	PHARMACY SOFTWARE AMENDMENTS
	2021 GENERAL SESSION
	STATE OF UTAH
	<b>Chief Sponsor: Rosemary Lesser</b>
	Senate Sponsor: Evan J. Vickers
LON	IG TITLE
Gene	eral Description:
	This bill amends provisions relating to an electronic prescription for a controlled
subst	ance.
High	lighted Provisions:
	This bill:
	<ul> <li>requires a pharmacy software system that receives electronic prescriptions for a</li> </ul>
contr	olled substance to allow an unfilled prescription to be transferred to a different
pharr	macy; and
	<ul> <li>makes technical changes.</li> </ul>
Mon	ey Appropriated in this Bill:
	None
Othe	er Special Clauses:
	None
Utah	Code Sections Affected:
AME	ENDS:
	58-37-6, as last amended by Laws of Utah 2020, Chapter 81
ENA	CTS:
	58-37-22, Utah Code Annotated 1953

27 Be it enacted by the Legislature of the state of Utah:

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28 Section 1. Section 58-37-6 is amended to read: 29 58-37-6. License to manufacture, produce, distribute, dispense, administer, or 30 conduct research -- Issuance by division -- Denial, suspension, or revocation -- Records 31 required -- Prescriptions. 32 (1) (a) The division may adopt rules relating to the licensing and control of the manufacture, distribution, production, prescription, administration, dispensing, conducting of 33 34 research with, and performing of laboratory analysis upon controlled substances within this 35 state. 36 (b) The division may assess reasonable fees to defray the cost of issuing original and 37 renewal licenses under this chapter pursuant to Section 63J-1-504. 38 (2) (a) (i) Every person who manufactures, produces, distributes, prescribes, dispenses, 39 administers, conducts research with, or performs laboratory analysis upon any controlled 40 substance in Schedules I through V within this state, or who proposes to engage in manufacturing, producing, distributing, prescribing, dispensing, administering, conducting 41 research with, or performing laboratory analysis upon controlled substances included in 42 43 Schedules I through V within this state shall obtain a license issued by the division. 44 (ii) The division shall issue each license under this chapter in accordance with a 45 two-vear renewal cycle established by rule. The division may by rule extend or shorten a 46 renewal period by as much as one year to stagger the renewal cycles it administers. 47 (b) Persons licensed to manufacture, produce, distribute, prescribe, dispense, 48 administer, conduct research with, or perform laboratory analysis upon controlled substances in 49 Schedules I through V within this state may possess, manufacture, produce, distribute, 50 prescribe, dispense, administer, conduct research with, or perform laboratory analysis upon 51 those substances to the extent authorized by their license and in conformity with this chapter. 52 (c) The following persons are not required to obtain a license and may lawfully possess 53 controlled substances included in Schedules II through V under this section: 54 (i) an agent or employee, except a sales representative, of any registered manufacturer, 55 distributor, or dispenser of any controlled substance, if the agent or employee is acting in the 56 usual course of the agent or employee's business or employment; however, nothing in this 57 subsection shall be interpreted to permit an agent, employee, sales representative, or detail man 58 to maintain an inventory of controlled substances separate from the location of the person's

59 employer's registered and licensed place of business; 60 (ii) a motor carrier or warehouseman, or an employee of a motor carrier or 61 warehouseman, who possesses a controlled substance in the usual course of the person's 62 business or employment; and 63 (iii) an ultimate user, or a person who possesses any controlled substance pursuant to a 64 lawful order of a practitioner. 65 (d) The division may enact rules waiving the license requirement for certain 66 manufacturers, producers, distributors, prescribers, dispensers, administrators, research 67 practitioners, or laboratories performing analysis if waiving the license requirement is 68 consistent with public health and safety. 69 (e) A separate license is required at each principal place of business or professional 70 practice where the applicant manufactures, produces, distributes, dispenses, conducts research 71 with, or performs laboratory analysis upon controlled substances. (f) The division may enact rules providing for the inspection of a licensee or applicant's 72 73 establishment, and may inspect the establishment according to those rules. 74 (3) (a) (i) Upon proper application, the division shall license a qualified applicant to manufacture, produce, distribute, conduct research with, or perform laboratory analysis upon 75 76 controlled substances included in Schedules I through V, unless it determines that issuance of a 77 license is inconsistent with the public interest. 78 (ii) The division may not issue a license to any person to prescribe, dispense, or 79 administer a Schedule I controlled substance except under Subsection (3)(a)(i). 80 (iii) In determining public interest under this Subsection (3)(a), the division shall 81 consider whether the applicant has: 82 (A) maintained effective controls against diversion of controlled substances and any 83 Schedule I or II substance compounded from any controlled substance into channels other than 84 legitimate medical, scientific, or industrial channels; 85 (B) complied with applicable state and local law; (C) been convicted under federal or state laws relating to the manufacture, distribution, 86 87 or dispensing of substances; 88 (D) past experience in the manufacture of controlled dangerous substances: 89 (E) established effective controls against diversion; and

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90 (F) complied with any other factors that the division establishes that promote the public 91 health and safety. 92 (b) Licenses granted under Subsection (3)(a) do not entitle a licensee to manufacture. 93 produce, distribute, conduct research with, or perform laboratory analysis upon controlled 94 substances in Schedule I other than those specified in the license. 95 (c) (i) Practitioners shall be licensed to administer, dispense, or conduct research with 96 substances in Schedules II through V if they are authorized to administer, dispense, or conduct 97 research under the laws of this state. 98 (ii) The division need not require a separate license for practitioners engaging in 99 research with nonnarcotic controlled substances in Schedules II through V where the licensee is 100 already licensed under this chapter in another capacity. 101 (iii) With respect to research involving narcotic substances in Schedules II through V, 102 or where the division by rule requires a separate license for research of nonnarcotic substances 103 in Schedules II through V, a practitioner shall apply to the division prior to conducting 104 research. 105 (iv) Licensing for purposes of bona fide research with controlled substances by a 106 practitioner considered qualified may be denied only on a ground specified in Subsection (4), 107 or upon evidence that the applicant will abuse or unlawfully transfer or fail to safeguard 108 adequately the practitioner's supply of substances against diversion from medical or scientific 109 use. 110 (v) Practitioners registered under federal law to conduct research in Schedule I 111 substances may conduct research in Schedule I substances within this state upon providing the 112 division with evidence of federal registration. 113 (d) Compliance by manufacturers, producers, and distributors with the provisions of 114 federal law respecting registration, excluding fees, entitles them to be licensed under this 115 chapter. 116 (e) The division shall initially license those persons who own or operate an 117 establishment engaged in the manufacture, production, distribution, dispensation, or 118 administration of controlled substances prior to April 3, 1980, and who are licensed by the 119 state. 120 (4) (a) Any license issued pursuant to Subsection (2) or (3) may be denied, suspended,

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121	placed on probation, or revoked by the division upon finding that the applicant or licensee has:
122	(i) materially falsified any application filed or required pursuant to this chapter;
123	(ii) been convicted of an offense under this chapter or any law of the United States, or
124	any state, relating to any substance defined as a controlled substance;
125	(iii) been convicted of a felony under any other law of the United States or any state
126	within five years of the date of the issuance of the license;
127	(iv) had a federal registration or license denied, suspended, or revoked by competent
128	federal authority and is no longer authorized to manufacture, distribute, prescribe, or dispense
129	controlled substances;
130	(v) had the licensee's license suspended or revoked by competent authority of another
131	state for violation of laws or regulations comparable to those of this state relating to the
132	manufacture, distribution, or dispensing of controlled substances;
133	(vi) violated any division rule that reflects adversely on the licensee's reliability and
134	integrity with respect to controlled substances;
135	(vii) refused inspection of records required to be maintained under this chapter by a
136	person authorized to inspect them; or
137	(viii) prescribed, dispensed, administered, or injected an anabolic steroid for the
138	purpose of manipulating human hormonal structure so as to:
139	(A) increase muscle mass, strength, or weight without medical necessity and without a
140	written prescription by any practitioner in the course of the practitioner's professional practice;
141	or
142	(B) improve performance in any form of human exercise, sport, or game.
143	(b) The division may limit revocation or suspension of a license to a particular
144	controlled substance with respect to which grounds for revocation or suspension exist.
145	(c) (i) Proceedings to deny, revoke, or suspend a license shall be conducted pursuant to
146	this section and in accordance with the procedures set forth in Title 58, Chapter 1, Division of
147	Occupational and Professional Licensing Act, and conducted in conjunction with the
148	appropriate representative committee designated by the director of the department.
149	(ii) Nothing in this Subsection (4)(c) gives the Division of Occupational and
150	Professional Licensing exclusive authority in proceedings to deny, revoke, or suspend licenses,
151	except where the division is designated by law to perform those functions, or, when not

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152 designated by law, is designated by the executive director of the Department of Commerce to 153 conduct the proceedings. 154 (d) (i) The division may suspend any license simultaneously with the institution of 155 proceedings under this section if it finds there is an imminent danger to the public health or 156 safety. 157 (ii) Suspension shall continue in effect until the conclusion of proceedings, including 158 judicial review, unless withdrawn by the division or dissolved by a court of competent 159 jurisdiction. 160 (e) (i) If a license is suspended or revoked under this Subsection (4), all controlled substances owned or possessed by the licensee may be placed under seal in the discretion of the 161 162 division. 163 (ii) Disposition may not be made of substances under seal until the time for taking an 164 appeal has lapsed, or until all appeals have been concluded, unless a court, upon application, orders the sale of perishable substances and the proceeds deposited with the court. 165 166 (iii) If a revocation order becomes final, all controlled substances shall be forfeited. 167 (f) The division shall notify promptly the Drug Enforcement Administration of all 168 orders suspending or revoking a license and all forfeitures of controlled substances. 169 (g) If an individual's Drug Enforcement Administration registration is denied, revoked, 170 surrendered, or suspended, the division shall immediately suspend the individual's controlled 171 substance license, which shall only be reinstated by the division upon reinstatement of the 172 federal registration, unless the division has taken further administrative action under 173 Subsection (4)(a)(iv), which would be grounds for the continued denial of the controlled 174 substance license. 175 (5) (a) A person licensed under Subsection (2) or (3) shall maintain records and 176 inventories in conformance with the record keeping and inventory requirements of federal and 177 state law and any additional rules issued by the division. 178 (b) (i) A physician, dentist, naturopathic physician, veterinarian, practitioner, or other 179 individual who is authorized to administer or professionally use a controlled substance shall 180 keep a record of the drugs received by the individual and a record of all drugs administered, 181 dispensed, or professionally used by the individual otherwise than by a prescription. 182 (ii) An individual using small quantities or solutions or other preparations of those

183	drugs for local application has complied with this Subsection (5)(b) if the individual keeps a
184	record of the quantity, character, and potency of those solutions or preparations purchased or
185	prepared by the individual, and of the dates when purchased or prepared.
186	(6) Controlled substances in Schedules I through V may be distributed only by a
187	licensee and pursuant to an order form prepared in compliance with division rules or a lawful
188	order under the rules and regulations of the United States.
189	(7) (a) An individual may not write or authorize a prescription for a controlled
190	substance unless the individual is:
191	(i) a practitioner authorized to prescribe drugs and medicine under the laws of this state
192	or under the laws of another state having similar standards; and
193	(ii) licensed under this chapter or under the laws of another state having similar
194	standards.
195	(b) An individual other than a pharmacist licensed under the laws of this state, or the
196	pharmacist's licensed intern, as required by Sections 58-17b-303 and 58-17b-304, may not
197	dispense a controlled substance.
198	(c) (i) A controlled substance may not be dispensed without the written prescription of
199	a practitioner, if the written prescription is required by the federal Controlled Substances Act.
200	(ii) That written prescription shall be made in accordance with Subsection (7)(a) and in
201	conformity with Subsection (7)(d).
202	(iii) In emergency situations, as defined by division rule, controlled substances may be
203	dispensed upon oral prescription of a practitioner, if reduced promptly to writing on forms
204	designated by the division and filed by the pharmacy.
205	(iv) Prescriptions reduced to writing by a pharmacist shall be in conformity with
206	Subsection (7)(d).
207	(d) Except for emergency situations designated by the division, an individual may not
208	issue, fill, compound, or dispense a prescription for a controlled substance unless the
209	prescription is signed by the prescriber in ink or indelible pencil or is signed with an electronic
210	signature of the prescriber as authorized by division rule, and contains the following
211	information:
212	(i) the name, address, and registry number of the prescriber;
213	(ii) the name, address, and age of the person to whom or for whom the prescription is

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214	issued;
215	(iii) the date of issuance of the prescription; and
216	(iv) the name, quantity, and specific directions for use by the ultimate user of the
217	controlled substance.
218	(e) A prescription may not be written, issued, filled, or dispensed for a Schedule I
219	controlled substance unless:
220	(i) the individual who writes the prescription is licensed under Subsection (2); and
221	(ii) the prescribed controlled substance is to be used in research.
222	(f) Except when administered directly to an ultimate user by a licensed practitioner,
223	controlled substances are subject to the restrictions of this Subsection (7)(f).
224	(i) A prescription for a Schedule II substance may not be refilled.
225	(ii) A Schedule II controlled substance may not be filled in a quantity to exceed a
226	one-month's supply, as directed on the daily dosage rate of the prescriptions.
227	(iii) (A) Except as provided in Subsection (7)(f)(iii)(B), a prescription for a Schedule II
228	or Schedule III controlled substance that is an opiate and that is issued for an acute condition
229	shall be completely or partially filled in a quantity not to exceed a seven-day supply as directed
230	on the daily dosage rate of the prescription.
231	(B) Subsection (7)(f)(iii)(A) does not apply to a prescription issued for a surgery when
232	the practitioner determined that a quantity exceeding seven days is needed, in which case the
233	practitioner may prescribe up to a 30-day supply, with a partial fill at the discretion of the
234	practitioner.
235	(C) Subsection $(7)(f)(iii)(A)$ does not apply to prescriptions issued for complex or
236	chronic conditions which are documented as being complex or chronic in the medical record.
237	(D) A pharmacist is not required to verify that a prescription is in compliance with
238	Subsection (7)(f)(iii).
239	(iv) A Schedule III or IV controlled substance may be filled only within six months of
240	issuance, and may not be refilled more than six months after the date of its original issuance or
241	be refilled more than five times after the date of the prescription unless renewed by the
242	practitioner.
243	(v) All other controlled substances in Schedule V may be refilled as the prescriber's
244	prescription directs, but they may not be refilled one year after the date the prescription was

245	issued unless renewed by the practitioner.
246	(vi) Any prescription for a Schedule II substance may not be dispensed if it is not
247	presented to a pharmacist for dispensing by a pharmacist or a pharmacy intern within 30 days
248	after the date the prescription was issued, or 30 days after the dispensing date, if that date is
249	specified separately from the date of issue.
250	(vii) A practitioner may issue more than one prescription at the same time for the same
251	Schedule II controlled substance, but only under the following conditions:
252	(A) no more than three prescriptions for the same Schedule II controlled substance may
253	be issued at the same time;
254	(B) no one prescription may exceed a 30-day supply; and
255	(C) a second or third prescription shall include the date of issuance and the date for
256	dispensing.
257	[(g) (i) Beginning January 1, 2022, each prescription issued for a controlled substance
258	shall be transmitted electronically as an electronic prescription unless the prescription is:]
259	[(A) for a patient residing in an assisted living facility as that term is defined in Section
260	26-21-2, a long-term care facility as that term is defined in Section 58-31b-102, or a
261	correctional facility as that term is defined in Section 64-13-1;]
262	[(B) issued by a veterinarian licensed under Title 58, Chapter 28, Veterinary Practice
263	Act;]
264	[(C) dispensed by a Department of Veterans Affairs pharmacy;]
265	[(D) issued during a temporary technical or electronic failure at the practitioner's or
266	pharmacy's location; or]
267	[(E) issued in an emergency situation.]
268	[(ii) The division, in collaboration with the appropriate boards that govern the licensure
269	of the licensees who are authorized by the division to prescribe or to dispense controlled
270	substances, shall make rules in accordance with Title 63G, Chapter 3, Utah Administrative
271	Rulemaking Act to:]
272	[(A) require that controlled substances prescribed or dispensed under Subsection
273	(7)(g)(i)(D) indicate on the prescription that the prescribing practitioner or the pharamacy is
274	experiencing a technical difficulty or an electronic failure;]
275	[(B) define an emergency situation for purposes of Subsection (7)(g)(i)(E);]

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276 (C) establish additional exemptions to the electronic prescription requirements 277 established in this Subsection (7)(g);] 278 (D) establish guidelines under which a prescribing practitioner or a pharmacy may 279 obtain an extension of up to two additional years to comply with Subsection (7)(g)(i);] 280 (E) establish a protocol to follow if the pharmacy that receives the electronic 281 prescription is not able to fill the prescription; and] 282 [(F) establish requirements that comply with federal laws and regulations for software 283 used to issue and dispense electronic prescriptions.] 284 [(h)] (g) An order for a controlled substance in Schedules II through V for use by an 285 inpatient or an outpatient of a licensed hospital is exempt from all requirements of this 286 Subsection (7) if the order is: 287 (i) issued or made by a prescribing practitioner who holds an unrestricted registration 288 with the federal Drug Enforcement Administration, and an active Utah controlled substance 289 license in good standing issued by the division under this section, or a medical resident who is 290 exempted from licensure under Subsection 58-1-307(1)(c); 291 (ii) authorized by the prescribing practitioner treating the patient and the prescribing 292 practitioner designates the quantity ordered; 293 (iii) entered upon the record of the patient, the record is signed by the prescriber 294 affirming the prescriber's authorization of the order within 48 hours after filling or 295 administering the order, and the patient's record reflects the quantity actually administered; and 296 (iv) filled and dispensed by a pharmacist practicing the pharmacist's profession within 297 the physical structure of the hospital, or the order is taken from a supply lawfully maintained by 298 the hospital and the amount taken from the supply is administered directly to the patient 299 authorized to receive it. 300 (ii) (h) A practitioner licensed under this chapter may not prescribe, administer, or 301 dispense a controlled substance to a child, without first obtaining the consent required in 302 Section 78B-3-406 of a parent, guardian, or person standing in loco parentis of the child except in cases of an emergency. For purposes of Subsection (7)[(i)](h), "child" has the same meaning 303 as defined in Section 78A-6-105, and "emergency" means any physical condition requiring the 304 305 administration of a controlled substance for immediate relief of pain or suffering. 306 (i) A practitioner licensed under this chapter may not prescribe or administer

dosages of a controlled substance in excess of medically recognized quantities necessary totreat the ailment, malady, or condition of the ultimate user.

309 [(k)] (j) A practitioner licensed under this chapter may not prescribe, administer, or 310 dispense any controlled substance to another person knowing that the other person is using a 311 false name, address, or other personal information for the purpose of securing the controlled 312 substance.

313 [(1)] (k) A person who is licensed under this chapter to manufacture, distribute, or
 314 dispense a controlled substance may not manufacture, distribute, or dispense a controlled
 315 substance to another licensee or any other authorized person not authorized by this license.

316 [(m)] (l) A person licensed under this chapter may not omit, remove, alter, or obliterate
317 a symbol required by this chapter or by a rule issued under this chapter.

318 [(n)] (m) A person licensed under this chapter may not refuse or fail to make, keep, or
 319 furnish any record notification, order form, statement, invoice, or information required under
 320 this chapter.

321 [(o)] (n) A person licensed under this chapter may not refuse entry into any premises
 322 for inspection as authorized by this chapter.

323 [(p)] (o) A person licensed under this chapter may not furnish false or fraudulent
 324 material information in any application, report, or other document required to be kept by this
 325 chapter or willfully make any false statement in any prescription, order, report, or record
 326 required by this chapter.

(8) (a) (i) Any person licensed under this chapter who is found by the division to have
violated any of the provisions of Subsections (7)(k) through (o) or Subsection (10) is subject to
a penalty not to exceed \$5,000. The division shall determine the procedure for adjudication of
any violations in accordance with Sections 58-1-106 and 58-1-108.

(ii) The division shall deposit all penalties collected under Subsection (8)(a)(i) [in] into
the General Fund as a dedicated credit to be used by the division under Subsection
58-37f-502(1).

334 (iii) The director may collect a penalty that is not paid by:

- 335 (A) referring the matter to a collection agency; or
- (B) bringing an action in the district court of the county where the person against

337 whom the penalty is imposed resides or in the county where the office of the director is located.

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338	(iv) A county attorney or the attorney general of the state shall provide legal assistance
339	and advice to the director in an action to collect a penalty.
340	(v) A court shall award reasonable attorney fees and costs to the prevailing party in an
341	action brought by the division to collect a penalty.
342	(b) Any person who knowingly and intentionally violates Subsections (7)(h) through (j)
343	or Subsection (10) is:
344	(i) upon first conviction, guilty of a class B misdemeanor;
345	(ii) upon second conviction, guilty of a class A misdemeanor; and
346	(iii) on third or subsequent conviction, guilty of a third degree felony.
347	(c) Any person who knowingly and intentionally violates Subsections (7)(k) through
348	(o) shall upon conviction be guilty of a third degree felony.
349	(9) Any information communicated to any licensed practitioner in an attempt to
350	unlawfully procure, or to procure the administration of, a controlled substance is not considered
351	to be a privileged communication.
352	(10) A person holding a valid license under this chapter who is engaged in medical
353	research may produce, possess, administer, prescribe, or dispense a controlled substance for
354	research purposes as licensed under Subsection (2) but may not otherwise prescribe or dispense
355	a controlled substance listed in Section 58-37-4.2.
356	Section 2. Section <b>58-37-22</b> is enacted to read:
357	58-37-22. Electronic prescriptions for controlled substances.
358	(1) Beginning January 1, 2022, each prescription issued for a controlled substance shall
359	be transmitted electronically as an electronic prescription unless the prescription is:
360	(a) for a patient residing in an assisted living facility as that term is defined in Section
361	26-21-2, a long-term care facility as that term is defined in Section 58-31b-102, or a
362	correctional facility as that term is defined in Section 64-13-1;
363	(b) issued by a veterinarian licensed under Title 58, Chapter 28, Veterinary Practice
364	<u>Act;</u>
365	(c) dispensed by a Department of Veterans Affairs pharmacy;
366	(d) issued during a temporary technical or electronic failure at the practitioner's or
367	pharmacy's location; or
368	(e) issued in an emergency situation.

369	(2) The division, in collaboration with the appropriate boards that govern the licensure
370	of the licensees who are authorized by the division to prescribe or to dispense controlled
371	substances, shall make rules in accordance with Title 63G, Chapter 3, Utah Administrative
372	Rulemaking Act to:
373	(a) require that controlled substances prescribed or dispensed under Subsection (1)(d)
374	indicate on the prescription that the prescribing practitioner or the pharmacy is experiencing a
375	technical difficulty or an electronic failure;
376	(b) define an emergency situation for purposes of Subsection (1)(e);
377	(c) establish additional exemptions to the electronic prescription requirements
378	established in this section;
379	(d) establish guidelines under which a prescribing practitioner or a pharmacy may
380	obtain an extension of up to two additional years to comply with Subsection (1);
381	(e) establish a protocol to follow if the pharmacy that receives the electronic
382	prescription is not able to fill the prescription; and
383	(f) establish requirements that comply with federal laws and regulations for software
384	used to issue and dispense electronic prescriptions.
385	(3) Beginning July 1, 2024, a pharmacy software program for receiving an electronic
386	prescription for a controlled substance shall be capable of electronically transferring a
387	prescription to a different pharmacy Ĥ→ : ←Ĥ
387a	$\hat{H} \rightarrow (\underline{a}) \leftarrow \hat{H}$ upon the request of the patient or the practitioner $\hat{H} \rightarrow \underline{;}$
387b	(b) with the approval of a pharmacist at the originating pharmacy; and
387c	(c) ← $\hat{H}$ if the
388	prescription is unfilled.