

## 1 VIRGINIA ACTS OF ASSEMBLY — CHAPTER

2 *An Act to amend and reenact §§ 8.01-413, 32.1-127.1:01, 32.1-127.1:03, 42.1-77, 54.1-111,*  
 3 *54.1-2403.2, 54.1-2403.3, 54.1-3401, 54.1-3434, and 54.1-3434.01 of the Code of Virginia, relating*  
 4 *to the practice of pharmacy; penalties.*

5 [S 560]

6 Approved

7 **Be it enacted by the General Assembly of Virginia:**

8 **1. That §§ 8.01-413, 32.1-127.1:01, 32.1-127.1:03, 42.1-77, 54.1-111, 54.1-2403.2, 54.1-2403.3,**  
 9 **54.1-3401, 54.1-3434, and 54.1-3434.01 of the Code of Virginia are amended and reenacted as**  
 10 **follows:**

11 § 8.01-413. Certain copies of health care provider's records or papers of patient admissible; right of  
 12 patient or his attorney to copies of such records or papers; subpoena; damages, costs and attorney's fees.

13 A. In any case where the hospital, nursing facility, physician's, or other health care provider's original  
 14 records or papers of any patient in a hospital or institution for the treatment of physical or mental illness  
 15 are admissible or would be admissible as evidence, any typewritten copy, photograph, photostatted copy,  
 16 or microphotograph or printout or other hard copy generated from computerized or other electronic  
 17 storage, microfilm, or other photographic, mechanical, electronic or chemical storage process thereof  
 18 shall be admissible as evidence in any court of this Commonwealth in like manner as the original, if the  
 19 printout or hard copy or microphotograph or photograph is properly authenticated by the employees  
 20 having authority to release or produce the original records.

21 Any hospital, nursing facility, physician, or other health care provider whose records or papers  
 22 relating to any such patient are subpoenaed for production under this section or the Rules of the  
 23 Supreme Court of Virginia may comply with the subpoena by a timely mailing to the clerk issuing the  
 24 subpoena properly authenticated copies, photographs or microphotographs in lieu of the originals. The  
 25 court whose clerk issued the subpoena may, after notice to such hospital, nursing facility, physician, or  
 26 other health care provider, enter an order requiring production of the originals, if available, of any stored  
 27 records or papers whose copies, photographs or microphotographs are not sufficiently legible. The party  
 28 requesting the subpoena shall be liable for the reasonable charges of the hospital, nursing facility,  
 29 physician, or other health care provider for the service of maintaining, retrieving, reviewing, preparing,  
 30 copying and mailing the items produced. Except for copies of X-ray photographs, however, such charges  
 31 shall not exceed fifty cents for each page up to fifty pages and twenty-five cents a page thereafter for  
 32 copies from paper and one dollar per page for copies from microfilm or other micrographic process,  
 33 plus all postage and shipping costs and a search and handling fee not to exceed ten dollars.

34 B. Copies of hospital, nursing facility, physician's, or other health care provider's records or papers  
 35 shall be furnished within fifteen days of such request to the patient or his attorney upon such patient's or  
 36 attorney's written request, which request shall comply with the requirements of § 32.1-127.1:03.  
 37 However, copies of a patient's records shall not be furnished to such patient where the patient's treating  
 38 physician has made a part of the patient's records a written statement that in his opinion the furnishing  
 39 to or review by the patient of such records would be injurious to the patient's health or well-being, but  
 40 in any such case such records shall be furnished to the patient's attorney within fifteen days of the date  
 41 of such request. A reasonable charge may be made for the service of maintaining, retrieving, reviewing  
 42 and preparing such copies. Except for copies of X-ray photographs, however, such charges shall not  
 43 exceed fifty cents per page for up to fifty pages and twenty-five cents a page thereafter for copies from  
 44 paper and one dollar per page for copies from microfilm or other micrographic process, plus all postage  
 45 and shipping costs and a search and handling fee not to exceed ten dollars. Any hospital, nursing  
 46 facility, physician, or other health care provider receiving such a request from a patient's attorney shall  
 47 require a writing signed by the patient confirming the attorney's authority to make the request.

48 C. Upon the failure of any hospital, nursing facility, physician, or other health care provider to  
 49 comply with any written request made in accordance with subsection B within the period of time  
 50 specified in that subsection and within the manner specified in § 32.1-127.1:03, the patient or his  
 51 attorney may by affidavit filed with the clerk of the circuit court wherein any eventual suit, if any,  
 52 would be required to be filed, upon payment of the fees required by subdivision (23) of § 14.1-112, and  
 53 fees for service, request that the clerk subpoena such records or papers. The clerk shall thereupon issue  
 54 a subpoena, returnable within twenty days of proper service, directing the hospital, nursing facility,  
 55 physician, or other health care provider to produce and furnish copies of the reports and papers to him,  
 56 whereupon, the clerk shall make the same available to the patient or his attorney. If the court finds that

57 a hospital, nursing facility, physician, or other health care provider willfully refused to comply with a  
 58 written request made in accordance with subsection B, either by willfully or arbitrarily refusing or by  
 59 imposing a charge in excess of the reasonable expense of making the copies and processing the request  
 60 for records, the court may award damages for all expenses incurred by the patient to obtain such copies,  
 61 including court costs and reasonable attorney's fees.

62 D. The provisions of subsections A, B, and C hereof shall apply to any health care provider whose  
 63 office is located within or without the Commonwealth if the records pertain to any patient who is a  
 64 party to a cause of action in any court in the Commonwealth of Virginia, and shall apply only to  
 65 requests made by an attorney, or his client, in anticipation of litigation or in the course of litigation.

66 E. Health care provider, as used in this section, shall have the same meaning as provided in  
 67 § 32.1-127.1:03 and shall also include an independent medical copy retrieval service contracted to  
 68 provide the service of retrieving, reviewing, and preparing such copies for distribution.

69 *F. Notwithstanding the authorization to admit as evidence patient records in the form of*  
 70 *microphotographs, prescription dispensing records maintained in or on behalf of any pharmacy*  
 71 *registered or permitted in Virginia shall only be stored in compliance with §§ 54.1-3410, 54.1-3411 and*  
 72 *54.1-3412.*

73 § 32.1-127.1:01. Record storage.

74 A. Medical records, as defined in § 42.1-77, may be stored by computerized or other electronic  
 75 process or microfilm, or other photographic, mechanical, or chemical process; however, the stored record  
 76 shall identify the location of any documents or information that could not be so technologically stored.  
 77 If the technological storage process creates an unalterable record, the nursing facility, hospital or other  
 78 licensed health care provider shall not be required to maintain paper copies of medical records that have  
 79 been stored by computerized or other electronic process, microfilm, or other photographic, mechanical,  
 80 or chemical process. Upon completing such technological storage, paper copies of medical records may  
 81 be destroyed in a manner that preserves the patient's confidentiality. However, any documents or  
 82 information that could not be so technologically stored shall be preserved.

83 *B. Notwithstanding the authority of this section to copy patient records in the form of microfilm,*  
 84 *prescription dispensing records maintained in or on behalf of any pharmacy registered or permitted in*  
 85 *Virginia shall only be stored in compliance with §§ 54.1-3410, 54.1-3411 and 54.1-3412.*

86 § 32.1-127.1:03. Patient Health Records Privacy.

87 A. There is hereby recognized a patient's right of privacy in the content of a patient's medical record.  
 88 Patient records are the property of the provider maintaining them, and, except when permitted by this  
 89 section or by another provision of state or federal law, no provider, or other person working in a health  
 90 care setting, may disclose the records of a patient.

91 Patient records shall not be removed from the premises where they are maintained without the  
 92 approval of the provider, except in accordance with a court order or subpoena consistent with § 8.01-413  
 93 C or with this section *or in accordance with the regulations relating to change of ownership of patient*  
 94 *records promulgated by a health regulatory board established in Title 54.1.*

95 No third party to whom disclosure of patient records was made by a provider shall redisclose or  
 96 otherwise reveal the records of a patient, beyond the purpose for which such disclosure was made,  
 97 without first obtaining the patient's specific consent to such redisclosure. This redisclosure prohibition  
 98 shall not, however, prevent any provider who receives records from another provider from making  
 99 subsequent disclosures permitted under this section.

100 B. As used in this section:

101 "Agent" means a person who has been appointed as a patient's agent under a power of attorney for  
 102 health care or an advance directive under the Health Care Decisions Act (§ 54.1-2981 et seq.)

103 "Guardian" means a court-appointed guardian of the person.

104 "Health services" includes but is not limited to examination, diagnosis, evaluation, treatment,  
 105 pharmaceuticals, aftercare, habilitation or rehabilitation and mental health therapy of any kind.

106 "Parent" means a biological, adoptive or foster parent.

107 "Patient" means a person who is receiving or has received health services from a provider.

108 "Provider" shall have the same meaning as set forth in the definition of "Health care provider" in  
 109 § 8.01-581.1, except that state-operated facilities shall also be considered providers for the purposes of  
 110 this section. Provider shall also include all ~~individuals~~ *persons* who are licensed ~~or~~ certified, *registered*  
 111 *or permitted* by any of the health regulatory boards within the Department of Health Professions, except  
 112 ~~individuals~~ *persons* regulated by the Board of Funeral Directors and Embalmers or the Board of  
 113 Veterinary Medicine.

114 "Record" means any written, printed or electronically recorded material maintained by a provider in  
 115 the course of providing health services to a patient concerning the patient and the services provided.  
 116 "Record" also includes the substance of any communication made by a patient to a provider in  
 117 confidence during or in connection with the provision of health services to a patient or information

118 otherwise acquired by the provider about a patient in confidence and in connection with the provision of  
119 health services to the patient.

120 C. The provisions of this section shall not apply to any of the following:

121 1. The status of and release of information governed by §§ 65.2-604 and 65.2-607 of the Virginia  
122 Workers Compensation Act; or

123 2. Except where specifically provided herein, the records of minor patients.

124 D. Providers may disclose the records of a patient:

125 1. As set forth in subsection E of this section, pursuant to the written consent of the patient or in the  
126 case of a minor patient, his custodial parent, guardian or other person authorized to consent to treatment  
127 of minors pursuant to § 54.1-2969; also, in emergency cases or situations where it is impractical to  
128 obtain the patient's written consent, pursuant to the patient's oral consent for a provider to discuss the  
129 patient's records with a third party specified by the patient;

130 2. In compliance with a subpoena issued in accord with subsection H of this section, pursuant to  
131 court order upon good cause shown or in compliance with a subpoena issued pursuant to subsection C  
132 of § 8.01-413;

133 3. In accord with subsection F of § 8.01-399 including, but not limited to, situations where disclosure  
134 is reasonably necessary to establish or collect a fee or to defend a provider or the provider's employees  
135 or staff against any accusation of wrongful conduct; also as required in the course of an investigation,  
136 audit, review or proceedings regarding a provider's conduct by a duly authorized law-enforcement,  
137 licensure, accreditation, or professional review entity;

138 4. In testimony in accordance with §§ 8.01-399 and 8.01-400.2;

139 5. In compliance with the provisions of § 8.01-413;

140 6. As required or authorized by any other provision of law including contagious disease, public  
141 safety, and suspected child or adult abuse reporting requirements, including but not limited to those  
142 contained in §§ 32.1-36, 32.1-36.1, 32.1-40, 32.1-41, 32.1-276.5, 32.1-283, 32.1-283.1, 37.1-98.2,  
143 53.1-40.10, 54.1-2403.3, 54.1-2906, 54.1-2907, 54.1-2966, 54.1-2966.1, 54.1-2967, 54.1-2968, 63.1-55.3  
144 and 63.1-248.11;

145 7. Where necessary in connection with the care of the patient;

146 8. In the normal course of business in accordance with accepted standards of practice within the  
147 health services setting; *however, the maintenance, storage, and disclosure of the mass of prescription*  
148 *dispensing records maintained in a pharmacy registered or permitted in Virginia shall only be*  
149 *accomplished in compliance with §§ 54.1-3410, 54.1-3411 and 54.1-3412;*

150 9. When the patient has waived his right to the privacy of the medical records;

151 10. When examination and evaluation of a patient is undertaken pursuant to judicial or administrative  
152 law order, but only to the extent as required by such;

153 11. To the guardian ad litem in the course of a guardianship proceeding of an adult patient  
154 authorized under §§ 37.1-128.1, 37.1-128.2 and 37.1-132;

155 12. To the attorney appointed by the court to represent a patient in a civil commitment proceeding  
156 under § 37.1-67.3;

157 13. To the attorney and/or guardian ad litem of a minor patient who represents such minor in any  
158 judicial or administrative proceeding, provided that the court or administrative hearing officer has  
159 entered an order granting the attorney or guardian ad litem this right and such attorney or guardian ad  
160 litem presents evidence to the provider of such order;

161 14. With regard to the Court Appointed Special Advocate (CASA) program, a minor's records in  
162 accord with § 9-173.12;

163 15. To an agent appointed under a patient's power of attorney or to an agent or decision maker  
164 designated in a patient's advance directive for health care or to any other person consistent with the  
165 provisions of the Health Care Decisions Act (§ 54.1-2981 et seq.);

166 16. To third-party payors and their agents pursuant to the deemed consent provisions of §§ 37.1-226  
167 and 37.1-227 when the patient has requested the provider to submit bills to the third-party payor for  
168 payment under a contract or insurance policy;

169 17. As is necessary to support an application for receipt of health care benefits from a governmental  
170 agency or as required by an authorized governmental agency reviewing such application or reviewing  
171 benefits already provided;

172 18. Upon the sale of a medical practice as provided in § 54.1-2405; *or upon a change of ownership*  
173 *or closing of a pharmacy pursuant to regulations of the Board of Pharmacy;*

174 19. In accord with § 54.1-2400.1 B, to communicate a patient's specific and immediate threat to  
175 cause serious bodily injury or death of an identified or readily identifiable person;

176 20. To the patient, except as provided in subsections E and F of this section and subsection B of  
177 § 8.01-413;

178 21. In the case of substance abuse records when permitted by and in conformity with requirements of

179 federal law found in 42 U.S.C. 290dd-2 and 42 C.F.R. Part 2;

180 22. In connection with the work of any entity established as set forth in § 8.01-581.16 to evaluate the  
181 adequacy or quality of professional services or the competency and qualifications for professional staff  
182 privileges; and

183 23. Records of a deceased or mentally incapacitated patient to the personal representative or executor  
184 of the deceased patient or the legal guardian or committee of the incompetent or incapacitated patient or  
185 if there is no such person appointed, to the following persons in the following order of priority: a  
186 spouse, an adult son or daughter, either parent, an adult brother or sister, or any other relative of the  
187 deceased patient in order of blood relationship; and

188 ~~24. Pursuant to a medical temporary detention order as set out in subsection M of § 37.1-134.5.~~

189 E. Requests for copies of medical records shall (i) be in writing, dated and signed by the requester;  
190 (ii) identify the nature of the information requested; and (iii) include evidence of the authority of the  
191 requester to receive such copies and identification of the person to whom the information is to be  
192 disclosed. Within fifteen days of receipt of a request for copies of medical records, the provider shall do  
193 one of the following: (i) furnish such copies to any requester authorized to receive them; (ii) inform the  
194 requester if the information does not exist or cannot be found; (iii) if the provider does not maintain a  
195 record of the information, so inform the requester and provide the name and address, if known, of the  
196 provider who maintains the record; or (iv) deny the request (a) under subsection F, (b) on the grounds  
197 that the requester has not established his authority to receive such records or proof of his identity, or (c)  
198 as otherwise provided by law. Procedures set forth in this section shall apply only to requests for  
199 records not specifically governed by other provisions of this Code or of federal law.

200 F. Except as provided in subsection B of § 8.01-413, copies of a patient's records shall not be  
201 furnished to such patient or anyone authorized to act on the patient's behalf where the patient's attending  
202 physician or the patient's clinical psychologist has made a part of the patient's record a written statement  
203 that, in his opinion, the furnishing to or review by the patient of such records would be injurious to the  
204 patient's health or well-being. If any custodian of medical records denies a request for copies of records  
205 based on such statement, the custodian shall permit examination and copying of the medical record by  
206 another such physician or clinical psychologist selected by the patient, whose licensure, training and  
207 experience relative to the patient's condition is at least equivalent to that of the physician or clinical  
208 psychologist upon whose opinion the denial is based. The person or entity denying the request shall  
209 inform the patient of the patient's right to select another reviewing physician or clinical psychologist  
210 under this subsection who shall make a judgment as to whether to make the record available to the  
211 patient. Any record copied for review by the physician or clinical psychologist selected by the patient  
212 shall be accompanied by a statement from the custodian of the record that the patient's attending  
213 physician or clinical psychologist determined that the patient's review of his record would be injurious to  
214 the patient's health or well-being.

215 G. A written consent to allow release of patient records may, but need not, be in the following form:

216  
217 CONSENT TO RELEASE OF CONFIDENTIAL HEALTH CARE INFORMATION  
218  
219 Patient Name.....  
220  
221 Provider Name.....  
222  
223 Person, agency or provider to whom  
224  
225 disclosure is to be made.....  
226  
227 Information or Records to be disclosed.....  
228

229 As the person signing this consent, I understand that I am giving my  
230  
231 permission to the above-named provider or other named third party for  
232  
233 disclosure of confidential health care records. I also understand that  
234  
235 I have the right to revoke this consent, but that my revocation is not  
236  
237 effective until delivered in writing to the person who is in possession

238  
 239 of my records. A copy of this consent and a notation concerning the  
 240  
 241 persons or agencies to whom disclosure was made shall be included  
 242  
 243 with my original records. The person who receives the records to  
 244  
 245 which this consent pertains may not redisclose them to anyone else  
 246  
 247 without my separate written consent unless such recipient is a  
 248  
 249 provider who makes a disclosure permitted by law.

250  
 251 This consent expires on (date).....  
 252  
 253 Signature of Patient ..... Date .....

254  
 255 H. 1. No party to an action shall request the issuance of a subpoena duces tecum for an opposing  
 256 party's medical records unless a copy of the request for the subpoena is provided to opposing counsel or  
 257 the opposing party if they are pro se, simultaneously with filing the request. No party to an action shall  
 258 request the issuance of a subpoena duces tecum for the medical records of a nonparty witness unless a  
 259 copy of the request for the subpoena is provided to the nonparty witness simultaneously with filing the  
 260 request.

261 In instances where medical records being subpoenaed are those of a pro se party or nonparty witness,  
 262 the party requesting the issuance of the subpoena shall deliver to the pro se party or nonparty witness  
 263 together with the copy of the request for subpoena, a statement informing them of their rights and  
 264 remedies. The statement shall include the following language and the heading shall be in boldface  
 265 capital letters:

266 **NOTICE TO PATIENT**

267 **The attached Request for Subpoena means that (insert name of party requesting subpoena) has**  
 268 **asked the court to issue a subpoena to your doctor or other health care providers (names of health**  
 269 **care providers inserted here) requiring them to produce your medical records. Your doctor or**  
 270 **other health care provider is required to respond by providing a copy of your medical records. If**  
 271 **you believe your records should not be disclosed and object to their disclosure, you have the right**  
 272 **to file a motion with the clerk of the court to quash the subpoena. You may contact the clerk's**  
 273 **office to determine the requirements that must be satisfied when filing a motion to quash and you**  
 274 **may elect to contact an attorney to represent your interest. If you elect to file a motion to quash,**  
 275 **it must be filed as soon as possible before the provider sends out the records in response to the**  
 276 **subpoena. If you elect to file a motion to quash, you must notify your doctor or other health care**  
 277 **provider(s) that you are filing the motion so that the provider knows to send the records to the**  
 278 **clerk of court in a sealed envelope or package for safekeeping while your motion is decided.**

279 2. Any party filing a request for a subpoena duces tecum for a patient's medical records shall include  
 280 a Notice to Providers in the same part of the request where the provider is directed where and when to  
 281 return the records. Such notice shall be in boldface capital letters and shall include the following  
 282 language:

283 **NOTICE TO PROVIDERS**

284 **IF YOU RECEIVE NOTICE THAT YOUR PATIENT HAS FILED A MOTION TO QUASH**  
 285 **(OBJECTING TO) THIS SUBPOENA, OR IF YOU FILE A MOTION TO QUASH THIS**  
 286 **SUBPOENA, SEND THE RECORDS ONLY TO THE CLERK OF THE COURT WHICH**  
 287 **ISSUED THE SUBPOENA USING THE FOLLOWING PROCEDURE: PLACE THE RECORDS**  
 288 **IN A SEALED ENVELOPE AND ATTACH TO THE SEALED ENVELOPE A COVER LETTER**  
 289 **TO THE CLERK OF COURT WHICH STATES THAT CONFIDENTIAL HEALTH CARE**  
 290 **RECORDS ARE ENCLOSED AND ARE TO BE HELD UNDER SEAL PENDING THE**  
 291 **COURT'S RULING ON THE MOTION TO QUASH THE SUBPOENA. THE SEALED**  
 292 **ENVELOPE AND THE COVER LETTER SHALL BE PLACED IN AN OUTER ENVELOPE OR**  
 293 **PACKAGE FOR TRANSMITTAL TO THE COURT.**

294 3. Health care providers shall provide a copy of all records as required by a subpoena duces tecum  
 295 or court order for such medical records. If the health care provider has, however, actual receipt of notice  
 296 that a motion to quash the subpoena has been filed or if the health care provider files a motion to quash  
 297 the subpoena for medical records, then the health care provider shall produce the records to the clerk of

298 the court issuing the subpoena, where the court shall place the records under seal until a determination  
 299 is made regarding the motion to quash. The securely sealed envelope shall only be opened on order of  
 300 the judge. In the event the court grants the motion to quash, the records shall be returned to the health  
 301 care provider in the same sealed envelope in which they were delivered to the court. In the event that a  
 302 judge orders the sealed envelope to be opened to review the records in camera, a copy of the judge's  
 303 order shall accompany any records returned to the provider. The records returned to the provider shall  
 304 be in a securely sealed envelope.

305 4. It is the duty of any party requesting a subpoena duces tecum for medical records to determine  
 306 whether the patient whose records are sought is pro se or a nonparty. Any request for a subpoena duces  
 307 tecum for the medical records of a nonparty or of a pro se party shall direct the provider (in boldface  
 308 type) not to produce the records until ten days after the date on which the provider is served with the  
 309 subpoena duces tecum and shall be produced no later than twenty days after the date of such service.

310 In the event that the individual whose records are being sought files a motion to quash the subpoena,  
 311 the court shall decide whether good cause has been shown by the discovering party to compel disclosure  
 312 of the patient's private records over the patient's objections. In determining whether good cause has been  
 313 shown, the court shall consider (i) the particular purpose for which the information was collected; (ii)  
 314 the degree to which the disclosure of the records would embarrass, injure, or invade the privacy of the  
 315 individual; (iii) the effect of the disclosure on the individual's future health care; (iv) the importance of  
 316 the information to the lawsuit or proceeding; and (v) any other relevant factor.

317 The provisions of this subsection have no application to subpoenas for medical records requested  
 318 under § 8.01-413, or issued by a duly authorized administrative agency conducting an investigation,  
 319 audit, review or proceedings regarding a provider's conduct. The provisions of this subsection apply to  
 320 the medical records of both minors and adults.

321 A subpoena for substance abuse records must conform to the requirements of federal law found in 42  
 322 C.F.R. Part 2, Subpart E.

323 Providers may testify about the medical records of a patient in compliance with §§ 8.01-399 and  
 324 8.01-400.2.

325 § 42.1-77. Definitions.

326 As used in this chapter:

327 "Agency" means all boards, commissions, departments, divisions, institutions, authorities, or parts  
 328 thereof, of the Commonwealth or its political subdivisions and includes the offices of constitutional  
 329 officers.

330 "Archival quality" means a quality of reproduction consistent with established standards specified by  
 331 state and national agencies and organizations responsible for establishing such standards, such as the  
 332 Association for Information and Image Management, the American Standards Association, and the  
 333 National Bureau of Standards.

334 "Board" means the State Library Board.

335 "Council" means the State Public Records Advisory Council.

336 "Custodian" means the public official in charge of an office having public records.

337 "Data" means symbols, or representations, of facts or ideas that can be communicated, interpreted, or  
 338 processed by manual or automated means.

339 "Database" means a set of data, consisting of one file or a group of integrated files, maintained as an  
 340 information system managed by a database management system.

341 "Database management system" means a set of software programs that controls the organization,  
 342 storage and retrieval of data in a database. It also controls the security and integrity of the database.

343 "Electronic record" means any information that is recorded in machine readable form.

344 "Electronic records system" means any information system that produces, processes, or stores records  
 345 by using a computer, and is also called an automated information system.

346 "Information system" means the organized collection, processing, transmission, and dissemination of  
 347 information in accordance with defined procedures, whether automated or manual.

348 "State Librarian" means the State Librarian or his designated representative.

349 "Public official" means all persons holding any office created by the Constitution of Virginia or by  
 350 any act of the General Assembly, the Governor and all other officers of the executive branch of the state  
 351 government, and all other officers, heads, presidents or chairmen of boards, commissions, departments,  
 352 and agencies of the state government or its political subdivisions.

353 "Public record" means recorded information that documents a transaction or activity by or with any  
 354 public officer, agency or employee of the state government or its political subdivisions. Regardless of  
 355 physical form or characteristic, the recorded information is a public record if it is produced, collected,  
 356 received or retained in pursuance of law or in connection with the transaction of public business.

357 The medium on which such information is recorded may be, but is not limited to paper, film,  
 358 magnetic, optical or solid state devices which can store electronic signals, tapes, mylar, linen, silk or

359 vellum. The general types of records may be, but are not limited to books, papers, letters, documents,  
 360 printouts, photographs, films, tapes, microfiche, microfilm, photostats, sound recordings, maps, drawings,  
 361 and any representations held in computer memory.

362 Nonrecord materials, meaning reference books and exhibit materials made or acquired and preserved  
 363 solely for reference use or exhibition purposes, extra copies of documents preserved only for  
 364 convenience or reference, and stocks of publications, shall not be included within the definition of  
 365 public records as used in this chapter.

366 "Archival records" means all noncurrent records of continuing and enduring value useful to the  
 367 citizens of the Commonwealth and necessary to the administrative functions of public agencies in the  
 368 conduct of services and activities mandated by law. In appraisal of public records deemed archival, the  
 369 terms "administrative," "legal," "fiscal," and "historical" shall be defined as:

370 1. "Administrative value" : Records shall be deemed of administrative value if they have continuing  
 371 utility in the operation of an agency.

372 2. "Legal value" : Records shall be deemed of legal value when they document actions taken in the  
 373 protection and proving of legal or civil rights and obligations of individuals and agencies.

374 3. "Fiscal value" : Records shall be deemed of fiscal value so long as they are needed to document  
 375 and verify financial authorizations, obligations and transactions.

376 4. "Historical value" : Records shall be deemed of historical value when they contain unique  
 377 information, regardless of age, which provides understanding of some aspect of the government and  
 378 promotes the development of an informed and enlightened citizenry.

379 "Medical records" means the documentation of health care services, whether physical or mental,  
 380 rendered by direct or indirect patient-provider interaction which is used as a mechanism for tracking the  
 381 patient's health care status. Medical records may be technologically stored by computerized or other  
 382 electronic process, or through microfilm or other similar photographic form or chemical process.  
 383 *Notwithstanding the authority provided by this definition to store medical records on microfilm or other*  
 384 *similar photographic form or chemical process, prescription dispensing records maintained in or on*  
 385 *behalf of any pharmacy registered or permitted in Virginia shall only be stored in compliance with*  
 386 *§§ 54.1-3410, 54.1-3411 and 54.1-3412.*

387 "Official records" means public records.

388 "Persons under a disability" means persons so defined under subsection A of § 8.01-229.

389 "Preservation" means maintaining archival records in their original physical form by stabilizing them  
 390 chemically or strengthening them physically to ensure their survival as long as possible in their original  
 391 form. It also means the reformatting of written, printed, electronic or visual archival information to  
 392 extend the life of the information.

393 "Retention and disposition schedule" means an approved timetable stating the retention time period  
 394 and disposition action of records series.

395 "Software programs" means the written specifications used to operate an electronic records system as  
 396 well as the documentation describing implementation strategies.

397 § 54.1-111. Unlawful acts; prosecution; proceedings in equity.

398 A. It shall be unlawful for any person, partnership, corporation or other entity to engage in any of  
 399 the following acts:

400 1. Practicing a profession or occupation without holding a valid license as required by statute or  
 401 regulation.

402 2. Making use of any designation provided by statute or regulation to denote a standard of  
 403 professional or occupational competence without being duly certified or licensed.

404 3. Making use of any titles, words, letters or abbreviations which may reasonably be confused with a  
 405 designation provided by statute or regulation to denote a standard of professional or occupational  
 406 competence without being duly certified or licensed.

407 4. Performing any act or function which is restricted by statute or regulation to persons holding a  
 408 professional or occupational license or certification, without being duly certified or licensed.

409 5. Failing to register as a practitioner of a profession or occupation as required by statute or  
 410 regulation.

411 6. Materially misrepresenting facts in an application for licensure, certification or registration.

412 7. Willfully refusing to furnish a regulatory board information or records required or requested  
 413 pursuant to statute or regulation.

414 8. Violating any statute or regulation governing the practice of any profession or occupation  
 415 regulated pursuant to this title.

416 9. *Refusing to process a request, tendered in accordance with the regulations of the relevant health*  
 417 *regulatory board or applicable statutory law, for patient records or prescription dispensing records after*  
 418 *the closing of a business or professional practice or the transfer of ownership of a business or*  
 419 *professional practice.*

420 Any person who willfully engages in any unlawful act enumerated in this section shall be guilty of a  
 421 Class 1 misdemeanor. The third or any subsequent conviction for violating this section during a  
 422 thirty-six-month period shall constitute a Class 6 felony.

423 B. In addition to the criminal penalties provided for in subsection A of this section, the Department  
 424 of Professional and Occupational Regulation or the Department of Health Professions, without  
 425 compliance with the Administrative Process Act (§ 9-6.14:1 et seq.), shall have the authority to enforce  
 426 the provisions of subsection A of this section and may institute proceedings in equity to enjoin any  
 427 person, partnership, corporation or any other entity from engaging in any unlawful act enumerated in  
 428 this section. Such proceedings shall be brought in the name of the Commonwealth by the appropriate  
 429 Department in the circuit court of the city or county in which the unlawful act occurred or in which the  
 430 defendant resides.

431 *C. This section shall not be construed to prohibit or prevent the owner of patient records from (i)*  
 432 *retaining copies of his patient records or prescription dispensing records after the closing of a business*  
 433 *or professional practice or the transfer of ownership of a business or professional practice or (ii)*  
 434 *charging a reasonable fee, not in excess of the amounts authorized in § 8.01-413, for copies of patient*  
 435 *records.*

436 § 54.1-2403.2. Record storage.

437 A. Medical records, as defined in § 42.1-77, may be stored by computerized or other electronic  
 438 process or microfilm, or other photographic, mechanical, or chemical process; however, the stored record  
 439 shall identify the location of any documents or information that could not be so technologically stored.  
 440 If the technological storage process creates an unalterable record, a health care provider licensed,  
 441 certified or registered by a health regulatory board within the Department shall not be required to  
 442 maintain paper copies of medical records that have been stored by computerized or other electronic  
 443 process, microfilm, or other photographic, mechanical, or chemical process. Upon completing such  
 444 technological storage, paper copies of medical records may be destroyed in a manner that preserves the  
 445 patient's confidentiality. However, any documents or information that could not be so technologically  
 446 stored shall be preserved.

447 *B. Notwithstanding the authority given in this section to store patient records in the form of*  
 448 *microfilm, prescription dispensing records maintained in or on behalf of any pharmacy registered or*  
 449 *permitted in Virginia shall only be stored in compliance with §§ 54.1-3410, 54.1-3411 and 54.1-3412.*

450 § 54.1-2403.3. Medical records; ownership; provision of copies.

451 Medical records maintained by any health care provider as defined in § 32.1-127.1:03 shall be the  
 452 property of such health care provider *or, in the case of a health care provider employed by another*  
 453 *health care provider, the property of the employer.* Such health care provider shall release copies of any  
 454 such medical records in compliance with § 32.1-127.1:03 or § 8.01-413, if the request is made for  
 455 purposes of litigation, or as otherwise provided by state or federal law.

456 § 54.1-3401. Definitions.

457 As used in this chapter, unless the context requires a different meaning:

458 "Administer" means the direct application of a controlled substance, whether by injection, inhalation,  
 459 ingestion or any other means, to the body of a patient or research subject by (i) a practitioner or by his  
 460 authorized agent and under his direction, or (ii) the patient or research subject at the direction and in the  
 461 presence of the practitioner.

462 "Advertisement" means all representations disseminated in any manner or by any means, other than  
 463 by labeling, for the purpose of inducing, or which are likely to induce, directly or indirectly, the  
 464 purchase of drugs or devices.

465 "Anabolic steroid" means any drug or hormonal substance, chemically and pharmacologically related  
 466 to testosterone, other than estrogens, progestins, and corticosteroids, that promotes muscle growth.

467 "Animal" means any nonhuman animate being endowed with the power of voluntary action.

468 "Agent" means an authorized person who acts on behalf of or at the direction of a manufacturer,  
 469 distributor, or dispenser. It does not include a common or contract carrier, public warehouseman, or  
 470 employee of the carrier or warehouseman.

471 "Board" means the Board of Pharmacy.

472 *"Change of ownership" of an existing entity permitted, registered or licensed by the Board means (i)*  
 473 *the sale or transfer of all or substantially all of the assets of the entity or of any corporation that owns*  
 474 *or controls the entity; (ii) the creation of a partnership by a sole proprietor, the dissolution of a*  
 475 *partnership, or change in partnership composition; (iii) the acquisition or disposal of fifty percent or*  
 476 *more of the outstanding shares of voting stock of a corporation owning the entity or of the parent*  
 477 *corporation of a wholly owned subsidiary owning the entity, except that this shall not apply to any*  
 478 *corporation the voting stock of which is actively traded on any securities exchange or in any*  
 479 *over-the-counter market; (iv) the merger of a corporation owning the entity or of the parent corporation*  
 480 *of a wholly-owned subsidiary owning the entity with another business or corporation; or (v) the*

481 *expiration or forfeiture of a corporation's charter.*

482 "Compound" means the taking of two or more ingredients and fabricating them into a single  
483 preparation, usually referred to as a dosage form.

484 "Controlled substance" means a drug, substance or immediate precursor in Schedules I through VI of  
485 this chapter. The term shall not include distilled spirits, wine, malt beverages, or tobacco as those terms  
486 are defined or used in Title 3.1 or Title 4.1.

487 "DEA" means the Drug Enforcement Administration, United States Department of Justice, or its  
488 successor agency.

489 "Deliver" or "delivery" means the actual, constructive, or attempted transfer of any item regulated by  
490 this chapter, whether or not there exists an agency relationship.

491 "Device" means instruments, apparatus, and contrivances, including their components, parts and  
492 accessories, intended for use in the diagnosis, cure, mitigation, treatment, or prevention of disease in  
493 man or animals or to affect the structure or any function of the body of man or animals.

494 "Dialysis solution" means either the commercially available, unopened, sterile solutions whose  
495 purpose is to be instilled into the peritoneal cavity during the medical procedure known as peritoneal  
496 dialysis, or commercially available solutions whose purpose is to be used in the performance of  
497 hemodialysis, not to include any solutions administered to the patient intravenously.

498 "Dispense" means to deliver a drug to an ultimate user or research subject by or pursuant to the  
499 lawful order of a practitioner, including the prescribing and administering, packaging, labeling or  
500 compounding necessary to prepare the substance for that delivery.

501 "Dispenser" means a practitioner who dispenses.

502 "Distribute" means to deliver other than by administering or dispensing a controlled substance.

503 "Distributor" means a person who distributes.

504 "Drug" means (i) articles or substances recognized in the official United States Pharmacopoeia  
505 National Formulary or official Homeopathic Pharmacopoeia of the United States, or any supplement to  
506 any of them; (ii) articles or substances intended for use in the diagnosis, cure, mitigation, treatment or  
507 prevention of disease in man or animals; (iii) articles or substances, other than food, intended to affect  
508 the structure or any function of the body of man or animals; or (iv) articles or substances intended for  
509 use as a component of any article specified in clause (i), (ii) or (iii). "Drug" does not include devices or  
510 their components, parts or accessories.

511 "Hashish oil" means any oily extract containing one or more cannabinoids, but shall not include any  
512 such extract with a tetrahydrocannabinol content of less than twelve percent by weight.

513 "Immediate precursor" means a substance which the Board of Pharmacy has found to be and by  
514 regulation designates as being the principal compound commonly used or produced primarily for use,  
515 and which is an immediate chemical intermediary used or likely to be used in the manufacture of a  
516 controlled substance, the control of which is necessary to prevent, curtail, or limit manufacture.

517 "Label" means a display of written, printed or graphic matter upon the immediate container of any  
518 article. A requirement made by or under authority of this chapter that any word, statement or other  
519 information appear on the label shall not be considered to be complied with unless such word, statement  
520 or other information also appears on the outside container or wrapper, if any, of the retail package of  
521 such article, or is easily legible through the outside container or wrapper.

522 "Labeling" means all labels and other written, printed or graphic matter on an article or any of its  
523 containers or wrappers, or accompanying such article.

524 "Manufacture" means the production, preparation, propagation, compounding, conversion or  
525 processing of any item regulated by this chapter, either directly or indirectly by extraction from  
526 substances of natural origin, or independently by means of chemical synthesis, or by a combination of  
527 extraction and chemical synthesis, and includes any packaging or repackaging of the substance or  
528 labeling or relabeling of its container. This term does not include the preparing, compounding,  
529 packaging or labeling of a controlled substance by a practitioner as an incident to his administering or  
530 dispensing of a controlled substance or marijuana in the course of his professional practice, or by a  
531 practitioner, or by his authorized agent under his supervision, for the purpose of, or as an incident to,  
532 research, teaching, or chemical analysis and not for sale.

533 "Manufacturer" means every person who manufactures.

534 "Marijuana" means any part of a plant of the genus *Cannabis* whether growing or not, its seeds or  
535 resin; and every compound, manufacture, salt, derivative, mixture, or preparation of such plant, its seeds,  
536 or its resin. Marijuana shall not include any oily extract containing one or more cannabinoids unless  
537 such extract contains less than twelve percent of tetrahydrocannabinol by weight, or the mature stalks of  
538 such plant, fiber produced from such stalk, oil or cake made from the seeds of such plant, any other  
539 compound, manufacture, salt, derivative, mixture or preparation of such mature stalks, fiber, oil, or cake,  
540 or the sterilized seed of such plant which is incapable of germination.

541 "Medical equipment supplier" means any person, as defined in § 1-13.19, engaged in the delivery to

542 the ultimate consumer, pursuant to the lawful order of a practitioner, of hypodermic syringes and  
 543 needles, medicinal oxygen, Schedule VI controlled devices, those Schedule VI controlled substances with  
 544 no medicinal properties which are used for the operation and cleaning of medical equipment and  
 545 solutions for peritoneal dialysis.

546 "Narcotic drug" means any of the following, whether produced directly or indirectly by extraction  
 547 from substances of vegetable origin, or independently by means of chemical synthesis, or by a  
 548 combination of extraction and chemical synthesis: (i) opium, opiates, and any salt, compound, derivative,  
 549 or preparation of opium or opiates; (ii) any salt, compound, isomer, derivative, or preparation thereof  
 550 which is chemically equivalent or identical with any of the substances referred to in clause (i), but not  
 551 including the isoquinoline alkaloids of opium; (iii) opium poppy and poppy straw; (iv) coca leaves and  
 552 any salt, compound, derivative, or preparation of coca leaves, and any salt, compound, isomer,  
 553 derivative, or preparation thereof which is chemically equivalent or identical with any of these  
 554 substances, but not including decocainized coca leaves or extraction of coca leaves which do not contain  
 555 cocaine or ecgonine.

556 "New drug" means: (i) any drug, except a new animal drug or an animal feed bearing or containing  
 557 a new animal drug, the composition of which is such that such drug is not generally recognized, among  
 558 experts qualified by scientific training and experience to evaluate the safety and effectiveness of drugs,  
 559 as safe and effective for use under the conditions prescribed, recommended, or suggested in the labeling,  
 560 except that such a drug not so recognized shall not be deemed to be a "new drug" if at any time prior  
 561 to the enactment of this chapter it was subject to the Food and Drugs Act of June 30, 1906, as  
 562 amended, and if at such time its labeling contained the same representations concerning the conditions  
 563 of its use; or (ii) any drug, except a new animal drug or an animal feed bearing or containing a new  
 564 animal drug, the composition of which is such that such drug, as a result of investigations to determine  
 565 its safety and effectiveness for use under such conditions, has become so recognized, but which has not,  
 566 otherwise than in such investigations, been used to a material extent or for a material time under such  
 567 conditions.

568 "Official compendium" means the official United States Pharmacopoeia National Formulary, official  
 569 Homeopathic Pharmacopoeia of the United States, or any supplement to any of them.

570 "Official written order" means an order written on a form provided for that purpose by the United  
 571 States Drug Enforcement Administration, under any laws of the United States making provision therefor,  
 572 if such order forms are authorized and required by federal law, and if no such order form is provided  
 573 then on an official form provided for that purpose by the Board of Pharmacy.

574 "Opiate" means any substance having an addiction-forming or addiction-sustaining liability similar to  
 575 morphine or being capable of conversion into a drug having such addiction-forming or  
 576 addiction-sustaining liability. It does not include, unless specifically designated as controlled under  
 577 Article 4 (§ 54.1-3437 et seq.) of this chapter, the dextrorotatory isomer of  
 578 3-methoxy-n-methylmorphinan and its salts (dextromethorphan). It does include its racemic and  
 579 levorotatory forms.

580 "Opium poppy" means the plant of the species *Papaver somniferum* L., except the seeds thereof.

581 "Original package" means the unbroken container or wrapping in which any drug or medicine is  
 582 enclosed together with label and labeling, put up by or for the manufacturer, wholesaler, or distributor  
 583 for use in the delivery or display of such article.

584 "Person" means both the plural and singular, as the case demands, and includes *an* individual,  
 585 partnership, corporation, association, governmental agency, trust, or other institution or entity.

586 "Poppy straw" means all parts, except the seeds, of the opium poppy, after mowing.

587 "Practitioner" means a physician, dentist, licensed nurse practitioner pursuant to § 54.1-2957.01,  
 588 licensed physician assistant pursuant to § 54.1-2952.1, pharmacist pursuant to § 54.1-3300, TPA-certified  
 589 optometrist pursuant to Article 5 (§ 54.1-3222 et seq.) of Chapter 32 of this title, veterinarian, scientific  
 590 investigator, or other person licensed, registered or otherwise permitted to distribute, dispense, prescribe  
 591 and administer, or conduct research with respect to, a controlled substance in the course of professional  
 592 practice or research in this Commonwealth.

593 "Prescriber" means a practitioner who is authorized pursuant to §§ 54.1-3303 and 54.1-3408 to issue  
 594 a prescription.

595 "Prescription" means an order for drugs or medical supplies, written or signed or transmitted by word  
 596 of mouth, telephone, telegraph or other means of communication to a pharmacist by a duly licensed  
 597 physician, dentist, veterinarian or other practitioner, authorized by law to prescribe and administer such  
 598 drugs or medical supplies.

599 "Prescription drug" means any drug required by federal law or regulation to be dispensed only  
 600 pursuant to a prescription, including finished dosage forms and active ingredients subject to § 503 (b) of  
 601 the federal Food, Drug, and Cosmetic Act.

602 "Production" or "produce" includes the manufacture, planting, cultivation, growing or harvesting of a

603 controlled substance or marijuana.

604 "Proprietary medicine" means a completely compounded nonprescription drug in its unbroken,  
605 original package which does not contain any controlled substance or marijuana as defined in this chapter  
606 and is not in itself poisonous, and which is sold, offered, promoted or advertised directly to the general  
607 public by or under the authority of the manufacturer or primary distributor, under a trademark, trade  
608 name or other trade symbol privately owned, and the labeling of which conforms to the requirements of  
609 this chapter and applicable federal law. However, this definition shall not include a drug which is only  
610 advertised or promoted professionally to licensed practitioners, a narcotic or drug containing a narcotic,  
611 a drug which may be dispensed only upon prescription or the label of which bears substantially the  
612 statement "Warning - may be habit-forming," or a drug intended for injection.

613 "Sale" includes barter, exchange, or gift, or offer therefor, and each such transaction made by any  
614 person, whether as individual, proprietor, agent, servant or employee.

615 "Warehouser" means any person, other than a wholesale distributor, engaged in the business of  
616 selling or otherwise distributing prescription drugs or devices to any person who is not the ultimate user  
617 or consumer. No person shall be subject to any state or local tax by reason of this definition.

618 "Wholesale distribution" means distribution of prescription drugs to persons other than consumers or  
619 patients, subject to the exceptions set forth in § 54.1-3401.1.

620 "Wholesale distributor" means any person engaged in wholesale distribution of prescription drugs  
621 including, but not limited to, manufacturers; repackers; own-label distributors; private-label distributors;  
622 jobbers; brokers; warehouses, including manufacturers' and distributors' warehouses, chain drug  
623 warehouses conducting wholesale distributions, and wholesale drug warehouses; independent wholesale  
624 drug traders; and retail pharmacies conducting wholesale distributions. No person shall be subject to any  
625 state or local tax as a wholesale merchant by reason of this definition.

626 The words "drugs" and "devices" as used in Chapter 33 (§ 54.1-3300 et seq.) of this title and in this  
627 chapter shall not include surgical or dental instruments, physical therapy equipment, X-ray apparatus or  
628 glasses or lenses for the eyes.

629 The terms "pharmacist," "pharmacy" and "practice of pharmacy" as used in this chapter shall be  
630 defined as provided in Chapter 33 of this title unless the context requires a different meaning.

631 § 54.1-3434. Permit to conduct pharmacy.

632 No person shall conduct a pharmacy without first obtaining a permit from the Board.

633 The application for such permit shall be made on a form provided by the Board and signed by a  
634 pharmacist who will be in full and actual charge of the pharmacy and who will be fully engaged in the  
635 practice of pharmacy at the location designated on the application.

636 The application shall (i) show the corporate name and trade name ~~and shall~~, (ii) list any pharmacist  
637 in addition to the pharmacist-in-charge practicing at the location indicated on the application, *and (iii)*  
638 *list the hours during which the pharmacy will be open to provide pharmacy services. Any change in the*  
639 *hours of operation, which is expected to last more than one week, shall be reported to the Board in*  
640 *writing and posted, at least fourteen days prior to the anticipated change, in a conspicuous place to*  
641 *provide notice to the public. The Board shall promulgate regulations to provide exceptions to this prior*  
642 *notification.*

643 If the owner is other than the pharmacist making the application, the type of ownership shall be  
644 indicated and shall list any partner or partners, and, if a corporation, then the corporate officers and  
645 directors. Further, if the owner is not a pharmacist, he shall not abridge the authority of the  
646 pharmacist-in-charge to exercise professional judgment relating to the dispensing of drugs in accordance  
647 with this act and Board regulations.

648 The permit shall be issued only to the pharmacist who signs the application as the  
649 pharmacist-in-charge and as such assumes the full responsibilities for the legal operation of the  
650 pharmacy. This permit and responsibilities shall not be construed to negate any responsibility of any  
651 pharmacist or other person.

652 Upon termination of practice by the pharmacist-in-charge, or upon any change in partnership  
653 composition, or upon the acquisition, *as defined in Board regulations*, of the existing corporation by  
654 another person *or the closing of a pharmacy*, the permit previously issued shall be immediately  
655 surrendered to the Board by the pharmacist-in-charge to whom it was issued, or by his legal  
656 representative, and an application for a new permit may be made in accordance with the requirements of  
657 this chapter.

658 The Board shall promulgate regulations (i) *defining acquisition of an existing permitted, registered or*  
659 *licensed facility or of any corporation under which the facility is directly or indirectly organized;* (ii)  
660 *providing for the transfer, confidentiality, integrity, and security of the pharmacy's prescription*  
661 *dispensing records and other patient records, regardless of where located; and (iii) establishing a*  
662 *reasonable time period for designation of a new pharmacist-in-charge. At the conclusion of ~~such~~the time*  
663 *period for designation of a new pharmacist-in-charge, a pharmacy which has failed to designate a new*

664 pharmacist-in-charge shall not operate as a pharmacy nor maintain a stock of prescription drugs on the  
 665 premises. The Director shall immediately notify the owner of record that the pharmacy no longer holds a  
 666 valid permit and that the owner shall make provision for the proper disposition of all Schedule II  
 667 through VI drugs and devices on the premises within fifteen days of receipt of this notice. At the  
 668 conclusion of the fifteen-day period, the Director or his authorized agent shall seize and indefinitely  
 669 secure all Schedule II through VI drugs and devices still on the premises, and notify the owner of such  
 670 seizure. The Director may properly dispose of the seized drugs and devices after six months from the  
 671 date of the notice of seizure if the owner has not claimed and provided for the proper disposition of the  
 672 property. The Board shall assess a fee of not less than the cost of storage of said drugs upon the owner  
 673 for reclaiming seized property.

674 The succeeding pharmacist-in-charge shall cause an inventory to be made of all Schedule I, II, III,  
 675 IV and V drugs on hand. Such inventory shall be completed as of the date he becomes  
 676 pharmacist-in-charge and prior to opening for business on that date.

677 The pharmacist to whom such permit is issued shall provide safeguards against diversion of all  
 678 controlled substances.

679 An application for a pharmacy permit shall be accompanied by a fee determined by the Board. All  
 680 permits shall expire on December 31 of each year.

681 Every pharmacy must be equipped so that prescriptions can be properly filled. The Board of  
 682 Pharmacy shall prescribe the minimum of such professional and technical equipment, which a pharmacy  
 683 shall at all times possess, and such list shall include as reference the latest revision of the United States  
 684 Pharmacopoeia Dispensing Information. No permit shall be issued or continued for the conduct of a  
 685 pharmacy until or unless there is compliance with the provisions of this chapter and regulations  
 686 promulgated by the Board.

687 Each day during which a person is in violation of this section shall constitute a separate offense.

688 § 54.1-3434.01. Notice of pharmacy closing; change of ownership; penalty.

689 A. Prior to the closing of a pharmacy for more than one week, the ~~pharmacist-in-charge~~ owner shall  
 690 either (i) post a conspicuous notice at least thirty days prior to the anticipated closing or (ii) mail a  
 691 notice, at least 14 *fourteen* days prior to the anticipated closing, to every current pharmacy customer  
 692 having refill authority. Each notice posted or mailed pursuant to this section shall indicate the date of  
 693 such closing, if available, and the name of the pharmacy to which prescriptions *and other required*  
 694 *prescription dispensing records and individual patient records* will be transferred unless patients indicate  
 695 their preference to the contrary. The Board of Pharmacy shall promulgate regulations providing *for a*  
 696 *definition of "closing of a pharmacy" and exceptions to this requirement the requirements of this section.*

697 B. *Upon any change of ownership of a pharmacy, regardless of how such change may be effectuated,*  
 698 *the prescription dispensing records and other patient records for at least two years immediately prior to*  
 699 *the change of ownership, shall be transferred, in accordance with Board regulations, to the new owner*  
 700 *in a manner to ensure the confidentiality, integrity, and security of the pharmacy's prescription*  
 701 *dispensing records and other patient records and the continuity of pharmacy services at substantially the*  
 702 *same level as that offered by the previous owner.*

703 *Refusing to process a request for the prescription dispensing records and other patient records*  
 704 *tendered in accordance with law or regulation shall constitute a closing and the requirements of this*  
 705 *section shall apply. Such refusal may constitute a violation of § 54.1-111 A 9, depending on the*  
 706 *circumstance.*

707 **2. That an emergency exists and this act is in force from its passage.**

708 **3. That the Board of Pharmacy shall promulgate regulations to implement this act to be effective**  
 709 **within 280 days of the enactment of this act.**