation**

4. Conduct likely to deceive, defraud, or harm the public.  
Range of action: from Censure to Revocation
5. Making a false or misleading statement regarding one's skill or the efficacy or value of the medicine, treatment, or remedy Dispensed in the treatment of any disease or other condition of the body or mind.  
Range of action: from Probation to Revocation
6. The use of any false, fraudulent, or deceptive statement in any document connected with the Practice of Pharmacy.  
Range of action: from Warning to Revocation
7. Practicing Pharmacy under a false or assumed name.  
Range of action: from Probation to Revocation

#### **Patient Confidentiality/Records**

8. Improper management of Pharmacy patient records, including illegal use or disclosure of Protected Health Information.  
Range of action: from Warning to Suspension

#### **Negligence/Incompetence/Disability/Malpractice**

9. Negligence in the Practice of Pharmacy as determined by the Board.  
Range of action: from Warning to Revocation
10. Being found mentally incompetent or insane by any court of competent jurisdiction.  
Range of action: from Suspension to Revocation
11. Being physically or mentally unable to engage safely in the Practice of Pharmacy.  
Range of action: from Probation to Revocation
12. Demonstration of incapacity or incompetence to practice Pharmacy.  
Range of action: from Probation to Revocation
13. Any adverse judgment, award, or settlement against the licensee resulting from a professional malpractice claim related to conduct that would constitute grounds for

action as defined in this section.  
Range of action: from Censure to Revocation

### **Sexual Misconduct**

14. Commission of any act of sexual abuse, misconduct, or exploitation related to a licensee's Practice of Pharmacy.  
Range of action: from Probation to Revocation

### **Drug- and Alcohol-Related Offenses**

15. Being dependent on or habituated to a Drug or intoxicant.  
Range of action: from Probation to Revocation
16. Dispensing, prescribing, selling, Administering, Distributing, ordering, or giving any Drug legally classified as a controlled substance or recognized as an addictive or dangerous Drug for any purposes other than medically accepted as therapeutic.  
Range of action: from Probation to Revocation
17. Except as otherwise permitted by law, Dispensing, prescribing, selling, Administering, Distributing, ordering, or giving to an habitué, addict, or any Person previously Drug dependent any Drug legally classified as a controlled substance or recognized as an addictive or dangerous Drug.  
Range of action: from Probation to Revocation
18. Violating any State or Federal law or regulation relating to controlled substances.  
Range of action: from Warning to Revocation

### **Misuse of License**

19. Aiding or abetting the Practice of Pharmacy by an unlicensed, incompetent, or impaired Person.  
Range of action: from Reprimand to Revocation
20. Allowing another Person or organization to use one's license to practice Pharmacy.  
Range of action: from Reprimand to Revocation

### **Disciplinary Action by Other Jurisdictions**

21. Disciplinary action of another state or jurisdiction against a license or other authorization to practice Pharmacy based upon conduct by the licensee similar to conduct that would constitute grounds for action as defined in this Section.  
Range of action: same as for similar offense in this State

### **Failure to Report to and/or Cooperate with Board**

22. Failure to report to the Board any adverse action taken by another licensing jurisdiction (United States or foreign), government agency, law enforcement agency, or court for conduct that would constitute grounds for action as defined in this Section.  
Range of action: from Censure to Revocation
23. Failure to report to the Board one's surrender of a license or authorization to practice Pharmacy in another state or jurisdiction while under disciplinary investigation by any of those authorities or bodies for conduct that would constitute grounds for action as defined in this section.  
Range of action: from Censure to Revocation

- 24. Failure to report to the Board any adverse judgment, settlement, or award arising from a malpractice claim related to conduct that would constitute grounds for action as defined in this section.  
Range of action: from Censure to Suspension
- 25. Failure to cooperate with a lawful investigation conducted by the Board.  
Range of action: from Censure to Revocation
- 26. Failure to furnish to the Board, its investigators, or representatives any information legally requested by the Board.  
Range of action: from Censure to Revocation

**Other Violations**

- 27. Violation of any provision(s) of the Pharmacy Practice Act, any rules and regulations of the Board, or any action, stipulation, or Agreement of the Board.  
Range of action: corresponds to related actions above

## Appendix D

### Community Pharmacy Quality-Related Event (QRE) Data Collection Form

QRE Prescription Data			
Attach copy of: <input type="checkbox"/> prescription <input type="checkbox"/> label <input type="checkbox"/> photo of vial (mark all available)			
Original Rx date:		Refill date: Date/time reported:	
Drug Prescribed (name, strength, and dosage form): Directions: Medication indication:			
Prescription was received by the pharmacy via: <input type="checkbox"/> telephone, by whom: <input type="checkbox"/> written <input type="checkbox"/> computer <input type="checkbox"/> fax			

QRE Data	
QRE Type: (select all that apply) A. Prescription processing error: <input type="checkbox"/> Incorrect drug (1) <input type="checkbox"/> Incorrect strength (2) <input type="checkbox"/> Incorrect dosage form (3) <input type="checkbox"/> Incorrect patient (4) <input type="checkbox"/> Inaccurate or incorrect packaging, labeling, or directions (5) <input type="checkbox"/> Other (6):	B. A failure to identify and manage <input type="checkbox"/> Over/under-utilization (1) <input type="checkbox"/> Therapeutic duplication (2) <input type="checkbox"/> Drug-disease contraindications (3) <input type="checkbox"/> Drug-drug interactions (4) <input type="checkbox"/> Incorrect duration of treatment (5) <input type="checkbox"/> Incorrect dosage (6) <input type="checkbox"/> Drug-allergy interaction (7) <input type="checkbox"/> Clinical abuse/misuse (8)



QRE Contributing Factors	
Day of the week and time of QRE:	
No. of new prescriptions:	No. of refill prescriptions:
RPh to tech ratio:	
RPh staff status: <input type="checkbox"/> regular staff	<input type="checkbox"/> part-time/substitute staff
length of employment:	
No. of hours RPh on duty:	Average No. of prescriptions filled per hour:
No. of other RPhs on duty:	No. of support staff on duty:
Automation <input type="checkbox"/> yes <input type="checkbox"/> no	Type:
<input type="checkbox"/> Computer software (lighting/noise/distractions/workspace) <input type="checkbox"/> Environmental <input type="checkbox"/> Equipment failure <input type="checkbox"/> Failure to supervise <input type="checkbox"/> Legibility <input type="checkbox"/> Increased Rx volume as compared to normal <input type="checkbox"/> Shift change <input type="checkbox"/> Policies and/or procedure not followed <input type="checkbox"/> Staff shortage <input type="checkbox"/> Sound alike/look alike medication <input type="checkbox"/> Other, explain	
Describe factors checked above and/or other preliminary root contributors:	
Counseling was offered: <input type="checkbox"/> yes <input type="checkbox"/> no	Counseling was given: <input type="checkbox"/> yes <input type="checkbox"/> no
Documentation of offer: <input type="checkbox"/> yes <input type="checkbox"/> no	Documentation of counseling: <input type="checkbox"/> yes <input type="checkbox"/> no

Pharmacist Information
Name of verifying pharmacist:
Name(s) of other person(s) and title(s) involved in processing the prescription:
Describe remedial action taken:

**If patient received medication, complete Patient and Prescriber Information sections.**

Patient Information	
Patient's name:	Prescription was dispensed to:
Address:	Telephone No.:
Patient DOB: Sex: M or F If minor, name of parent(s)/guardian(s):	
Who discovered the error/relationship to patient?:	
Did patient ingest medication? <input type="checkbox"/> yes <input type="checkbox"/> no If yes, how many doses?:	
<input type="checkbox"/> Not harmed <input type="checkbox"/> Received treatment and or increased monitoring <input type="checkbox"/> Seriously harmed, explain <input type="checkbox"/> Did not survive, explain	

Prescriber Information
Was the prescriber informed: <input type="checkbox"/> yes <input type="checkbox"/> no If yes, provide date:
Prescriber's name: Telephone No.:
Prescriber's instructions/comments:

Report Affirmation
Additional comments:
Name and title of preparer of this report:
Signature: Date:

## Community Pharmacy Continuous Quality Improvement Program Inspection Form

General Information	
Pharmacy name:	License No.:
Address:	Phone No.:
Pharmacist-in-charge (PIC):	PIC License No.:
Date/time:	Date of previous inspection: Attach copy of previous inspection
<b>Purpose of inspection</b> <input type="checkbox"/> Complaint <input type="checkbox"/> Routine <input type="checkbox"/> Follow-up <input type="checkbox"/> New pharmacy <input type="checkbox"/> Change in owner <input type="checkbox"/> Other Comment:	

Pharmacy Staff			
(Include pharmacist, intern, certified pharmacy technician, certified pharmacy technician candidate, and cashier)			
Name	Title	License No.	Present

<b>P=Present                      A=Absent                      N/A= Not applicable</b>				
<b>CQI Program</b>		<b>P</b>	<b>A</b>	<b>N/A</b>
	Policy and procedures in place			
	Periodic CQI meetings held			
	Quality-Related Events (QRE) recorded			
	Sentinel Events			
	Workload compiled			
	Staffing needs analyzed/addressed			
	Outcome-based certified pharmacy technician training conducted			
	Technology utilized in current/updated			
	Pharmacist Care Services initiatives in place			
	Consumer survey policy in place			
	Professional performance evaluation policy in place			
<b>Comments:</b>  				
<b>Recommendations:</b>  				
<b>Report Affirmation</b>				
<b>Additional comments:</b>  				
Pharmacist signature:		Date:		
Surveyor signature:		Date:		

## Community Pharmacy Quality Self-Audit

Each pharmacy shall conduct a quality self-audit at least quarterly and upon change of pharmacist-in-charge. The goals of the quality self-audit are to monitor changes in the number of quality-related events (QRE) over time, to evaluate compliance with CQI procedures, and to develop a plan for improved adherence with the CQI Program.

General Information	
Date:	Quarterly <input type="checkbox"/> Change of pharmacist-in-charge <input type="checkbox"/>
Pharmacy name:	Address:
Telephone:	License No.:
Pharmacist-in-charge:	Date of previous self-audit:

Pharmacy Staff			
(Include pharmacist, intern, certified pharmacy technician and certified pharmacy technician candidate, cashier)			
Name	Title	License No.	Start Date

Staffing/Workload Date	
Staffing	Yes/No/Answer
Number of pharmacist hours allocated per week	
Number of certified pharmacy technician hours allocated per week	
Number of certified pharmacy technician candidate hours allocated per week	
Number of other pharmacy support staff hours allocated per week	
Number of certified pharmacy technicians	
Number of certified pharmacy technician candidates	
Outcome-based certified pharmacy technician training program (If yes, check all applicable)	
<input type="checkbox"/> Cash register <input type="checkbox"/> Prescription intake <input type="checkbox"/> Prescription filling	
<input type="checkbox"/> Inventory <input type="checkbox"/> Returning stock bottles to shelf <input type="checkbox"/> Clean room	
<input type="checkbox"/> Computer data entry <input type="checkbox"/> Pharmaceutical calculations <input type="checkbox"/> Knowledge of practice settings	
<input type="checkbox"/> Identifying drugs, doses, routes of Administration, dosage forms, etc	
<input type="checkbox"/> Pharmaceutical and medical terminology <input type="checkbox"/> Other	
Workload	Yes/No/Answer
Number of hours pharmacy department is open during the week	
Average number of prescriptions filled per week	
Usual ratio of pharmacist to technicians	
Policy is in place that requires increased staffing if workload increases	
Automation	Yes/No/Answer
Type	

CQI Program Data	
General	Yes/No/Answer
Pharmacy has a CQI policy and procedure manual	

Employees must verify review of policy and procedure manual	
<b>Periodic CQI Meetings</b>	<b>Yes/No/Answer</b>
Pharmacy holds CQI meetings (if yes, indicate frequency)	
Average length of CQI meetings in minutes	
Staff attending CQI meetings <input type="checkbox"/> Pharmacists <input type="checkbox"/> Technicians <input type="checkbox"/> Manager <input type="checkbox"/> Pharmacy supervisor <input type="checkbox"/> Owner <input type="checkbox"/> Other	

<b>Program Documentation Methods</b>	<b>Yes/No/Answer</b>
QRE forms utilized	
Method used to document interaction in relation to CQI program <input type="checkbox"/> Computer database <input type="checkbox"/> On prescription <input type="checkbox"/> Custom-made form <input type="checkbox"/> Standard form <input type="checkbox"/> Other	
Method used to verify drug product with prescription label <input type="checkbox"/> Bar code <input type="checkbox"/> NDC code <input type="checkbox"/> Name of product <input type="checkbox"/> Other	
First time refills are checked against hardcopy	
<b>Consumer Surveys</b>	<b>Yes/No/Answer</b>
Consumer survey policy in place, if yes, indicate frequency	
Other technique in place to evaluate performance, if yes, describe	
Method of conducting consumer survey <input type="checkbox"/> Distributed at time of dispensing <input type="checkbox"/> Mail <input type="checkbox"/> Telephone <input type="checkbox"/> Other	
Consumer survey feedback utilized to improve delivery of pharmacy services	
<b>Outcome-Based Professional Performance Evaluation</b>	
Frequency <input type="checkbox"/> Annually <input type="checkbox"/> Biannually <input type="checkbox"/> Quarterly <input type="checkbox"/> Other	

<p>Staff required to have outcome-based professional performance evaluations</p> <p><input type="checkbox"/> All employees   <input type="checkbox"/> Full-time pharmacists   <input type="checkbox"/> Part-time pharmacists   <input type="checkbox"/> Other</p>
<p>Self-audit includes:</p>
<p><input type="checkbox"/> Number of overridden drug-drug interaction warnings</p>
<p><input type="checkbox"/> Number of patients that received duplicative drug therapy</p>
<p><input type="checkbox"/> Number of patients that received extensive counseling</p>
<p><input type="checkbox"/> Number of QREs tracked over time. Indicate time period</p>



**QRE Incidents**

Utilizing QRE Data Collection Sheets, compile the data below.

Date								
QRE type (eg, A(1) = incorrect drug dispensed)								
Drug name and strength								
Rx received via:								
New or refill								
Day of week/time								
RPh to tech ratio								
RPh staff status								
No. of hrs RPh on duty								
No. of other pharmacists on duty								
No. of other support staff								
Average No. of prescriptions/hour								

Responsible pharmacist's name								
Patient received medication								
Prescriber notified								
Counseling offered								
Counseling accepted								

## **Appendix E**

### **Model Prescription Monitoring Program Act**

#### **Section 1. Short Title.**

This Act shall be known and may be cited as the Model Prescription Monitoring Program Act.

#### **Section 2. Legislative Findings.**

(Insert State-appropriate mission/purposes.)

#### **Section 3. Purpose.**

(Insert State-appropriate mission/purposes.)

#### **Section 4. Definitions.**

- (a) “Dispenser” means a person authorized in this State to distribute to the ultimate user a substance monitored by the Prescription Monitoring Program, but does not include:
  - (1) a licensed hospital or institutional facility Pharmacy that distributes such substances for the purposes of inpatient care;
  - (2) a licensed nurse or medication aide who administers such a substance at the direction of a licensed physician;
- (b) “Drug of Concern” means any prescription or over-the-counter medication that demonstrates a potential for abuse, particularly those identified by Boards of Pharmacy, law enforcement, and addiction treatment professionals.
- (c) “Electronic Health Information Systems” means an electronic data intermediary, gateway, or hub that facilitates secure delivery of electronic health information to Practitioners or Dispensers, including:
  - (1) health information exchanges;
  - (2) health information networks;
  - (3) pharmacy software systems;
  - (4) electronic medical (health) record software applications; or
  - (5) electronic prescribing software applications.
- (d) “Interoperability” means the sharing of Prescription Monitoring Program Information with another PMP, or the integration of Prescription Monitoring Program Information into the Electronic Health Information Systems.
- (e) “Prescription Monitoring Program Information” means information submitted to and maintained by the Prescription Monitoring Program.<sup>185</sup>
- (f) “Prescription Monitoring Program (PMP)” means a program established under Section 5 of this Act.

---

<sup>185</sup> This reporting exception also applies to situations where a patient, who has been dispensed controlled substance medications during a stay in an institutional facility, is allowed to retain any remaining medication upon discharge.

## **Section 5. Establishment of a Prescription Monitoring Program.**

- (a) The Board of Pharmacy shall establish and maintain an electronic system for monitoring all controlled substances in Schedules II through V, all State-specified controlled substances in Schedules II through V, and State-specified Drugs of Concern dispensed to patients in this State.
- (b) The Board of Pharmacy may contract with a vendor to establish and maintain the electronic monitoring system pursuant to guidelines, which the Board of Pharmacy shall promulgate.
- (c) The Board of Pharmacy shall promulgate rules or establish policy to include the following:
  - (1) using the PMP to improve patient care and to facilitate the goal of reducing misuse, abuse, overdose, addiction to and diversion of controlled substances and Drugs of Concern;
  - (2) implementing security and safeguards necessary to ensure information is released only to authorized individuals;
  - (3) developing criteria for referring Prescription Monitoring Program information to a law enforcement agency
  - (4) developing criteria for referring Prescription Monitoring Program Information to a licensing boards, or other state or federal agency charged with the regulation of prescribing, dispensing, or administering a controlled substance or Drug of Concern;
  - (5) designing and implementing training, education, and/or instruction in the appropriate access to and use of the PMP;
  - (6) adopting the most recent version of the American Society for Automation in Pharmacy (ASAP) technical standards for electronic reporting of Prescription Monitoring Program Information; and
  - (7) incorporating technological improvement to facilitate the interoperability of the PMP with other state PMPs and Electronic Health Information Systems and to facilitate Prescribers' and Dispensers' access to and use of the PMP.

## **Section 6. Reporting of Prescription Monitoring Program Information.**

- (a) Each Dispenser shall submit to the Board of Pharmacy, by electronic means, or other format specified in a waiver granted by the Board of Pharmacy, within 24 hours, information specified by the Board of Pharmacy, including:
  - (1) identification Number of Dispenser;
  - (2) identification number of the Prescriber;<sup>186</sup>
  - (3) patient name, address, and telephone number;<sup>187</sup>
  - (4) patient gender;
  - (5) patient date of birth;
  - (6) identification of the drug by a national drug code number;
  - (7) quantity dispensed;
  - (8) number of days supplied;
  - (9) number of refills ordered;
  - (10) whether drug was dispensed as a refill or as a new prescription;

---

<sup>186</sup> It is recommended that Boards of Pharmacy consider using practitioner's NPI number for identification purposes when applicable. Consider using state license numbers for veterinarians.

<sup>187</sup> For veterinary prescriptions, use the pet owner's name, address, telephone number, gender, and date of birth.

- (11) date prescription was dispensed;<sup>188</sup>
  - (12) if a refill, date of the original dispensing;
  - (13) prescription number;
  - (14) date the prescription was issued by the Prescriber;
  - (15) method of payment for the prescription; and
  - (16) such other information as may be required by State law.
- (b) Each Dispenser shall ensure that information reported to the PMP is correct and shall submit corrections when necessary.
  - (c) Each Dispenser shall reverse information for any prescription that was not dispensed.

## **Section 7. Access to Prescription Monitoring Program Information/Confidentiality.**

- (a) Except as indicated in paragraphs (b), (c), and (d) of this Section 7, Prescription Monitoring Program Information submitted to the Board of Pharmacy shall be considered Protected Health Information and not subject to public or open records laws.
- (b) The Board of Pharmacy shall review the Prescription Monitoring Program Information. If there is reasonable cause to believe a violation of law (or breach of professional or occupational standards) may have occurred, the Board shall notify the appropriate law enforcement, or professional or occupational licensing, certification, or regulatory agency or entity, and provide Prescription Monitoring Program Information required for an investigation.<sup>189</sup>
- (c) The Board of Pharmacy may provide Prescription Monitoring Program Information for public research, policy or education purposes, to the extent all information has been De-identified.
- (d) The following persons may access the Prescription Monitoring Program Information in the same or similar manner, and for the same or similar purposes, as those persons are authorized to access similar Protected Health Information under federal and State law and regulation:
  - (1) Practitioners (or agents thereof) or Dispensers (or agents thereof) who certify, under the procedures determined by the State, that the requested information is for the purpose of providing medical or pharmaceutical treatment or evaluating the need for such treatment to a bona fide current patient, or verifying PMP information for prescriptions issued by practitioners;
  - (2) Boards of Pharmacy or vendors/contractors for the purpose of establishing and maintaining the Prescription Monitoring Program;
  - (3) other state licensing, certification, or regulatory agencies that license, certify, or regulate health care professionals authorized to prescribe, administer, and dispense controlled substances, which certify, under the procedures determined by the State, that the requested information is related to an individual investigation or proceeding involving the unlawful diversion or misuse of a

---

<sup>188</sup> It is recommended that the date prescription was dispensed be clarified to mean the date of delivery to the patient.

<sup>189</sup> This section is intended to allow boards of pharmacy to evaluate Prescription Monitoring Program information and determine appropriate information to provide to law enforcement entities. It is not intended to allow law enforcement officials open access to all data.

- reportable substance, and such information will further the purpose of the investigation or assist in the proceeding;
- (4) local, State, or Federal law enforcement, narcotics control, licensure, disciplinary, or program authorities, which certify, under the procedures determined by the State, that the requested information is related to an individual investigation or proceeding involving the unlawful diversion or misuse of a reportable substance, and such information will further the purpose of the investigation or assist in the proceeding;
  - (5) entities that serve as sources of data impacting the identification and reporting of prescription drug injuries and deaths, such as, but not limited to, coroners' offices, to help address the prescription drug epidemic and improve patient care;
  - (6) other appropriate entities;<sup>190</sup> and
  - (7) Patients who certify, under the procedures determined by the State, that the requested information is for the purpose of obtaining and reviewing their own records.
- (e) The Board of Pharmacy shall be immune from civil liability arising from inaccuracy of any of the information submitted to the Board of Pharmacy pursuant to this Act.

#### **Section 8. Interoperability.**

- (a) The Board of Pharmacy shall execute a memorandum of understanding to participate in a single national hub capable of facilitating interoperability among Prescription Monitoring Programs and between Prescription Monitoring Programs and Electronic Health Information Systems.
- (b) The Board of Pharmacy shall ensure that access to Prescription Monitoring Program Information by other state Prescription Monitoring Programs is limited to persons described in Section 7(d).
- (c) The Board of Pharmacy shall establish the technological connectivity and infrastructure to facilitate the secure delivery of Prescription Monitoring Program Information to authorized users of Prescription Monitoring Programs through other states' Prescription Monitoring Programs or Electronic Health Information Systems.
- (d) Any such gateway, hub, or any Electronic Health Information System that facilitates the integration of Prescription Monitoring Program Information into a patient's medical record shall:
  - (1) verify the identity of the individual requesting the Information;
  - (2) verify the credential of the individual requesting the Information;
  - (3) provide the Board of Pharmacy with an audit trail for each request; and
  - (4) maintain the security and confidentiality of such information.

#### **Section 9. Unlawful Acts and Penalties.**

- (a) A Dispenser who knowingly fails to submit Prescription Monitoring Program Information to the Board of Pharmacy as required by this Act shall be subject to (insert appropriate administrative, civil, or criminal penalty).

---

<sup>190</sup> It is recommended that other appropriate entities include drug courts, district attorneys' offices, addiction treatment professionals, or other similar entities, and only for the purpose of ensuring appropriate patient treatment, as opposed to efforts to search for information without knowledge of whether such information exists.

- (b) A person who knowingly accesses or uses Prescription Monitoring Program Information without authorization in violation of this Act shall be subject to (insert appropriate administrative, civil, or criminal penalty).
- (c) A person authorized to have Prescription Monitoring Program Information pursuant to this Act who knowingly discloses such information in violation of this Act shall be subject to (insert appropriate administrative, civil, or criminal penalty).
- (d) A person authorized to have Prescription Monitoring Program Information pursuant to this Act who uses such information in a manner or for a purpose in violation of this Act shall be subject to (insert appropriate administrative, civil, or criminal penalty).

#### **Section 10. Evaluation, Data Analysis, and Reporting.**

- (a) The Board of Pharmacy shall design and implement an evaluation component to identify cost benefits of the Prescription Monitoring Program, and other information relevant to policy, research, and education involving substances monitored by the PMP.
- (b) The Board of Pharmacy shall report to the (insert appropriate State decision makers, eg, legislature) on a periodic basis, no less than annually, about the cost-benefits and other information noted in paragraph (a).

#### **Section 11. Rules and Regulations.**

The Board of Pharmacy shall promulgate rules and regulations necessary to implement the provisions of this Act.

#### **Section 12. Severability.**

If any provision of this Act or application thereof to any person or circumstance is held invalid, the invalidity does not affect other provisions or applications of the Act which can be given effect without the invalid provisions or applications, and to this end the provisions of this Act are severable.

#### **Section 13. Effective Date.**

This Act shall be effective on (insert specific date or reference to normal State method of determination of the effective date).

## **Appendix F**

### **Model Rules for the Practice of Telepharmacy**

- (a) General Requirements
  - (1) The Pharmacy shall:
    - (i) obtain a resident or nonresident permit issued by the Board prior to engaging in the Practice of Telepharmacy;
    - (ii) comply with appropriate federal and state controlled substance laws and rules for each Pharmacy if controlled substances are maintained;
    - (iii) maintain additional policies and procedures specific to Telepharmacy.
- (b) Remote Dispensing Site Requirements
  - (1) Shall submit an application to the Board.
  - (2) The Pharmacist-in-Charge of supervising pharmacy shall be responsible for all operations.
  - (3) Shall have a written contract or agreement that outlines the services provided and the responsibilities of each party in complying with federal and state pharmacy laws and rules.
  - (4) The Pharmacist-in-Charge shall oversee monthly inspections, maintenance and reconciliation of all controlled substances, including maintaining a perpetual inventory for all Schedule II controlled substances.
  - (5) A Pharmacist must be designated to be available within ( \_ ) hours, in case of emergency.
  - (6) Unless staffed by a Pharmacist, a Remote Dispensing Site must be staffed by at least one (1) Certified Pharmacy Technician. All Certified Pharmacy Technicians and Certified Pharmacy Technician Candidates shall be under the supervision of a Pharmacist at the supervising Pharmacy at all times that the remote site is operational. The Pharmacist shall supervise Telepharmacy operations electronically from the supervising Pharmacy.
  - (7) The Remote Dispensing Site and the supervising Pharmacy must utilize a common electronic record-keeping system that must be capable of the following:
    - (i) Electronic records must be available to, and accessible from, both the supervising pharmacy and the Remote Dispensing Site at all times of operations; and
    - (ii) Prescriptions dispensed at the Remote Dispensing Site must be distinguishable from those dispensed from the supervising pharmacy.
  - (8) Controlled substance records must be maintained at the registered location unless specific approval is granted for central storage as permitted by, and in compliance with, state and federal law.
  - (9) A supervising Pharmacy of a Remote Dispensing Site must maintain a video and audio communication system that provides for effective communication between the supervising Pharmacy and the Remote Dispensing Site personnel and patients or caregivers. The system must provide an adequate number of views of the entire site, facilitate adequate pharmacist supervision and allow the appropriate exchanges of visual, verbal, and written communications for patient counseling and other matters involved in the lawful transaction or



Delivery of Drugs. The Remote Dispensing Site must retain a recording of facility surveillance, excluding patient communications, for a minimum of ( ) days.

- (i) Adequate supervision by the pharmacist in this setting is maintaining uninterrupted visual supervision and auditory communication with the site and full supervisory control of the automated system, if applicable, and must not be delegated to another person or entity.
  - (ii) Each component of the communication system must be in good working order. Unless a pharmacist is present onsite, the Remote Dispensing Site must be, or remain, closed to the public if any component of the communication system is malfunctioning, until system corrections or repairs are completed.
  - (iii) The video and audio communication system used to counsel and interact with each patient or patient's caregiver must be secure and compliant with state and federal confidentiality requirements.
- (10) Unless a Pharmacist is present, a Remote Dispensing Site must not be open or its employees allowed access to it during times the supervising Pharmacy is closed. The security system must allow for tracking of entries into the Remote Dispensing Site, and the Pharmacist-in-Charge must periodically review the provision of access and record of entries.
- (11) If Drugs are maintained or dispensed from the Remote Dispensing Site, Drug transfers to the Remote Dispensing Site must comply with applicable state and federal requirements.
- (12) A Remote Dispensing Site must display a sign, easily visible to the public, that informs patients:
  - (i) this is a remote site
  - (ii) location of supervising Pharmacy; and
  - (iii) that a Pharmacist will counsel the patient using audio and video communication systems each time a new medication is delivered, and on a refill, if necessary, at a Remote Dispensing Site.
- (13) The Remote Dispensing Site must use Telepharmacy Technology that confirms that the Drug selected to fill the prescription is the same as indicated on the prescription label and Prescription Drug Order.

## A

Accreditation Council for Pharmacy Education, 16, 43, 48, 49, 51, 65, 66, 67  
active ingredients, 9, 20, 104, 107, 123  
administer, 9, 14, 21, 24, 31, 39, 85, 86, 102, 104, 106, 158, 163, 165  
adulterate, 9, 10, 17, 28, 45, 70, 76, 105, 113, 118, 124  
affiliated entity, 10  
American Society of Health-System Pharmacists, 51, 73  
authenticate, 14, 119  
authentication of product history, 26, 101  
authorized distributor of record, 121  
automated pharmacy system, 10, 24, 68, 69, 73, 74, 88, 90, 91, 92, 93

## B

beyond-use date, 10, 94, 95  
bioburden, 10, 14  
biological product, 10, 24, 26, 29, 32  
biosimilar product, 10, 18, 26  
board of pharmacy, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 21, 22, 23, 24, 25, 26, 27, 28, 31, 32, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, 50, 51, 52, 53, 54, 55, 56, 57, 58, 59, 60, 61, 62, 63, 65, 66, 67, 69, 71, 72, 73, 74, 75, 76, 80, 81, 83, 84, 86, 87, 88, 89, 91, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 104, 105, 107, 108, 109, 110, 111, 112, 113, 114, 115, 116, 117, 118, 119, 120, 122, 123, 124, 125, 126, 127, 128, 130, 131, 132, 133, 137, 149, 150, 151, 163, 164, 165, 166, 167, 168

## C

cease and desist, 10, 43, 48, 53, 56, 57  
censure, 10, 43, 53, 149, 150, 151  
chart order, 11, 69, 70  
co-licensee, 12, 21  
component, 12, 14, 101  
compounding, 10, 12, 18, 19, 21, 27, 32, 43, 58, 59, 60, 61, 71, 104, 105, 106, 107, 134, 135, 137, 138, 143  
consumer survey, 12, 88, 156, 159  
continuous quality improvement program, 12, 26, 86, 87, 88, 90, 155, 156, 157, 158, 159  
contraband drug, 12, 117, 119, 120  
counterfeit drug, 13, 17, 28  
critical areas, 13  
critical surfaces, 13

## D

database, centralized performance, 11, 23, 44  
de-identified health information, 13  
designated representative, 14, 108, 111, 112, 113, 121, 124, 127

device, 8, 9, 10, 11, 12, 13, 14, 18, 19, 20, 21, 22, 29, 30, 31, 32, 33, 39, 40, 41, 43, 44, 45, 58, 59, 60, 61, 62, 63, 69, 70, 71, 72, 73, 82, 96, 97, 108, 110, 111, 112, 113, 114, 116, 117, 118, 119, 123, 137  
digital signature, 14, 15, 76, 84  
disaster areas, 13, 28, 97, 98, 100  
dispense, 9, 11, 14, 15, 16, 18, 21, 22, 24, 25, 26, 29, 32, 40, 43, 47, 58, 65, 70, 71, 73, 76, 77, 79, 80, 81, 82, 83, 84, 89, 90, 91, 92, 94, 96, 97, 98, 99, 100, 101, 102, 104, 106, 116, 122, 138, 139, 149, 163, 164, 165, 166  
dispensing, 8, 9, 10, 11, 14, 15, 19, 20, 21, 22, 24, 25, 26, 27, 28, 31, 43, 47, 58, 63, 66, 69, 70, 71, 76, 77, 78, 79, 80, 81, 82, 83, 84, 89, 92, 93, 94, 97, 98, 99, 101, 102, 104, 105, 106, 123, 147, 150, 159, 165  
distribute, 8, 9, 10, 12, 13, 14, 17, 18, 21, 22, 26, 27, 28, 29, 30, 31, 32, 33, 41, 43, 55, 58, 59, 60, 62, 63, 69, 71, 73, 82, 99, 102, 104, 106, 107, 108, 109, 110, 111, 112, 113, 116, 117, 118, 119, 122, 123, 124, 125, 126, 127, 128, 129, 130, 132, 133, 138, 139, 147, 150, 159  
diversion activity, 14, 115, 116, 118  
drug, 8, 9, 10, 11, 12, 13, 14, 15, 16, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 29, 31, 32, 33, 39, 40, 41, 43, 44, 45, 46, 47, 54, 55, 58, 59, 60, 61, 62, 63, 66, 68, 69, 70, 71, 72, 73, 74, 75, 76, 77, 78, 79, 80, 81, 82, 83, 84, 85, 86, 87, 89, 92, 93, 94, 95, 96, 97, 98, 99, 100, 101, 102, 104, 105, 106, 107, 108, 109, 110, 111, 112, 113, 114, 115, 116, 117, 118, 119, 120, 122, 123, 124, 137, 148, 150, 152, 158, 159, 160, 161, 164, 169  
Drug Distributor Accreditation, 107, 110, 126  
Drug Enforcement Administration, 27, 75, 76, 77, 83, 87, 97, 98, 100, 113, 114, 115, 116, 117, 118  
drug of concern, 14, 15, 72, 73, 114, 115, 163, 164  
drug utilization review, 11, 15, 23, 25, 27, 85, 89, 98, 140

## E

electronic signature, 14, 15, 76  
electronic transmission, 34, 76, 77, 78

## F

Federal Food, Drug, and Cosmetic Act, 10, 16, 21, 121, 122, 124, 125, 130  
Food and Drug Administration, 10, 12, 14, 16, 19, 21, 26, 30, 32, 33, 59, 73, 79, 101, 104, 106, 107, 108, 110, 116, 117, 118, 122, 123, 124, 125, 130, 131, 136, 142, 143

## G

global exchange pharmacy student, 16, 67

## H

Health Insurance Portability and Accountability Act, 13, 16, 17, 25

## I

individually identifiable health information, 17, 25, 34  
institutional pharmacy, 11, 18, 68, 69, 70  
interchangeable product, 18, 80

## L

labeling, 12, 13, 18, 19, 25, 26, 27, 45, 71, 79, 80, 81,  
94, 95, 101, 102, 104, 105, 106, 113, 114, 118,  
123, 128, 132  
long-term care facility, 17, 18, 19, 77

## M

medication therapy management, 19, 20, 22, 86  
medication-assisted treatment, 19, 87

## N

NABP, 1, 2, 3, 8, 9, 20, 21, 32, 39, 42, 43, 47, 48, 49,  
59, 65, 72, 99, 104, 110, 114, 115, 117, 126  
NABP Clearinghouse, 49, 99  
NABP Information Sharing Network, 21, 104  
nonprescription drug, 18, 21  
normal distribution channel, 21  
nuclear pharmacy, 21, 25, 58, 101, 102, 103

## P

patient counseling, 11, 19, 21, 71, 80, 85, 86, 87, 92,  
96, 98  
patient intervention program, 16, 21, 86  
pedigree, 12, 119  
pharmacist care, 9, 16, 18, 22, 23, 24, 39, 40, 43, 44,  
47, 58, 69, 71, 72, 79, 85, 88, 91, 92, 95, 96, 103,  
104, 156  
pharmacist-in-charge, 11, 22, 58, 59, 61, 68, 69, 70,  
72, 73, 74, 75, 82, 88, 90, 92, 105, 155, 157, 168,  
169  
pharmacy benefits manager, 13, 16, 17, 19, 23, 58, 60,  
61, 62  
pharmacy intern, 23, 24, 43, 49, 50, 53, 54, 56, 57, 65,  
66, 67, 74, 76, 77, 78, 89, 90, 99, 155, 157  
pharmacy practice, collaborative, 12, 22, 24, 86, 87;  
agreement, 12, 22, 86, 87  
pharmacy technician, 8, 11, 26, 43, 48, 50, 51, 52, 53,  
54, 56, 57, 73, 74, 75, 76, 77, 78, 79, 82, 86, 89,  
90, 95, 99, 105, 155, 156, 157, 158, 159, 161, 168  
pharmacy warehouse, chain, 11, 21  
pharmacy, closed, 12  
pharmacy, coordinating, 88, 89, 90, 91, 92, 168  
practice of pharmacy, 8, 9, 11, 12, 22, 23, 39, 40, 41,  
43, 44, 47, 48, 49, 50, 53, 54, 55, 56, 57, 58, 60,  
62, 65, 66, 67, 71, 75, 76, 79, 96, 149, 150  
Prescription Drug Marketing Act of 1987, 62  
prescription drug order, 11, 12, 14, 15, 19, 24, 25, 26,  
27, 31, 32, 66, 71, 76, 77, 78, 79, 80, 81, 83, 84,  
85, 89, 90, 91, 96, 97, 98, 102, 104, 106,

prescription monitoring program, 85, 140, 163, 164,  
165, 166, 167  
prescription processing, centralized, 11, 39, 63, 69, 70  
protected health information, 13, 16, 17, 19, 21, 25,  
34, 43, 44, 55, 61, 63, 71, 73, 78, 79, 82, 86, 87,  
92, 96, 149, 165

## Q

quality-related event, 11, 12, 26, 87, 88, 152, 153, 156,  
157, 159, 160, 161

## R

reference product, 10, 26, 80  
remote dispensing site, 26, 91, 168, 169  
remote pharmacy, 89, 91, 92  
repackage, 16, 19, 26, 29, 31, 32, 33, 58, 59, 61, 62,  
96, 104, 108, 109, 110, 111, 112, 114, 115, 116,  
117, 118, 132, 142

## S

standard of care, 22, 28, 96, 99  
sterile pharmaceutical, 27, 28, 59, 104, 105  
substances, added, 9, 12  
suspicious order, 28, 114, 115, 118

## T

telepharmacy, 23, 47, 58, 91, 168, 169  
therapeutic interchange, 28, 69  
third-party logistics provider, 21, 28, 29, 32, 33, 58,  
62, 96, 108, 109, 110, 111, 112, 113, 114, 115,  
116, 117, 118, 120

## U

United States Pharmacopeia, 10, 14, 31, 59, 70, 71,  
94, 97, 104, 105, 113, 114, 121, 122, 123, 128,  
129, 143

## V

veterinary dispensing, 31  
virtual manufacturer, 19, 31  
virtual wholesale distributor, 31, 33

## W

wholesale distribution, 11, 12, 13, 14, 15, 16, 21, 29,  
30, 31, 32, 33, 40, 46, 58, 60, 61, 62, 63, 96, 108,  
109, 110, 111, 112, 113, 114, 115, 116, 117, 118,  
119, 120, 121, 122, 123, 124, 125, 126, 127, 128,  
129, 130, 131, 133, 134