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Evan J. Vickers proposes the following substitute bill:

Cannabinoid Amendments

2025 GENERAL SESSION STATE OF UTAH

Chief Sponsor: Jennifer Dailey-Provost

Senate Sponsor: Evan J. Vickers

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3	LONG TITLE
4	General Description:

This bill amends provisions related to hemp and medical cannabis regulation.

Highlighted Provisions:

This bill:

- defines terms;
- prohibits certain cannabinoids from being used in cannabinoid products;
- - amends background check requirements for cannabinoid processor licenses;
 - amends qualifications for obtaining a cannabinoid processor license;
 - requires industrial hemp retailers to maintain a video surveillance system;
- ▶ amends provisions related to cannabinoid product enforcement;
 - requires a person to have a cannabis processor license to transport hemp concentrate;
- 17 removes the requirement that certain cannabinoid products be in a medicinal dosage form;
 - allows for additional medical cannabis pharmacies;
- reates a new medical cannabis pharmacy license for independent medical cannabis
- 20 pharmacies;
- creates ownership restrictions for independent medical cannabis pharmacies;
- 22 adjusts fees for certain medical cannabis pharmacy licenses;
- ≥ amends provisions regarding cannabis production and sanitation;
 - modifies provisions related to enforcement and appeals;
- 25 amends provisions related to closed-door medical cannabis pharmacies;
- ≥ allows a cannabis processing facility to have a website that includes product information;
- 27 amends provisions regarding when the department may seize products and test products;
- 28 amends provisions related to information a medical cannabis pharmacy must have

29	available to a patient purchasing medical cannabis;
30	 creates a reporting requirement for the department;
31	 repeals sections related to the state central patient portal; and
32	 makes technical and conforming changes.
33	Money Appropriated in this Bill:
34	None
35	Other Special Clauses:
36	None
37	Utah Code Sections Affected:
38	AMENDS:
39	4-41-102, as last amended by Laws of Utah 2024, Chapter 35
40	4-41-103.2 , as last amended by Laws of Utah 2023, Chapter 146
41	4-41-103.3, as last amended by Laws of Utah 2023, Chapters 146, 327
42	4-41-105, as last amended by Laws of Utah 2024, Chapter 35
43	4-41-404, as last amended by Laws of Utah 2019, Chapter 23
44	4-41a-102, as last amended by Laws of Utah 2024, Chapters 217, 238 and 240
45	4-41a-403, as last amended by Laws of Utah 2023, Chapter 327
46	4-41a-501, as last amended by Laws of Utah 2023, Chapter 313
47	4-41a-701, as last amended by Laws of Utah 2023, Chapters 313, 317
48	4-41a-801, as renumbered and amended by Laws of Utah 2018, Third Special Session,
49	Chapter 1
50	4-41a-802, as last amended by Laws of Utah 2024, Chapter 217
51	4-41a-1001, as last amended by Laws of Utah 2024, Chapters 217, 238 and 240
52	4-41a-1003, as last amended by Laws of Utah 2023, Chapter 435 and renumbered and
53	amended by Laws of Utah 2023, Chapters 273, 307 and last amended by Coordination Clause,
54	Laws of Utah 2023, Chapter 307
55	4-41a-1005, as last amended by Laws of Utah 2024, Chapter 217
56	4-41a-1101, as last amended by Laws of Utah 2024, Chapter 217
57	4-41a-1201, as enacted by Laws of Utah 2023, Chapter 273
58	4-41a-1202, as last amended by Laws of Utah 2024, Chapters 217, 240
59	4-41a-1203, as renumbered and amended by Laws of Utah 2023, Chapters 273, 307 and
60	last amended by Coordination Clause, Laws of Utah 2023, Chapter 307
61	4-41a-1206, as enacted by Laws of Utah 2024, Chapter 238
62	26B-1-435 , as last amended by Laws of Utah 2024, Chapters 238, 240

63 **26B-4-201**, as last amended by Laws of Utah 2024, Chapters 217, 240 64 **26B-4-202**, as last amended by Laws of Utah 2024, Chapters 217, 240 65 26B-4-214, as last amended by Laws of Utah 2024, Chapter 240 66 **26B-4-222**, as last amended by Laws of Utah 2024, Chapter 240 67 **58-37-3.6**, as last amended by Laws of Utah 2024, Chapter 35 68 **58-85-102**, as last amended by Laws of Utah 2018, Third Special Session, Chapter 1 69 **63N-3-1301**, as enacted by Laws of Utah 2024, Chapter 35 70 **77-39-101**, as last amended by Laws of Utah 2024, Chapter 35 71 **ENACTS:** 72 **4-41-405**, Utah Code Annotated 1953 73 **4-41a-1006.** Utah Code Annotated 1953 74 REPEALS: 75 26B-4-236, as last amended by Laws of Utah 2023, Chapters 273, 317 and renumbered 76 and amended by Laws of Utah 2023, Chapter 307 and last amended by Coordination Clause, 77 Laws of Utah 2023, Chapter 307 78 79 *Be it enacted by the Legislature of the state of Utah:* 80 Section 1. Section **4-41-102** is amended to read: 81 4-41-102. Definitions. 82 As used in this chapter: 83 (1) "Adulterant" means any poisonous or deleterious substance in a quantity that may be 84 injurious to human health, including: 85 (a) pesticides; 86 (b) heavy metals; 87 (c) solvents: 88 (d) microbial life; 89 (e) artificially derived cannabinoids; 90 (f) toxins; or 91 (g) foreign matter. 92 (2)(a) "Artificially derived cannabinoid" means a chemical substance that is created by a 93 chemical reaction that changes the molecular structure of any chemical substances 94 derived from the cannabis plant. 95 (b) "Artificially derived cannabinoid" does not include:

(i) a naturally occurring chemical substance that is separated from the cannabis plant

97	by a chemical or mechanical extraction process; or
98	(ii) cannabinoids that are produced by decarboxylation from a naturally occurring
99	cannabinoid acid without the use of a chemical catalyst.
100	(3) "Cannabidiol" or "CBD" means the cannabinoid identified as CAS# 13956-29-1.
101	(4) "Cannabidiolic acid" or "CBDA" means the cannabinoid identified as CAS# 1244-58-2.
102	(5) "Cannabinoid processor license" means a license that the department issues to a person
103	for the purpose of processing a cannabinoid product.
104	(6) "Cannabinoid product" means a product that:
105	(a) contains or is represented to contain one or more naturally occurring cannabinoids;
106	(b) contains less than the cannabinoid product THC level, by dry weight;
107	(c) contains a combined amount of total THC and any THC analog that does not exceed
108	10% of the total cannabinoid content;
109	(d) does not exceed a total of THC and any THC analog that is greater than:
110	(i) 5 milligrams per serving; and
111	(ii) 150 milligrams per package; and
112	(e) unless the product is in an oil based suspension, has a serving size that:
113	(i) is an integer; and
114	(ii) is a discrete unit of the cannabinoid product.
115	(7) "Cannabinoid product class" means a group of cannabinoid products that:
116	(a) have all ingredients in common; and
117	(b) are produced by or for the same company.
118	(8) "Cannabinoid product THC level" means a combined concentration of total THC and
119	any THC analog of less than 0.3% on a dry weight basis if laboratory testing confirms a
120	result within a measurement of uncertainty that includes the combined concentration of
121	0.3%.
122	(9) "Cannabis" means the same as that term is defined in Section 26B-4-201.
123	(10) "Delta-9-tetrahydrocannabinol" or "delta-9-THC" means the cannabinoid identified as
124	CAS# 1972-08-3, the primary psychotropic cannabinoid in cannabis.
125	(11) "Industrial hemp" means any part of a cannabis plant, whether growing or not, with a
126	concentration of less than 0.3% tetrahydrocannabinol by dry weight.
127	(12) "Industrial hemp producer registration" means a registration that the department issues
128	to a person for the purpose of processing industrial hemp or an industrial hemp product.
129	(13)(a) "Industrial hemp product" means a product made by processing industrial hemp
130	plants or industrial hemp parts.

131	(b) "Industrial hemp product" does not include cannabinoid material or a cannabinoid
132	product.
133	[(13)] (14) "Industrial hemp retailer permit" means a permit that the department issues to a
134	retailer who sells any viable industrial hemp seed or cannabinoid product.
135	[(14)(a) "Industrial hemp product" means a product made by processing industrial hemp
136	plants or industrial hemp parts.]
137	[(b) "Industrial hemp product" does not include cannabinoid material.]
138	(15) "Key participant" means any of the following:
139	(a) a licensee;
140	(b) an operation manager;
141	(c) a site manager; or
142	(d) an employee who has access to any industrial hemp material with a THC
143	concentration above 0.3%.
144	(16) "Licensee" means a person possessing a cannabinoid processor license that the
145	department issues under this chapter.
146	(17) "Newly identified cannabinoid" means a cannabinoid that:
147	(a) is not expressly identified by chemical name or CAS number in this chapter; and
148	(b) is identified by the department under Section 4-41-405.
149	[(17)] (18) "Non-compliant material" means:
150	(a) a hemp plant that does not comply with this chapter, including a cannabis plant with
151	a concentration of 0.3% tetrahydrocannabinol or greater by dry weight;[-and]
152	(b) a cannabinoid product, chemical, or compound with a concentration that exceeds the
153	cannabinoid product THC level[-] ; and
154	(c) a cannabinoid product containing any of the following:
155	(i) delta-9-tetrahydrocannabiphorol (THCP), the cannabinoid identified as CAS#
156	<u>54763-99-4;</u>
157	(ii) delta-8-tetrahydrocannabiphorol (THCP), the cannabinoid identified as CAS#
158	<u>51768-60-6;</u>
159	(iii) delta-9-tetrahyrdocannabinol (THC) acetate, the cannabinoid identified as CAS#
160	23132-17-4;
161	(iv) delta-8-tetrahydrocannabinol (THC) acetate, the cannabinoid identified as CAS#
162	23050-54-6;
163	(v) 9(s)-hexahydrocannabinol (HHC), the cannabinoid identified as CAS#
164	36403-91-5; or

165	(vi) 9(r)-hexahyrdocannabinol (HHC), the cannabinoid identified as CAS#
166	<u>36403-90-4.</u>
167	[(18)] (19) "Permittee" means a person possessing a permit that the department issues under
168	this chapter.
169	[(19)] <u>(20)</u> "Person" means:
170	(a) an individual, partnership, association, firm, trust, limited liability company, or
171	corporation; and
172	(b) an agent or employee of an individual, partnership, association, firm, trust, limited
173	liability company, or corporation.
174	[(20)] (21) "Retailer permittee" means a person possessing an industrial hemp retailer permit
175	that the department issues under this chapter.
176	[(21)] (22) "Tetrahydrocannabinol" or "THC" means a delta-9-tetrahydrocannabinol, the
177	cannabinoid identified as CAS# 1972-08-3.
178	[(22)] (23)(a) "THC analog" means a substance that is structurally or pharmacologically
179	substantially similar to, or is represented as being similar to, delta-9-THC.
180	(b) "THC analog" does not include the following substances or the naturally occurring
181	acid forms of the following substances:
182	(i) cannabichromene (CBC), the cannabinoid identified as CAS# 20675-51-8;
183	(ii) cannabicyclol (CBL), the cannabinoid identified as CAS# 21366-63-2;
184	(iii) cannabidiol (CBD), the cannabinoid identified as CAS# 13956-29-1;
185	(iv) cannabidivarol (CBDV), the cannabinoid identified as CAS# 24274-48-4;
186	(v) cannabielsoin (CBE), the cannabinoid identified as CAS# 52025-76-0;
187	(vi) cannabigerol (CBG), the cannabinoid identified as CAS# 25654-31-3;
188	(vii) cannabigerovarin (CBGV), the cannabinoid identified as CAS# 55824-11-8;
189	(viii) cannabinol (CBN), the cannabinoid identified as CAS# 521-35-7;
190	(ix) cannabivarin (CBV), the cannabinoid identified as CAS# 33745-21-0; or
191	(x) delta-9-tetrahydrocannabivarin (THCV), the cannabinoid identified as CAS#
192	31262-37-0.
193	[(23)] (24) "Total cannabidiol" or "total CBD" means the combined amounts of cannabidiol
194	and cannabidiolic acid, calculated as "total CBD = CBD + (CBDA x 0.877)".
195	[(24)] (25) "Total tetrahydrocannabinol" or "total THC" means the sum of the determined
196	amounts of delta-9-THC, tertrahydrocannabinolic acid, calculated as "total THC =
197	delta-9-THC + (THCA x 0.877)".
198	[(25)] (26) "Transportable industrial hemp concentrate" means any amount of a natural

199	cannabinoid in a purified state that:
200	(a) is the product of any chemical or physical process applied to naturally occurring
201	biomass that concentrates or isolates the cannabinoids contained in the biomass;
202	(b) is derived from a cannabis plant that, based on sampling that was collected no more
203	than 30 days before the day on which the cannabis plant was harvested, contains a
204	combined concentration of total THC and any THC analog of less than 0.3% on a dry
205	weight basis;
206	(c) has a THC and THC analog concentration total that is less than 20% when
207	concentrated from the cannabis plant to the purified state; and
208	(d) is intended to be processed into a cannabinoid product.
209	Section 2. Section 4-41-103.2 is amended to read:
210	4-41-103.2 . Cannabinoid processor license.
211	(1) The department or a licensee of the department may process a cannabinoid product.
212	(2) A person seeking a cannabinoid processor license shall provide to the department:
213	(a) the legal description and global positioning coordinates sufficient for locating the
214	facility the person uses to process industrial hemp; and
215	(b) written consent allowing a representative of the department and local law
216	enforcement to enter all premises where the person processes or stores industrial
217	hemp for the purpose of:
218	(i) conducting a physical inspection; or
219	(ii) ensuring compliance with the requirements of this chapter.
220	[(3) An individual who has been convicted of a drug-related felony within the last 10 years
221	is not eligible to obtain a cannabinoid processor license.]
222	[(4)] (3) The department may set a fee in accordance with Subsection 4-2-103(2) for the
223	application for a cannabinoid processor license.
224	[(5)] (4) A licensee may only market a cannabinoid product that the licensee processes.
225	(5)(a) An applicant for a cannabis processor license shall:
226	(i) be at least 18 years old; and
227	(ii) submit a nationwide criminal history from the Federal Bureau of Investigation to
228	the department.
229	(b) The department shall reject an individual's application for a cannabis processor
230	license if the criminal history described in Subsection (5)(a)(ii) was not completed in
231	the previous 90 days before the day the applicant submits the license application to
232	the department.

233	(6) An applicant is not eligible to receive a cannabis processor license if the applicant has:
234	(a) been convicted of a felony; or
235	(b) been convicted of a drug-related misdemeanor within the previous ten years.
236	[(6)(a) Each applicant for a license to process cannabinoid products shall submit to the
237	department, at the time of application, from each key participant:]
238	[(i) a fingerprint card in a form acceptable to the Department of Public Safety;]
239	[(ii) a signed waiver in accordance with Subsection 53-10-108(4) acknowledging the
240	registration of the individual's fingerprints in the Federal Bureau of Investigation
241	Next Generation Identification System's Rap Back Service; and]
242	[(iii) consent to a fingerprint background check by:]
243	[(A) the Bureau of Criminal Identification; and]
244	[(B) the Federal Bureau of Investigation.]
245	[(b) The Bureau of Criminal Identification shall:]
246	[(i) check the fingerprints the applicant submits under Subsection (6)(a) against the
247	applicable state, regional, and national criminal records databases, including the
248	Federal Bureau of Investigation Next Generation Identification System;]
249	[(ii) report the results of the background check to the department;]
250	[(iii) maintain a separate file of fingerprints that applicants submit under Subsection
251	(6)(a) for search by future submissions to the local and regional criminal records
252	databases, including latent prints;]
253	[(iv) request that the fingerprints be retained in the Federal Bureau of Investigation
254	Next Generation Identification System's Rap Back Service for search by future
255	submissions to national criminal records databases, including the Next Generation
256	Identification System and latent prints; and]
257	[(v) establish a privacy risk mitigation strategy to ensure that the department only
258	receives notifications for an individual with whom the department maintains an
259	authorizing relationship.]
260	[(c) The department shall:]
261	[(i) assess an individual who submits fingerprints under Subsection (6)(a) a fee in an
262	amount that the department sets in accordance with Section 63J-1-504 for the
263	services that the Bureau of Criminal Identification or another authorized agency
264	provides under this section; and]
265	[(ii) remit the fee described in Subsection (6)(c)(i) to the Bureau of Criminal
266	Identification.]

267	Section 3. Section 4-41-103.3 is amended to read:
268	4-41-103.3 . Industrial hemp retailer permit.
269	(1) Except as provided in Subsection [(4)] (5), a retailer permittee of the department may
270	market or sell a cannabinoid product or a viable industrial hemp seed.
271	(2) A person seeking an industrial hemp retailer permit shall provide to the department:
272	(a) the name of the person that is seeking to market or sell a cannabinoid product or a
273	viable industrial hemp seed;
274	(b) the address of each location where a cannabinoid product or a viable industrial hemp
275	seed will be sold; and
276	(c) written consent allowing a representative of the department to enter all premises
277	where the person is selling a cannabinoid product or a viable industrial hemp seed for
278	the purpose of:
279	(i) conducting a physical inspection; or
280	(ii) ensuring compliance with the requirements of this chapter.
281	(3) Beginning January 1, 2026, an industrial hemp retailer permittee shall:
282	(a) maintain a video surveillance system that:
283	(i) is able to monitor who purchases a cannabinoid product from the permittee;
284	(ii) is tamper proof; and
285	(iii) stores a video record for at least 45 days; and
286	(b) provide the department access to the video surveillance system upon request.
287	[(3)] (4) The department may set a fee in accordance with Subsection 4-2-103(2) for the
288	application for an industrial hemp retailer permit.
289	[(4)] (5) Any marketing for a cannabinoid product or a viable industrial hemp seed shall
290	include a notice to consumers that the product is hemp and is not cannabis or medical
291	cannabis, as those terms are defined in Section 26B-4-201.
292	Section 4. Section 4-41-105 is amended to read:
293	4-41-105 . Unlawful acts.
294	(1) It is unlawful for a person to handle, process, or market living industrial hemp plants,
295	viable hemp seeds, leaf materials, or floral materials derived from industrial hemp
296	without the appropriate license or permit issued by the department under this chapter.
297	(2)(a) It is unlawful for any person to:
298	(i) distribute, sell, or market a cannabinoid product that is:
299	(A) not registered with the department under Section 4-41-104; or
300	(B) noncompliant material:

301	(ii) except as provided in Subsection (2)(b), transport into or out of the state extracted
302	material or final product that contains 0.3% or more of total THC and any THC
303	analog;
304	(iii) sell or use a cannabinoid product that is:
305	(A) added to a conventional food or beverage, as the department further defines in
306	rules described in Section 4-41-403;
307	(B) marketed or manufactured to be enticing to children, as further defined in
308	rules described in Section 4-41-403; or
309	(C) smokable flower; or
310	(iv) knowingly or intentionally sell or give a cannabinoid product that contains THC
311	or a THC analog in the course of business to an individual who is not at least 21
312	years old.
313	(b) A person may transport transportable industrial hemp concentrate if the person:
314	(i) complies with rules created by the department under Section 4-41-103.1 related to
315	transportable industrial hemp concentrate; and
316	(ii)(A) has [an industrial hemp producer registration] a cannabinoid processor
317	<u>license</u> ; or
318	(B) the equivalent to [an industrial hemp producer registration] a cannabinoid
319	<u>processor license</u> from another state.
320	(3) The department may seize and destroy non-compliant material.
321	(4) Nothing in this chapter authorizes any person to violate federal law, regulation, or any
322	provision of this title.
323	Section 5. Section 4-41-404 is amended to read:
324	4-41-404 . Department duties.
325	The department [shall assess the fine described in Subsection 4-41-403(4)] may take an
326	enforcement action in accordance with Section 4-41-106 against any person who offers an
327	unregistered cannabinoid product for sale in this state.
328	Section 6. Section 4-41-405 is enacted to read:
329	4-41-405 . Newly identified cannabinoid.
330	(1) For a newly identified cannabinoid, the department may:
331	(a) establish a maximum allowable concentration that a cannabinoid product may
332	contain of the newly identified cannabinoid;
333	(b) prohibit the newly identified cannabinoid from appearing in a cannabinoid product;
334	or

335	(c) modify the maximum allowable concentration described in Subsection (1)(a) as
336	necessary if it would not create a threat to public health.
337	(2) In accordance with Title 63G, Chapter 3, Utah Administrative Rulemaking Act, the
338	department shall make rules to implement Subsection (1).
339	Section 7. Section 4-41a-102 is amended to read:
340	4-41a-102 . Definitions.
341	As used in this chapter:
342	(1) "Adulterant" means any poisonous or deleterious substance in a quantity that may be
343	injurious to health, including:
344	(a) pesticides;
345	(b) heavy metals;
346	(c) solvents;
347	(d) microbial life;
348	(e) artificially derived cannabinoid;
349	(f) toxins; or
350	(g) foreign matter.
351	(2) "Advertise" or "advertising" means information provided by a person in any medium:
352	(a) to the public; and
353	(b) that is not age restricted to an individual who is at least 21 years old.
354	(3) "Advisory board" means the Medical Cannabis Policy Advisory Board created in
355	Section 26B-1-435.
356	(4)(a) "Anticompetitive business practice" means any practice that reduces the amount
357	of competition in the medical cannabis market that would be considered an attempt to
358	monopolize, as defined in Section 76-10-3103.
359	(b) "Anticompetitive business practice" may include:
360	(i) agreements that may be considered unreasonable when competitors interact to the
361	extent that they are:
362	(A) no longer acting independently; or
363	(B) when collaborating are able to wield market power together;
364	(ii) monopolizing or attempting to monopolize trade by:
365	(A) acting to maintain or acquire a dominant position in the market; or
366	(B) preventing new entry into the market; or
367	(iii) other conduct outlined in rule.
368	(5)(a) "Artificially derived cannabinoid" means a chemical substance that is created by a

369	chemical reaction that changes the molecular structure of any chemical substance
370	derived from the cannabis plant.
371	(b) "Artificially derived cannabinoid" does not include:
372	(i) a naturally occurring chemical substance that is separated from the cannabis plant
373	by a chemical or mechanical extraction process; or
374	(ii) a cannabinoid that is produced by decarboxylation from a naturally occurring
375	cannabinoid acid without the use of a chemical catalyst.
376	(6) "Batch" means a quantity of:
377	(a) cannabis extract produced on a particular date and time and produced between
378	completion of equipment and facility sanitation protocols until the next required
379	sanitation cycle during which lots of cannabis are used;
380	(b) cannabis product produced on a particular date and time and produced between
381	completion of equipment and facility sanitation protocols until the next required
382	sanitation cycle during which cannabis extract is used; or
383	(c) cannabis flower packaged on a particular date and time and produced between
384	completion of equipment and facility sanitation protocols until the next required
385	sanitation cycle during which lots of cannabis are being used.
386	[(6)] (7) "Cannabis Research Review Board" means the Cannabis Research Review Board
387	created in Section 26B-1-420.
388	[(7)] (8) "Cannabis" means the same as that term is defined in Section 26B-4-201.
389	[(8)] <u>(9)</u> "Cannabis concentrate" means:
390	(a) the product of any chemical or physical process applied to naturally occurring
391	biomass that concentrates or isolates the cannabinoids contained in the biomass; and
392	(b) any amount of a natural cannabinoid or artificially derived cannabinoid in an
393	artificially derived cannabinoid's purified state.
394	[(9)] (10) "Cannabis cultivation byproduct" means any portion of a cannabis plant that is not
395	intended to be sold as a cannabis plant product.
396	[(10)] (11) "Cannabis cultivation facility" means a person that:
397	(a) possesses cannabis;
398	(b) grows or intends to grow cannabis; and
399	(c) sells or intends to sell cannabis to a cannabis cultivation facility, a cannabis
400	processing facility, or a medical cannabis research licensee.
401	[(11)] (12) "Cannabis cultivation facility agent" means an individual who
402	holds a valid cannabis production establishment agent registration card with a cannabis

403	cultivation facility designation.
404	[(12)] (13) "Cannabis derivative product" means a product made using cannabis concentrate.
405	[(13)] (14) "Cannabis plant product" means any portion of a cannabis plant intended to be
406	sold in a form that is recognizable as a portion of a cannabis plant.
407	[(14)] (15) "Cannabis processing facility" means a person that:
408	(a) acquires or intends to acquire cannabis from a cannabis production establishment;
409	(b) possesses cannabis with the intent to manufacture a cannabis product;
410	(c) manufactures or intends to manufacture a cannabis product from unprocessed
411	cannabis or a cannabis extract; and
412	(d) sells or intends to sell a cannabis product to a medical cannabis pharmacy or a
413	medical cannabis research licensee.
414	[(15)] (16) "Cannabis processing facility agent" means an individual who
415	holds a valid cannabis production establishment agent registration card with a cannabis
416	processing facility designation.
417	[(16)] (17) "Cannabis product" means the same as that term is defined in Section 26B-4-201.
418	[(17)] (18) "Cannabis production establishment" means a cannabis cultivation facility, a
419	cannabis processing facility, or an independent cannabis testing laboratory.
420	[(18)] (19) "Cannabis production establishment agent" means a cannabis cultivation facility
421	agent, a cannabis processing facility agent, or an independent cannabis testing laboratory
422	agent.
423	[(19)] (20) "Cannabis production establishment agent registration card" means a registration
424	card that the department issues that:
425	(a) authorizes an individual to act as a cannabis production establishment agent; and
426	(b) designates the type of cannabis production establishment for which an individual is
427	authorized to act as an agent.
428	[(20)] (21) "Closed-door medical cannabis pharmacy" means a facility operated by a home
429	delivery medical cannabis pharmacy for delivering [eannabis or a medical cannabis
430	product] medical cannabis.
431	[(21)] (22) "Community location" means a public or private elementary or secondary school,
432	a church, a public library, a public playground, or a public park.
433	[(22)] (23) "Cultivation space" means, quantified in square feet, the horizontal area in which
434	a cannabis cultivation facility cultivates cannabis, including each level of horizontal area
435	if the cannabis cultivation facility hangs, suspends, stacks, or otherwise positions plants
436	above other plants in multiple levels.

437	[(23)] (24) "Delivery address" means:
438	(a) for a medical cannabis cardholder who is not a facility:
439	(i) the medical cannabis cardholder's home address; or
440	(ii) an address designated by the medical cannabis cardholder that:
441	(A) is the medical cannabis cardholder's workplace; and
442	(B) is not a community location; or
443	(b) for a medical cannabis cardholder that is a facility, the facility's address.
444	[(24)] (25) "Department" means the Department of Agriculture and Food.
445	[(25)] (26) "Family member" means a parent, step-parent, spouse, child, sibling,
446	step-sibling, uncle, aunt, nephew, niece, first cousin, mother-in-law, father-in-law,
447	brother-in-law, sister-in-law, son-in-law, daughter-in-law, grandparent, or grandchild.
448	[(26)] (27) "Government issued photo identification" means the same as that term is defined
449	in Section 26B-4-201, including expired identification in accordance with Section
450	26B-4-244.
451	[(27)] (28) "Home delivery medical cannabis pharmacy" means a medical cannabis
452	pharmacy that the department authorizes, as part of the pharmacy's license, to deliver
453	medical cannabis shipments to a delivery address to fulfill electronic orders[that the
454	state central patient portal facilitates].
455	[(28)] (29)(a) "Independent cannabis testing laboratory" means a person that:
456	(i) conducts a chemical or other analysis of cannabis or a cannabis product; or
457	(ii) acquires, possesses, and transports cannabis or a cannabis product with the intent
458	to conduct a chemical or other analysis of the cannabis or cannabis product.
459	(b) "Independent cannabis testing laboratory" includes a laboratory that the department
460	or a research university operates in accordance with Subsection 4-41a-201(14).
461	[(29)] (30) "Independent cannabis testing laboratory agent" means an individual who
462	holds a valid cannabis production establishment agent registration card with an
463	independent cannabis testing laboratory designation.
464	[(30)] (31) "Inventory control system" means a system described in Section 4-41a-103.
465	[(31)] (32) "Licensing board" or "board" means the Cannabis Production Establishment and
466	Pharmacy Licensing Advisory Board created in Section 4-41a-201.1.
467	[(32)] (33) "Medical cannabis" or "medical cannabis product" means the same as that term is
468	defined in Section 26B-4-201.
469	[(33)] (34) "Medical cannabis card" means the same as that term is defined in Section
470	26B-4-201.

- 471 [(34)] (35) "Medical cannabis courier" means a courier that:
- 472 (a) the department licenses in accordance with Section 4-41a-1201; and
- (b) contracts with a home delivery medical cannabis pharmacy to deliver medical cannabis shipments to fulfill electronic orders[-that the state central patient portal facilitates].
- 476 [(35)] (36) "Medical cannabis courier agent" means an individual who:
- 477 (a) is an employee of a medical cannabis courier; and
- (b) who holds a valid medical cannabis courier agent registration card.
- 479 [(36)] (37) "Medical cannabis pharmacy" means the same as that term is defined in Section 480 26B-4-201.
- 481 [(37)] (38) "Medical cannabis pharmacy agent" means the same as that term is defined in Section 26B-4-201.
- 483 [(38)] (39) "Medical cannabis research license" means a license that the department issues to 484 a research university for the purpose of obtaining and possessing medical cannabis for 485 academic research.
- 486 [(39)] (40) "Medical cannabis research licensee" means a research university that the 487 department licenses to obtain and possess medical cannabis for academic research, in 488 accordance with Section 4-41a-901.
- [(40)] (41) "Medical cannabis shipment" means a shipment of medical cannabis that a home delivery medical cannabis pharmacy or a medical cannabis courier delivers to a delivery address to fulfill an electronic medical cannabis order[-that the state central patient portal facilitates].
- 493 [(41)] (42) "Medical cannabis treatment" means the same as that term is defined in Section 494 26B-4-201.
- 495 [(42)] (43) "Medicinal dosage form" means the same as that term is defined in Section 496 26B-4-201.
- 497 [(43)] (44) "Pharmacy ownership limit" means an amount equal to 30% of the total number 498 of medical cannabis pharmacy licenses issued by the department rounded down to the 499 nearest whole number.
- 500 [(44)] (45) "Pharmacy medical provider" means the same as that term is defined in Section 26B-4-201.
- 502 [(45)] (46) "Qualified medical provider" means the same as that term is defined in Section 26B-4-201.
- 504 [(46)] (47) "Qualified Production Enterprise Fund" means the fund created in Section

505	4-41a-104.
506	[(47)] (48) "Recommending medical provider" means the same as that term is defined in
507	Section 26B-4-201.
508	[(48)] (49) "Research university" means the same as that term is defined in Section
509	53B-7-702 and a private, nonprofit college or university in the state that:
510	(a) is accredited by the Northwest Commission on Colleges and Universities;
511	(b) grants doctoral degrees; and
512	(c) has a laboratory containing or a program researching a schedule I controlled
513	substance described in Section 58-37-4.
514	[(49)] (50) "State electronic verification system" means the system described in Section
515	26B-4-202.
516	[(50)] (51) "Targeted marketing" means the promotion of [a cannabis product,] medical
517	cannabis, a medical cannabis brand, or a medical cannabis device using any of the
518	following methods:
519	(a) electronic communication to an individual who is at least 21 years old and has
520	requested to receive promotional information;
521	(b) an in-person marketing event that is:
522	(i) held inside a medical cannabis pharmacy; and
523	(ii) in an area where only a medical cannabis cardholder may access the event;
524	(c) other marketing material that is physically available or digitally displayed in a
525	medical cannabis pharmacy; or
526	(d) a leaflet a medical cannabis pharmacy places in the opaque package or box that is
527	provided to an individual when obtaining medical cannabis:
528	(i) in the medical cannabis pharmacy;
529	(ii) at the medical cannabis pharmacy's drive-through pick up window; or
530	(iii) in a medical cannabis shipment.
531	[(51)] (52) "Tetrahydrocannabinol" or "THC" means the same as that term is defined in
532	Section 4-41-102.
533	[(52)] (53) "THC analog" means the same as that term is defined in Section 4-41-102.
534	[(53)] (54) "Total composite tetrahydrocannabinol" means all detectable forms of
535	tetrahydrocannabinol.
536	[(54)] (55) "Total tetrahydrocannabinol" or "total THC" means the same as that term is
537	defined in Section 4-41-102.
538	Section 8. Section 4-41a-403 is amended to read:

539	4-41a-403 . Advertising.
540	(1) Except as provided in this section and Section 4-41a-604, a cannabis production
541	establishment may not advertise to the general public in any medium.
542	(2) A cannabis production establishment may advertise an employment opportunity at the
543	cannabis production establishment.
544	(3)(a) A cannabis production establishment may maintain a website that:
545	[(a)] (i) contains information about the establishment and employees; and
546	[(b)] (ii) except as provided in Subsection (3)(b), does not advertise any medical
547	cannabis, cannabis products, or medical cannabis devices.
548	(b) A cannabis processing facility may:
549	(i) if the website has age verification mechanisms that effectively prevent access by
550	individuals under 21 years of age, maintain a website that contains:
551	(A) educational information regarding medical cannabis produced by the cannabis
552	processing facility, including the certificate of analysis that is created by an
553	independent cannabis testing facility; and
554	(B) where medical cannabis produced by the cannabis processing facility may be
555	purchased in the state; and
556	(ii) engage in targeted marketing in accordance with Section 4-41a-604 for
557	advertising a particular medical cannabis product, medical cannabis device, or
558	medical cannabis brand.
559	(4)(a) Notwithstanding any municipal or county ordinance prohibiting signage, a
560	cannabis production establishment may use signage on the outside of the cannabis
561	production establishment that:
562	(i) includes only:
563	(A) in accordance with Subsection (4)(b), the cannabis production establishment's
564	name, logo, and hours of operation; and
565	(B) a green cross; and
566	(ii) complies with local ordinances regulating signage.
567	(b) The department shall define standards for a cannabis production establishment's
568	name and logo to ensure a medical rather than recreational disposition.
569	(5)(a) A cannabis production establishment may hold an educational event for the public
570	or medical providers in accordance with this Subsection (5) and the rules described in
571	Subsection (5)(c).
572	(b) A cannabis production establishment may not include in an educational event

573	described in Subsection (5)(a):
574	(i) any topic that conflicts with this chapter or Title 26B, Chapter 4, Part 2,
575	Cannabinoid Research and Medical Cannabis;
576	(ii) any gift items or merchandise other than educational materials, as those terms are
577	defined by the department;
578	(iii) any marketing for a specific product from the cannabis production establishment
579	or any other statement, claim, or information that would violate the federal Food,
580	Drug, and Cosmetic Act, 21 U.S.C. Sec. 301, et seq.; or
581	(iv) a presenter other than the following:
582	(A) a cannabis production establishment agent;
583	(B) a pharmacist licensed under Title 58, Chapter 17b, Pharmacy Practice Act;
584	(C) an advanced practice registered nurse licensed under Title 58, Chapter 31b,
585	Nurse Practice Act;
586	(D) a physician licensed under Title 58, Chapter 67, Utah Medical Practice Act, or
587	Title 58, Chapter 68, Utah Osteopathic Medical Practice Act;
588	(E) a physician assistant licensed under Title 58, Chapter 70a, Utah Physician
589	Assistant Act; or
590	(F) a state employee.
591	(c) The department shall make rules, in accordance with Title 63G, Chapter 3, Utah
592	Administrative Rulemaking Act, to define the elements of and restrictions on the
593	educational event described in Subsection (5)(a), including a minimum age of 21
594	years old for attendees.
595	Section 9. Section 4-41a-501 is amended to read:
596	4-41a-501. Cannabis cultivation facility Operating requirements.
597	(1) A cannabis cultivation facility shall ensure that any cannabis growing at the cannabis
598	cultivation facility is not visible from the ground level of the cannabis cultivation facility
599	perimeter.
600	(2) A cannabis cultivation facility shall use a unique identifier that is connected to the
601	facility's inventory control system to identify:
602	(a) beginning at the time a cannabis plant is eight inches tall and has a root ball, each
603	cannabis plant;
604	(b) each unique harvest of cannabis plants;
605	(c) each batch of cannabis the facility transfers to a medical cannabis pharmacy, a
606	cannabis processing facility, or an independent cannabis testing laboratory; and

607	(d) any excess, contaminated, or deteriorated cannabis of which the cannabis cultivation
608	facility disposes.
609	(3) A cannabis cultivation facility shall identify cannabis biomass as cannabis byproduct of
610	cannabis plant product before transferring the cannabis biomass from the facility.
611	(4) A cannabis cultivation facility shall either:
612	(a) ensure that a cannabis processing facility chemically or physically processes
613	cannabis cultivation byproduct to produce a cannabis concentrate for incorporation
614	into cannabis derivative products; or
615	(b) destroy cannabis cultivation byproduct in accordance with Section 4-41a-405.
616	(5) A cannabis cultivation facility may utilize radiation-based methods and equipment for
617	quality assurance or remediation purposes.
618	(6) The department shall make rules establishing:
619	(a) the records a cannabis cultivation facility must keep regarding each batch, amount
620	product treated, and the methods used; and
621	(b) disclosure requirements to a cannabis processor receiving the material subject to the
622	radiation including the methods and equipment used.
623	Section 10. Section 4-41a-701 is amended to read:
624	4-41a-701. Cannabis and cannabis product testing.
625	(1) In accordance with Title 63G, Chapter 3, Utah Administrative Rulemaking Act, the
626	department may make rules to:
627	(a) determine required adulterant tests for a cannabis plant product, cannabis
628	concentrate, or cannabis product;
629	(b) determine the amount of any adulterant that is safe for human consumption;
630	(c) immediately ban or limit the presence of any ingredient in a medical cannabis
631	product after receiving a recommendation to do so from a public health authority
632	under Section 26B-1-102;
633	(d) establish protocols for a recall of [eannabis or a eannabis product] medical cannab
634	by a cannabis production establishment; or
635	(e) allow the propagation of testing results forward to derived product if the processin
636	steps the cannabis production establishment uses to produce the product are unlike
637	to change the results of the test.
638	(2)(a) The department may require testing for a toxin if:
639	[(a)] (i) the department receives information indicating the potential presence of a
640	toxin; or

641	[(b)] (ii) the department's inspector has reason to believe a toxin may be present based
642	on the inspection of a facility.
643	(b) The department may not require a cannabis processor to test a cannabis batch or a
644	cannabis product batch a third time if the cannabis batch or cannabis product has
645	previously met all testing requirements after being tested by:
646	(i) an independent cannabis testing laboratory that is not the department; and
647	(ii) the department.
648	(3)(a) A cannabis production establishment may not:
649	(i) incorporate cannabis concentrate into a cannabis derivative product until an
650	independent cannabis testing laboratory tests the cannabis concentrate in
651	accordance with department rule; or
652	(ii) transfer cannabis or a cannabis product to a medical cannabis pharmacy until an
653	independent cannabis testing laboratory tests a representative sample of the
654	cannabis or cannabis product in accordance with department rule.
655	(b) A medical cannabis pharmacy may not offer any cannabis or cannabis product for
656	sale unless an independent cannabis testing laboratory has tested a representative
657	sample of the cannabis or cannabis product in accordance with department rule.
658	(4) Before the sale of a medical cannabis product, an independent cannabis testing
659	laboratory shall:
660	(a) identify and quantify any cannabinoid known to be present in [a] the medical
661	cannabis product; and
662	(b) test terpene profiles for the following products:
663	(i) raw cannabis; or
664	(ii) a cannabis product:
665	(A) contained in a vaporizer cartridge; or
666	(B) in concentrate form; and
667	(c) record the five highest terpene profiles tested under Subsection (4)(b).
668	(5) The department shall establish by rule, in accordance with Title 63G, Chapter 3, Utah
669	Administrative Rulemaking Act, the standards, methods, practices, and procedures for
670	the testing of cannabis and cannabis products by independent cannabis testing
671	laboratories.
672	(6) The department may require an independent cannabis testing laboratory to participate in
673	a proficiency evaluation that the department conducts or that an organization that the
674	denartment approves conducts

675	Section 11. Section 4-41a-801 is amended to read:
676	4-41a-801 . Enforcement Fine Citation.
677	(1) If a person that is a cannabis production establishment or a cannabis production
678	establishment agent violates this chapter, the department may:
679	(a) revoke the person's license or cannabis production establishment agent registration
680	card;
681	(b) decline to renew the person's license or cannabis production establishment agent
682	registration card; or
683	(c) assess the person an administrative penalty that the department establishes by rule in
684	accordance with Title 63G, Chapter 3, Utah Administrative Rulemaking Act.
685	(2) The department shall deposit an administrative penalty imposed under this section into
686	the General Fund.
687	(3)(a) The department may take an action described in Subsection (3)(b) if the
688	department concludes, upon investigation, that, for a person that is a cannabis
689	production establishment or a cannabis production establishment agent:
690	(i) the person has violated the provisions of this chapter, a rule made under this
691	chapter, or an order issued under this chapter; or.
692	(ii) the person produced cannabis or a cannabis product batch that contains a
693	substance, other than cannabis, that poses a significant threat to human health.
694	(b) If the department makes the determination about a person described in Subsection
695	(3)(a), the department shall:
696	(i) issue the person a written administrative citation;
697	(ii) attempt to negotiate a stipulated settlement;
698	[(iii) seize, embargo, or destroy the cannabis or cannabis product batch;]
699	[(iv)] (iii) order the person to cease and desist from the action that creates a violation; [
700	and] <u>or</u>
701	[(v)] (iv) direct the person to appear before an adjudicative proceeding conducted
702	under Title 63G, Chapter 4, Administrative Procedures Act.
703	(c) If the department concludes, upon investigation, that a cannabis production
704	establishment or a cannabis production establishment agent has produced a cannabis
705	batch or a cannabis product batch that contains a substance that poses a significant
706	threat to human health, the department shall seize, embargo, or destroy the cannabis
707	batch or cannabis product batch.

(4) The department may, for a person subject to an uncontested citation, a stipulated

709	settlement, or a finding of a violation in an adjudicative proceeding under this section,
710	for a fine amount not already specified in law, assess the person, who is not an
711	individual, a fine of up to \$5,000 per violation, in accordance with a fine schedule that
712	the department establishes by rule in accordance with Title 63G, Chapter 3, Utah
713	Administrative Rulemaking Act.
714	(5) The department may not revoke a [eannabis production establishment's-]license without
715	first directing the [eannabis production establishment] licensee to appear before an
716	adjudicative proceeding conducted under Title 63G, Chapter 4, Administrative
717	Procedures Act.
718	(6) If within $[2\theta]$ $\underline{30}$ calendar days after the day on which a department serves a citation for
719	a violation of this chapter, the person that is the subject of the citation fails to request a
720	hearing to contest the citation, the citation becomes the department's final order.
721	(7) The department may, for a person who fails to comply with a citation under this section:
722	(a) refuse to issue or renew the person's license or cannabis production establishment
723	agent registration card; or
724	(b) suspend, revoke, or place on probation the person's license or cannabis production
725	establishment registration card.
726	(8)(a) Except where a criminal penalty is expressly provided for a specific violation of
727	this chapter, if an individual:
728	(i) violates a provision of this chapter, the individual is:
729	(A) guilty of an infraction; and
730	(B) subject to a \$100 fine; or
731	(ii) intentionally or knowingly violates a provision of this chapter or violates this
732	chapter three or more times, the individual is:
733	(A) guilty of a class B misdemeanor; and
734	(B) subject to a \$1,000 fine.
735	(b) An individual who is guilty of a violation described in Subsection (8)(a) is not guilty
736	of a violation of Title 58, Chapter 37, Utah Controlled Substances Act, for the
737	conduct underlying the violation described in Subsection (8)(a).
738	(9) Nothing in this section prohibits the department from referring potential criminal
739	activity to law enforcement.
740	(10) An appeal of administrative action taken under this chapter shall be heard by an
741	administrative law judge as an informal proceeding in accordance with Title 63G,
742	Chapter 4, Administrative Procedures Act.

743	Section 12. Section 4-41a-802 is amended to read:
744	4-41a-802 . Report.
745	(1) At or before the November interim meeting each year, the department shall report to the
746	Health and Human Services Interim Committee on:
747	(a) the number of applications and renewal applications that the department receives
748	under this chapter;
749	(b) the number of each type of cannabis production facility that the department licenses
750	in each county;
751	(c) the amount of cannabis that licensees grow;
752	(d) the amount of cannabis that licensees manufacture into cannabis products;
753	(e) the number of licenses the department revokes under this chapter;
754	(f) the department's operation of an independent cannabis testing laboratory under
755	Section 4-41a-201, including:
756	(i) the cannabis and cannabis products the department tested; and
757	(ii) the results of the tests the department performed;
758	(g) the expenses incurred and revenues generated under this chapter; and
759	(h) an analysis of product availability in medical cannabis pharmacies in consultation
760	with the Department of Health and Human Services.
761	(2) The department may not include personally identifying information in the report
762	described in this section.
763	(3) The department shall report to the working group described in Section 36-12-8.2 as
764	requested by the working group.
765	(4)(a) Before August 1, of each year, the department shall provide a report to the
766	working group described in Section 36-12-8.2 that provides the following for each
767	fine issued by the department under this chapter:
768	(i) the date of the fine;
769	(ii) the reference to the statute or rule that was violated for each fine issued; and
770	(iii) a short description explaining why the fine was issued.
771	(b) The report described in Subsection (4)(a) may not include identifying information of
772	the person that was subject to the fine.
773	Section 13. Section 4-41a-1001 is amended to read:
774	4-41a-1001 . Medical cannabis pharmacy License Eligibility.
775	(1) A person may not:

(a) operate as a medical cannabis pharmacy without a license that the department issues

777	under this part;
778	(b) obtain a medical cannabis pharmacy license if obtaining the license would cause the
779	person to exceed the pharmacy ownership limit;
780	(c) obtain a partial ownership share of a medical cannabis pharmacy if obtaining the
781	partial ownership share would cause the person to exceed the pharmacy ownership
782	limit; or
783	(d) enter into any contract or agreement that allows the person to directly or indirectly
784	control the operations of a medical cannabis pharmacy if the person's control of the
785	medical cannabis pharmacy would cause the person to effectively exceed the
786	pharmacy ownership limit.
787	(2)(a)(i) Subject to Subsections (4) and (5) and to Section 4-41a-1005, the department
788	shall issue a license to operate a medical cannabis pharmacy through the licensing
789	board created under Section 4-41a-201.1.
790	(ii) The department may not issue a license to operate a medical cannabis pharmacy
791	to an applicant who is not eligible for a license under this section.
792	(b) An applicant is eligible for a license under this section if the applicant submits to the
793	department:
794	(i) subject to Subsection (2)(c), a proposed name and address where the applicant will
795	operate the medical cannabis pharmacy;
796	(ii) the name and address of an individual who:
797	(A) for a publicly traded company, has a financial or voting interest of 10% or
798	greater in the proposed medical cannabis pharmacy;
799	(B) for a privately held company, a financial or voting interest in the proposed
800	medical cannabis pharmacy; or
801	(C) has the power to direct or cause the management or control of a proposed
802	medical cannabis pharmacy;
803	(iii) for each application that the applicant submits to the department, a statement
804	from the applicant that the applicant will obtain and maintain:
805	(A) a performance bond in the amount of \$100,000 issued by a surety authorized
806	to transact surety business in the state; or
807	(B) a liquid cash account in the amount of \$100,000 with a financial institution;
808	(iv) an operating plan that:
809	(A) complies with Section 4-41a-1004;
810	(B) includes operating procedures to comply with the operating requirements for a

811	medical cannabis pharmacy described in this part and with a relevant municipal
812	or county law that is consistent with Section 4-41a-1106; and
813	(C) the department approves;
814	(v) an application fee in an amount that, subject to Subsection 4-41a-104(5), the
815	department sets in accordance with Section 63J-1-504; and
816	(vi) a description of any investigation or adverse action taken by any licensing
817	jurisdiction, government agency, law enforcement agency, or court in any state for
818	any violation or detrimental conduct in relation to any of the applicant's
819	cannabis-related operations or businesses.
820	(c)(i) A person may not locate a medical cannabis pharmacy:
821	(A) within 200 feet of a community location; or
822	(B) in or within 600 feet of a district that the relevant municipality or county has
823	zoned as primarily residential.
824	(ii) The proximity requirements described in Subsection (2)(c)(i) shall be measured
825	from the nearest entrance to the medical cannabis pharmacy establishment by
826	following the shortest route of ordinary pedestrian travel to the property boundary
827	of the community location or residential area.
828	(iii) The department may grant a waiver to reduce the proximity requirements in
829	Subsection (2)(c)(i) by up to 20% if the department determines that it is not
830	reasonably feasible for the applicant to cite the proposed medical cannabis
831	pharmacy without the waiver.
832	(iv) An applicant for a license under this section shall provide evidence of
833	compliance with the proximity requirements described in Subsection (2)(c)(i).
834	(d) The department may not issue a license to an eligible applicant that the department
835	has selected to receive a license until the selected eligible applicant complies with the
836	bond or liquid cash requirement described in Subsection (2)(b)(iii).
837	(e) If the department receives more than one application for a medical cannabis
838	pharmacy within the same city or town, the department shall consult with the local
839	land use authority before approving any of the applications pertaining to that city or
840	town.
841	(f) In considering the issuance of a medical cannabis pharmacy license under this
842	section, the department may consider the extent to which the pharmacy can increase
843	efficiency and reduce cost to patients of medical cannabis.
844	[(3) If the department selects an applicant-]

845	(3)(a) After an entity has been selected for a medical cannabis pharmacy license under
846	this section, the department shall:
847	[(a)] (i) charge the applicant an initial license fee in an amount that, subject to
848	Subsection 4-41a-104(5), the department sets in accordance with Section
849	63J-1-504;
850	[(b)] (ii) notify the Department of Public Safety of the license approval and the names
851	of each individual described in Subsection (2)(b)(ii); and
852	[(e)] (iii) charge the licensee a fee in an amount that, subject to Subsection 4-41a-104
853	(5), the department sets in accordance with Section 63J-1-504, for any change in
854	location, ownership, or company structure.
855	(b) For a fee described in Subsection (3)(a)(i), a license fee for a medical cannabis
856	pharmacy located in a medically underserved area as determined by the federal
857	Health Resources and Services Administration shall be 50% less than what is charged
858	for other medical cannabis pharmacies.
859	(4) The department may not issue a license to operate a medical cannabis pharmacy to an
860	applicant if an individual described in Subsection (2)(b)(ii):
861	(a) has been convicted under state or federal law of:
862	(i) a felony in the preceding 10 years; or
863	(ii) after December 3, 2018, a misdemeanor for drug distribution;
864	(b) is younger than 21 years old; or
865	(c) after September 23, 2019, until January 1, 2023, is actively serving as a legislator.
866	(5)[(a)] If an applicant for a medical cannabis pharmacy license under this section holds
867	another license under this chapter, the department may not give preference to the
868	applicant based on the applicant's status as a holder of the license.
869	[(b) If an applicant for a medical cannabis pharmacy license under this section holds a
870	license to operate a cannabis cultivation facility under this section, the department
871	may give consideration to the applicant's status as a holder of the license if:]
872	[(i) the applicant demonstrates that a decrease in costs to patients is more likely to
873	result from the applicant's vertical integration than from a more competitive
874	marketplace; and]
875	[(ii) the department finds multiple other factors, in addition to the existing license,
876	that support granting the new license.]
877	(6) The licensing board may revoke a license under this part:
878	(a) if the medical cannabis pharmacy does not begin operations within one year after the

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879	day on which the department issues an announcement of the department's intent to	
880	award a license to the medical cannabis pharmacy;	
881	(b) after the third the same violation of this chapter in any of the licensee's licensed	
882	cannabis production establishments or medical cannabis pharmacies;	
883	(c) if an individual described in Subsection (2)(b)(ii) is convicted, while the license is	
884	active, under state or federal law of:	
885	(i) a felony; or	
886	(ii) after December 3, 2018, a misdemeanor for drug distribution;	
887	(d) if the licensee fails to provide the information described in Subsection (2)(b)(vi) at	
888	the time of application, or fails to supplement the information described in	
889	Subsection (2)(b)(vi) with any investigation or adverse action that occurs after the	
890	submission of the application within 14 calendar days after the licensee receives	
891	notice of the investigation or adverse action;	
892	(e) if the medical cannabis pharmacy demonstrates a willful or reckless disregard for the	ıe
893	requirements of this chapter or the rules the department makes in accordance with	
894	this chapter;	
895	(f) if, after a change of ownership described in Subsection (11)(c), the department	
896	determines that the medical cannabis pharmacy no longer meets the minimum	
897	standards for licensure and operation of the medical cannabis pharmacy described i	n
898	this chapter; or	
899	(g) if through an investigation conducted under Subsection 4-41a-201.1(11) and in	
900	accordance with Title 63G, Chapter 4, Administrative Procedures Act, the board	
901	finds that the licensee has participated in anticompetitive business practices.	
902	(7)(a) A person who receives a medical cannabis pharmacy license under this chapter, if	
903	the municipality or county where the licensed medical cannabis pharmacy will be	
904	located requires a local land use permit, shall submit to the department a copy of the	
905	licensee's approved application for the land use permit within 120 days after the day	
906	on which the department issues the license.	
907	(b) If a licensee fails to submit to the department a copy the licensee's approved land us	se
908	permit application in accordance with Subsection (7)(a), the department may revoke	e
909	the licensee's license.	

(9) The department shall begin accepting applications under this part on or before March 1,

(8) The department shall deposit the proceeds of a fee imposed by this section into the

Qualified Production Enterprise Fund.

913	2020.
914	(10)(a) The department's authority to issue a license under this section is plenary and is
915	not subject to review.
916	(b) Notwithstanding Subsection (2), the decision of the department to award a license to
917	an applicant is not subject to:
918	(i) Title 63G, Chapter 6a, Part 16, Protests; or
919	(ii) Title 63G, Chapter 6a, Part 17, Procurement Appeals Board.
920	(11)(a) A medical cannabis pharmacy license is not transferrable or assignable.
921	(b) A medical cannabis pharmacy shall report in writing to the department no later than
922	10 business days before the date of any change of ownership of the medical cannabis
923	pharmacy.
924	(c) If the ownership of a medical cannabis pharmacy changes by 50% or more:
925	(i) concurrent with the report described in Subsection (11)(b), the medical cannabis
926	pharmacy shall submit a new application described in Subsection (2)(b), subject t
927	Subsection (2)(c);
928	(ii) within 30 days of the submission of the application, the department shall:
929	(A) conduct an application review; and
930	(B) award a license to the medical cannabis pharmacy for the remainder of the
931	term of the medical cannabis pharmacy's license before the ownership change
932	if the medical cannabis pharmacy meets the minimum standards for licensure
933	and operation of the medical cannabis pharmacy described in this chapter; an
934	(iii) if the department approves the license application, notwithstanding Subsection
935	(3), the medical cannabis pharmacy shall pay a license fee that the department set
936	in accordance with Section 63J-1-504 in an amount that covers the department's
937	cost of conducting the application review.
938	Section 14. Section 4-41a-1003 is amended to read:
939	4-41a-1003 . Renewal - Notice of available license.
940	(1)(a) The department shall renew a license [under Sections 4-41a-1001 through
941	4-41a-1005] issued under this part every year if, at the time of renewal:
942	[(a)] (i) the licensee meets the requirements of Section 4-41a-1001;
943	[(b)] (ii) the licensee pays the department a license renewal fee in an amount that,
944	subject to Subsection 4-41a-1004(5), the department sets in accordance with
945	Section 63J-1-504; and
946	[(e)] (iii) if the medical cannabis pharmacy changes the operating plan described in

947	Section 4-41a-1004 that the department approved under Subsection
948	4-41a-1001(2)(b)(iv), the department approves the new operating plan.
949	(b) A license fee for a medical cannabis pharmacy located in a county of the third,
950	fourth, fifth, or sixth class shall be 50% less than what is charged for other medical
951	cannabis pharmacies.
952	(2)(a) If a licensed medical cannabis pharmacy abandons the medical cannabis
953	pharmacy's license, the department shall publish notice of an available license[-], for
954	the geographic area in which the medical cannabis pharmacy license is available, as a
955	class A notice under Section 63G-30-102, for at least seven days.
956	(b) The department may establish criteria, in collaboration with the Division of
957	Professional Licensing and the Board of Pharmacy and in accordance with Title 63G,
958	Chapter 3, Utah Administrative Rulemaking Act, to identify the medical cannabis
959	pharmacy actions that constitute abandonment of a medical cannabis pharmacy
960	license.
961	(3) If the department has not completed the necessary processes to make a determination on
962	a license renewal under Subsections (1)(a) and (c) before the expiration of a license, the
963	department may issue a conditional medical cannabis pharmacy license to a licensed
964	medical cannabis pharmacy that has applied for license renewal under this section and
965	paid the fee described in Subsection (1)(b).
966	Section 15. Section 4-41a-1005 is amended to read:
967	4-41a-1005 . Maximum number of licenses.
968	(1)(a) [Except as provided in Subsection (1)(b) or (d), if a sufficient number of
969	applicants apply, the department] The licensing board shall issue up to [15] 17 medical
970	cannabis pharmacy licenses in accordance with this section including the two medical
971	cannabis pharmacy licenses in accordance with Section 4-41a-1006.
972	[(b) If an insufficient number of qualified applicants apply for the available number of
973	medical cannabis pharmacy licenses, the department shall issue a medical cannabis
974	pharmacy license to each qualified applicant.]
975	[(c) The department may issue the licenses described in Subsection (1)(a) in accordance
976	with this Subsection (1)(c).]
977	[(i) Using one procurement process, the department may issue eight licenses to an
978	initial group of medical cannabis pharmacies and six licenses to a second group of
979	medical cannabis pharmacies.]
980	[(ii) The department shall:]

981	(A) divide the state into no less than four geographic regions, set by the
982	department in rule;]
983	[(B) issue at least one license in each geographic region during each phase of
984	issuing licenses; and]
985	[(C) complete the process of issuing medical cannabis pharmacy licenses no later
986	than July 1, 2020.]
987	[(iii) In issuing a 15th license under Subsection (1), the department shall ensure that
988	the license recipient will locate the medical cannabis pharmacy within Dagget,
989	Duchesne, Uintah, Carbon, Sevier, Emery, Grand, or San Juan County.]
990	[(d)(i) The department may issue licenses to operate a medical cannabis pharmacy in
991	addition to the licenses described in Subsection (1)(a) if the department
992	determines, in consultation with the Department of Health and Human Services
993	and after an annual or more frequent analysis of the current and anticipated market
994	for medical cannabis, that each additional license is necessary to provide an
995	adequate supply, quality, or variety of medical cannabis to medical cannabis
996	cardholders.]
997	[(ii) The department shall:]
998	[(A) in accordance with Title 63G, Chapter 3, Utah Administrative Rulemaking
999	Act, make rules to establish criteria and processes for the consultation,
1000	analysis, and application for a license described in Subsection (1)(d)(i); and]
1001	[(B) report to the Executive Appropriations Committee of the Legislature before
1002	each time the department issues an additional license under Subsection
1003	(1)(d)(i) regarding the results of the consultation and analysis described in
1004	Subsection (1)(d)(i) and the application of the criteria described in Subsection
1005	(1)(d)(ii)(A).]
1006	(2)(a) [If there are more qualified applicants than there are available licenses for medical
1007	eannabis pharmacies, the department] The licensing board shall:
1008	(i) evaluate each applicant and award the license to the applicant that best
1009	demonstrates:
1010	(A) experience with establishing and successfully operating a business that
1011	involves complying with a regulatory environment, tracking inventory, and
1012	training, evaluating, and monitoring employees;
1013	(B) an operating plan that will best ensure the safety and security of patrons and
1014	the community;

1015	(C) positive connections to the local community;
1016	(D) the suitability of the proposed location and the location's accessibility for
1017	qualifying patients;
1018	(E) the extent to which the applicant can increase efficiency and reduce the cost of
1019	medical cannabis for patients; and
1020	(F) a strategic plan described in Subsection 4-41a-1004(7) that has a
1021	comparatively high likelihood of success; and
1022	(ii) ensure a geographic dispersal among licensees that is sufficient to reasonably
1023	maximize access to the largest number of medical cannabis cardholders.
1024	(b) In making the evaluation described in Subsection (2)(a), the licensing board may
1025	give increased consideration to applicants who indicate a willingness to:
1026	(i) site a medical cannabis pharmacy in an area or population center designated as a
1027	medically underserved area or population as determined by the federal Health
1028	Resources and Services Administration; and
1029	(ii) operate as a home delivery medical cannabis pharmacy that accepts electronic
1030	medical cannabis orders.
1031	[(b) In making the evaluation described in Subsection (2)(a), the department may give
1032	increased consideration to applicants who indicate a willingness to:]
1033	[(i) operate as a home delivery medical cannabis pharmacy that accepts electronic
1034	medical cannabis orders that the state central patient portal facilitates; and]
1035	[(ii) accept payments through:]
1036	[(A) a payment provider that the Division of Finance approves, in consultation
1037	with the state treasurer, in accordance with Section 4-41a-108; or]
1038	[(B) a financial institution in accordance with Subsection 4-41a-108(4).]
1039	(3) The [department] licensing board may conduct a face-to-face interview with an applicant
1040	for a license that the [department] <u>licensing board</u> evaluates under Subsection (2).
1041	Section 16. Section 4-41a-1006 is enacted to read:
1042	$\underline{4\text{-}41a\text{-}1006}$. Independent medical cannabis licenses.
1043	(1)(a) Subject to the requirements of Subsection (3) and the criteria established for
1044	obtaining a medical cannabis pharmacy license under this chapter, the licensing
1045	board shall:
1046	(i) before January 1, 2026, select one entity to receive a medical cannabis pharmacy
1047	license; and
1048	(ii) before January 1, 2027, but not before January 1, 2026, select one entity to

1049	receive a medical cannabis pharmacy license.
1050	(b) When selecting entities under this section, if there is a conflict between the criteria
1051	established for obtaining a medical cannabis pharmacy license under the other
1052	sections of this chapter and this section, this section controls.
1053	(2) For the license described in Subsection (1)(a)(ii), the licensing board may not select an
1054	entity:
1055	(a) that owns any interest in or operates a medical cannabis production establishment; or
1056	(b) that is owned, partially or entirely, or operated by a medical cannabis production
1057	establishment.
1058	(3) The licensing board:
1059	(a) may not select an entity to receive a license under this section if the entity owns a
1060	financial interest in a medical cannabis pharmacy or is owned by an entity that owns
1061	a financial interest in a medical cannabis pharmacy; and
1062	(b) shall select an entity that will site a medical cannabis pharmacy license issued under
1063	this section in an area:
1064	(i) designated as a medically underserved area as determined by the federal Health
1065	Resources and Services Administration; and
1066	(ii) located in a county of the third, fourth, fifth, or sixth class.
1067	(4) A license described in this section may not be transferred to another entity unless that
1068	entity meets the requirements of Subsections (2) and (3) that the transferring entity met
1069	when obtaining the license.
1070	(5) Notwithstanding Subsection (4), for a license described in Subsection (1)(a)(i), an
1071	applicant shall commit to not alienating or otherwise transferring control of the license
1072	or of the entity that holds the license to another person for at least 15 years from the day
1073	the license is issued under this chapter.
1074	(6) The department shall provide regular updates to the Medical Cannabis Governance
1075	Structure Working Group created in Section 36-12-8.2 regarding the application and
1076	selection process for licenses issued under this section.
1077	Section 17. Section 4-41a-1101 is amended to read:
1078	4-41a-1101 . Operating requirements General.
1079	(1)(a) A medical cannabis pharmacy shall operate:
1080	(i) at the physical address provided to the department under Section 4-41a-1001; and
1081	(ii) in accordance with the operating plan provided to the department under Section
1082	4-41a-1001 and, if applicable, Section 4-41a-1004.

1083		(b) A medical cannabis pharmacy shall notify the department before a change in the
1084		medical cannabis pharmacy's physical address or operating plan.
1085	(2)	An individual may not enter a medical cannabis pharmacy unless the individual:
1086		(a) is at least 18 years old or is an emancipated minor under Section 80-7-105; and
1087		(b) except as provided in Subsection (4):
1088		(i) possesses a valid:
1089		(A) medical cannabis pharmacy agent registration card;
1090		(B) pharmacy medical provider registration card; or
1091		(C) medical cannabis card;
1092		(ii) is an employee of the department performing an inspection under Section
1093		4-41a-1103; or
1094		(iii) is another individual as the department provides.
1095	(3)	A medical cannabis pharmacy may not employ an individual who is younger than 21
1096		years old.
1097	(4)	Notwithstanding Subsection (2)(a), a medical cannabis pharmacy may authorize an
1098		individual who is not a medical cannabis pharmacy agent or pharmacy medical provider
1099		to access the medical cannabis pharmacy if the medical cannabis pharmacy tracks and
1100		monitors the individual at all times while the individual is at the medical cannabis
1101		pharmacy and maintains a record of the individual's access.
1102	(5)	A medical cannabis pharmacy shall operate in a facility that has:
1103		(a) a single, secure public entrance;
1104		(b) a security system with a backup power source that:
1105		(i) detects and records entry into the medical cannabis pharmacy; and
1106		(ii) provides notice of an unauthorized entry to law enforcement when the medical
1107		cannabis pharmacy is closed; and
1108		(c) a lock on each area where the medical cannabis pharmacy stores [eannabis or a
1109		cannabis product] medical cannabis.
1110	(6)	A medical cannabis pharmacy shall post, both clearly and conspicuously in the medical
1111		cannabis pharmacy, the limit on the purchase of cannabis described in Subsection
1112		4-41a-1102(2).
1113	(7)	Except for an emergency situation described in Subsection 26B-4-213(3)(c), a medical
1114		cannabis pharmacy may not allow any individual to consume cannabis on the property
1115		or premises of the medical cannabis pharmacy.
1116	(8)	A medical cannabis pharmacy may not sell [eannabis or a cannabis product] medical

1117	<u>cannabis</u> without first indicating on the [cannabis or cannabis product] <u>medical cannabis</u>
1118	label the name of the medical cannabis pharmacy.
1119	(9)(a) Each medical cannabis pharmacy shall retain in the pharmacy's records the
1120	following information regarding each recommendation underlying a transaction:
1121	(i) the recommending medical provider's name, address, and telephone number;
1122	(ii) the patient's name and address;
1123	(iii) the date of issuance;
1124	(iv) directions of use and dosing guidelines or an indication that the recommending
1125	medical provider did not recommend specific directions of use or dosing
1126	guidelines; and
1127	(v) if the patient did not complete the transaction, the name of the medical cannabis
1128	cardholder who completed the transaction.
1129	(b)(i) Except as provided in Subsection (9)(b)(iii), a medical cannabis pharmacy may
1130	not sell medical cannabis unless the medical cannabis has a label securely affixed
1131	to the container indicating the following minimum information:
1132	(A) the name, address, and telephone number of the medical cannabis pharmacy;
1133	(B) the unique identification number that the medical cannabis pharmacy assigns;
1134	(C) the date of the sale;
1135	(D) the name of the patient;
1136	(E) the name of the recommending medical provider who recommended the
1137	medical cannabis treatment;
1138	(F) directions for use and cautionary statements, if any;
1139	(G) the amount dispensed and the cannabinoid content;
1140	(H) the suggested use date;
1141	(I) for unprocessed cannabis flower, the legal use termination date; and
1142	(J) any other requirements that the department determines, in consultation with the
1143	Division of Professional Licensing and the Board of Pharmacy.
1144	(ii) A medical cannabis pharmacy is exempt from the requirement to provide the
1145	following information under Subsection (9)(b)(i) if the information is already
1146	provided on the product label that a cannabis production establishment affixes:
1147	(A) a unique identification number;
1148	(B) directions for use and cautionary statements;
1149	(C) amount and cannabinoid content; and
1150	(D) a suggested use date.

1151	(iii) If the size of a medical cannabis container does not allow sufficient space to
1152	include the labeling requirements described in Subsection (9)(b)(i), the medical
1153	cannabis pharmacy may provide the following information described in
1154	Subsection (9)(b)(i) on a supplemental label attached to the container or an
1155	informational enclosure that accompanies the container:
1156	(A) the cannabinoid content;
1157	(B) the suggested use date; and
1158	(C) any other requirements that the department determines.
1159	(iv) A medical cannabis pharmacy may sell medical cannabis to another medical
1160	cannabis pharmacy without a label described in Subsection (9)(b)(i).
1161	(10) A pharmacy medical provider or medical cannabis pharmacy agent shall:
1162	(a) upon receipt of an order from a limited medical provider in accordance with
1163	Subsections 26B-4-204(1)(b) through (d):
1164	(i) for a written order or an electronic order under circumstances that the department
1165	determines, contact the limited medical provider or the limited medical provider's
1166	office to verify the validity of the recommendation; and
1167	(ii) for an order that the pharmacy medical provider or medical cannabis pharmacy
1168	agent verifies under Subsection (10)(a)(i) or an electronic order that is not subject
1169	to verification under Subsection (10)(a)(i), enter the limited medical provider's
1170	recommendation or renewal, including any associated directions of use, dosing
1171	guidelines, or caregiver indication, in the state electronic verification system;
1172	(b) in processing an order for a holder of a conditional medical cannabis card described
1173	in Subsection 26B-4-213(1)(b) that appears irregular or suspicious in the judgment of
1174	the pharmacy medical provider or medical cannabis pharmacy agent, contact the
1175	recommending medical provider or the recommending medical provider's office to
1176	verify the validity of the recommendation before processing the cardholder's order;
1177	(c) unless the medical cannabis cardholder has had a consultation under Subsection
1178	26B-4-231(5), verbally offer to a medical cannabis cardholder at the time of a
1179	purchase of [eannabis, a cannabis product,] medical cannabis or a medical cannabis
1180	device, personal counseling with the pharmacy medical provider; and
1181	(d) provide a telephone number or website by which the cardholder may contact a
1182	pharmacy medical provider for counseling.
1183	(11)(a) A medical cannabis pharmacy may create a medical cannabis disposal program
1184	that allows an individual to deposit unused or excess medical cannabis or cannabis

1185	residue from a medical cannabis device in a locked box or other secure receptacle
1186	within the medical cannabis pharmacy.
1187	(b) A medical cannabis pharmacy with a disposal program described in Subsection
1188	(11)(a) shall ensure that only a medical cannabis pharmacy agent or pharmacy
1189	medical provider can access deposited medical cannabis.
1190	(c) A medical cannabis pharmacy shall dispose of any deposited medical cannabis by:
1191	(i) rendering the deposited medical cannabis unusable and unrecognizable before
1192	transporting deposited medical cannabis from the medical cannabis pharmacy; and
1193	(ii) disposing of the deposited medical cannabis in accordance with:
1194	(A) federal and state law, rules, and regulations related to hazardous waste;
1195	(B) the Resource Conservation and Recovery Act, 42 U.S.C. Sec. 6991 et seq.;
1196	(C) Title 19, Chapter 6, Part 5, Solid Waste Management Act; and
1197	(D) other regulations that the department makes in accordance with Title 63G,
1198	Chapter 3, Utah Administrative Rulemaking Act.
1199	(12) A medical cannabis pharmacy:
1200	(a) shall employ a pharmacist who is licensed under Title 58, Chapter 17b, Pharmacy
1201	Practice Act, as a pharmacy medical provider;
1202	(b) may employ a physician who has the authority to write a prescription and is licensed
1203	under Title 58, Chapter 67, Utah Medical Practice Act, or Title 58, Chapter 68, Utah
1204	Osteopathic Medical Practice Act, as a pharmacy medical provider;
1205	(c) shall ensure that a pharmacy medical provider described in Subsection (12)(a) works
1206	onsite during all business hours;
1207	(d) shall designate one pharmacy medical provider described in Subsection (12)(a) as the
1208	pharmacist-in-charge to oversee the operation of and generally supervise the medical
1209	cannabis pharmacy;[and]
1210	(e) shall allow the pharmacist-in-charge to determine which [cannabis and cannabis
1211	products] medical cannabis products the medical cannabis pharmacy maintains in the
1212	medical cannabis pharmacy's inventory[-]; and
1213	(f) for each medical cannabis product sold by the medical cannabis pharmacy, shall:
1214	(i) allow a medical cannabis cardholder located in the pharmacy to view the back
1215	panel of the product when requested; and
1216	(ii) beginning July 1, 2025, include a picture of the back panel of the product on the
1217	medical cannabis pharmacy's website.
1218	(13) The department shall establish by rule, in accordance with Title 63G, Chapter 3, Utah

1219	Administrative Rulemaking Act, protocols for a recall of cannabis and cannabis products
1220	by a medical cannabis pharmacy.
1221	Section 18. Section 4-41a-1201 is amended to read:
1222	4-41a-1201. Medical cannabis home delivery designation.
1223	(1) The department may designate a medical cannabis pharmacy as a home delivery
1224	medical cannabis pharmacy if the department determines that the medical cannabis
1225	pharmacy's operating plan demonstrates the functional and technical ability to:
1226	(a) safely conduct transactions for medical cannabis shipments;
1227	(b) accept electronic medical cannabis orders[-that the state central patient portal
1228	facilitates]; and
1229	(c) accept payments through:
1230	(i) a payment provider that the Division of Finance approves, in consultation with the
1231	state treasurer, in accordance with Section 26-61a-603; or
1232	(ii) a financial institution in accordance with Subsection 26-61a-603(4).
1233	(2) An applicant seeking a designation as a home delivery medical cannabis pharmacy shall
1234	identify in the applicant's operating plan any information relevant to the department's
1235	evaluation described in Subsection (1), including:
1236	(a) the name and contact information of the payment provider;
1237	(b) the nature of the relationship between the prospective licensee and the payment
1238	provider;
1239	(c) the processes of the following to safely and reliably conduct transactions for medical
1240	cannabis shipments:
1241	(i) the prospective licensee; and
1242	(ii) the electronic payment provider or the financial institution described in
1243	Subsection (1)(c); and
1244	(d) the ability of the licensee to comply with the department's rules regarding the secure
1245	transportation and delivery of medical cannabis [or medical cannabis product] to a
1246	medical cannabis cardholder.
1247	(3) Notwithstanding any county or municipal ordinance, a medical cannabis pharmacy that
1248	the department designates as a home delivery medical cannabis pharmacy may deliver
1249	medical cannabis shipments in accordance with this part.
1250	Section 19. Section 4-41a-1202 is amended to read:
1251	4-41a-1202 . Home delivery of medical cannabis shipments Medical cannabis
1252	couriers License

1286

1253	(1) The department shall make rules, in accordance with Title 63G, Chapter 3, Utah
1254	Administrative Rulemaking Act, to ensure the safety, security, and efficiency of a home
1255	delivery medical cannabis pharmacy's fulfillment of electronic medical cannabis orders[
1256	that the state central patient portal facilitates], including rules regarding the safe and
1257	controlled delivery of medical cannabis shipments.
1258	(2) A person may not operate as a medical cannabis courier without a license that the
1259	department issues under this section.
1260	(3)(a) Subject to Subsections (5) and (6), the department shall issue a license to operate
1261	as a medical cannabis courier to an applicant who is eligible for a license under this
1262	section.
1263	(b) An applicant is eligible for a license under this section if the applicant submits to the
1264	department:
1265	(i) the name and address of an individual who:
1266	(A) has a financial or voting interest of 10% or greater in the proposed medical
1267	cannabis courier; or
1268	(B) has the power to direct or cause the management or control of a proposed
1269	cannabis production establishment;
1270	(ii) an operating plan that includes operating procedures to comply with the operating
1271	requirements for a medical cannabis courier described in this chapter; and
1272	(iii) an application fee in an amount that, subject to Subsection 4-41a-104(5), the
1273	department sets in accordance with Section 63J-1-504.
1274	(4) If the department determines that an applicant is eligible for a license under this section,
1275	the department shall:
1276	(a) charge the applicant an initial license fee in an amount that, subject to Subsection
1277	4-41a-104(5), the department sets in accordance with Section 63J-1-504; and
1278	(b) notify the Department of Public Safety of the license approval and the names of each
1279	individual described in Subsection (3)(b)(i).
1280	(5) The department may not issue a license to operate as a medical cannabis courier to an
1281	applicant if an individual described in Subsection (3)(b)(i):
1282	(a) has been convicted under state or federal law of:
1283	(i) a felony in the preceding 10 years; or
1284	(ii) after September 23, 2019, a misdemeanor for drug distribution; or
1285	(b) is younger than 21 years old.

(6) The department may revoke a license under this part if:

1287	(a) the medical cannabis courier does not begin operations within one year after the day
1288	on which the department issues the initial license;
1289	(b) the medical cannabis courier makes the same violation of this chapter three times;
1290	(c) an individual described in Subsection (3)(b)(i) is convicted, while the license is
1291	active, under state or federal law of:
1292	(i) a felony; or
1293	(ii) after September 23, 2019, a misdemeanor for drug distribution; or
1294	(d) after a change of ownership described in Subsection (14)(c), the department
1295	determines that the medical cannabis courier no longer meets the minimum standards
1296	for licensure and operation of the medical cannabis courier described in this chapter.
1297	(7) The department shall deposit the proceeds of a fee imposed by this section in the
1298	Qualified Production Enterprise Fund.
1299	(8) The department's authority to issue a license under this section is plenary and is not
1300	subject to review.
1301	(9) Each applicant for a license as a medical cannabis courier shall submit, at the time of
1302	application, from each individual who has a financial or voting interest of 10% or
1303	greater in the applicant or who has the power to direct or cause the management or
1304	control of the applicant:
1305	(a) a fingerprint card in a form acceptable to the Department of Public Safety;
1306	(b) a signed waiver in accordance with Subsection 53-10-108(4) acknowledging the
1307	registration of the individual's fingerprints in the Federal Bureau of Investigation
1308	Next Generation Identification System's Rap Back Service; and
1309	(c) consent to a fingerprint background check by:
1310	(i) the Bureau of Criminal Identification; and
1311	(ii) the Federal Bureau of Investigation.
1312	(10) The Bureau of Criminal Identification shall:
1313	(a) check the fingerprints the applicant submits under Subsection (9) against the
1314	applicable state, regional, and national criminal records databases, including the
1315	Federal Bureau of Investigation Next Generation Identification System;
1316	(b) report the results of the background check to the department;
1317	(c) maintain a separate file of fingerprints that applicants submit under Subsection (9)
1318	for search by future submissions to the local and regional criminal records databases,
1319	including latent prints;
1320	(d) request that the fingerprints be retained in the Federal Bureau of Investigation Next

1321	Generation Identification System's Rap Back Service for search by future	
1322	submissions to national criminal records databases, including the Next Ge	eneration
1323	Identification System and latent prints; and	
1324	(e) establish a privacy risk mitigation strategy to ensure that the department of	nly
1325	receives notifications for an individual with whom the department mainta	ins an
1326	authorizing relationship.	
1327	(11) The department shall:	
1328	(a) assess an individual who submits fingerprints under Subsection (9) a fee i	n an
1329	amount that the department sets in accordance with Section 63J-1-504 for	the
1330	services that the Bureau of Criminal Identification or another authorized a	igency
1331	provides under this section; and	
1332	(b) remit the fee described in Subsection (11)(a) to the Bureau of Criminal Id	entification
1333	(12) The department shall renew a license under this section every year if, at the t	ime of
1334	renewal:	
1335	(a) the licensee meets the requirements of this section; and	
1336	(b) the licensee pays the department a license renewal fee in an amount that,	subject to
1337	Subsection 4-41a-104(5), the department sets in accordance with Section	63J-1-504.
1338	(13) A person applying for a medical cannabis courier license shall submit to the	
1339	department a proposed operating plan that complies with this section and that	includes:
1340	(a) a description of the physical characteristics of any proposed facilities, incl	uding a
1341	floor plan and an architectural elevation, and delivery vehicles;	
1342	(b) a description of the credentials and experience of each officer, director, or	owner of
1343	the proposed medical cannabis courier;	
1344	(c) the medical cannabis courier's employee training standards;	
1345	(d) a security plan; and	
1346	(e) storage and delivery protocols, both short and long term, to ensure that me	edical
1347	cannabis shipments are stored and delivered in a manner that is sanitary a	nd
1348	preserves the integrity of the cannabis.	
1349	(14)(a) A medical cannabis courier license is not transferable or assignable.	
1350	(b) A medical cannabis courier shall report in writing to the department no la	ter than 10
1351	business days before the date of any change of ownership of the medical of	cannabis
1352	courier.	
1353	(c) If the ownership of a medical cannabis courier changes by 50% or more:	
1354	(i) concurrent with the report described in Subsection (14)(b), the medical	l cannabis

1355	courier shall submit a new application described in Subsection (3)(b);
1356	(ii) within 30 days of the submission of the application, the department shall:
1357	(A) conduct an application review; and
1358	(B) award a license to the medical cannabis courier for the remainder of the term
1359	of the medical cannabis courier's license before the ownership change if the
1360	medical cannabis courier meets the minimum standards for licensure and
1361	operation of the medical cannabis courier described in this chapter; and
1362	(iii) if the department approves the license application, notwithstanding Subsection
1363	(4), the medical cannabis courier shall pay a license fee that the department sets in
1364	accordance with Section 63J-1-504 in an amount that covers the board's cost of
1365	conducting the application review.
1366	(15)(a) Except as provided in Subsection(15)(b), a person may not advertise regarding
1367	the transportation of medical cannabis.
1368	(b) Notwithstanding Subsection (14)(a) and subject to Section 4-41a-109, a licensed
1369	home delivery medical cannabis pharmacy or a licensed medical cannabis courier
1370	may advertise:
1371	(i) a green cross;
1372	(ii) the pharmacy's or courier's name and logo; and
1373	(iii) that the pharmacy or courier is licensed to transport medical cannabis shipments.
1374	Section 20. Section 4-41a-1203 is amended to read:
1375	4-41a-1203. Medical cannabis shipment transportation.
1376	(1) The department shall ensure that each home delivery medical cannabis pharmacy is
1377	capable of delivering, directly or through a medical cannabis courier, medical cannabis
1378	shipments in a secure manner.
1379	(2)(a) A home delivery medical cannabis pharmacy may contract with a licensed
1380	medical cannabis courier to deliver medical cannabis shipments to fulfill electronic
1381	medical cannabis orders[-that the state central patient portal facilitates].
1382	(b) If a home delivery medical cannabis pharmacy enters into a contract described in
1383	Subsection (2)(a), the pharmacy shall:
1384	(i) impose security and personnel requirements on the medical cannabis courier
1385	sufficient to ensure the security and safety of medical cannabis shipments; and
1386	(ii) provide regular oversight of the medical cannabis courier.
1387	(3) Notwithstanding Subsection 4-41a-404(1), an individual may transport a medical
1388	cannabis shipment if the individual is:

1389	(a) a registered pharmacy medical provider;
1390	(b) a registered medical cannabis pharmacy agent; or
1391	(c) a registered agent of the medical cannabis courier described in Subsection (2).
1392	(4) An individual transporting a medical cannabis shipment under Subsection (3) shall
1393	comply with the requirements of Subsection 4-41a-404(3).
1394	(5) In addition to the requirements in Subsections (3) and (4), the department may establish
1395	by rule, in collaboration with the Division of Professional Licensing and the Board of
1396	Pharmacy and in accordance with Title 63G, Chapter 3, Utah Administrative
1397	Rulemaking Act, requirements for transporting medical cannabis shipments that are
1398	related to safety for human consumption of [eannabis or a cannabis product] medical
1399	<u>cannabis</u> .
1400	(6)(a) It is unlawful for an individual to transport a medical cannabis shipment with a
1401	manifest that does not meet the requirements of Subsection (4).
1402	(b) Except as provided in Subsection (6)(d), an individual who violates Subsection (6)(a)
1403	is:
1404	(i) guilty of an infraction; and
1405	(ii) subject to a \$100 fine.
1406	(c) An individual who is guilty of a violation described in Subsection (6)(b) is not guilty
1407	of a violation of Title 58, Chapter 37, Utah Controlled Substances Act, for the
1408	conduct underlying the violation described in Subsection (6)(b).
1409	(d) If the individual described in Subsection (6)(a) is transporting more cannabis,
1410	cannabis product, or medical cannabis devices than the manifest identifies, except for
1411	a de minimis administrative error:
1412	(i) this chapter does not apply; and
1413	(ii) the individual is subject to penalties under Title 58, Chapter 37, Utah Controlled
1414	Substances Act.
1415	Section 21. Section 4-41a-1206 is amended to read:
1416	4-41a-1206. Closed-door medical cannabis pharmacy.
1417	(1)(a) Subject to Subsections (1)(b) and (c), a home delivery medical cannabis pharmacy
1418	may open a single closed-door medical cannabis pharmacy.
1419	(b) A home delivery medical cannabis pharmacy may not open a closed-door medical
1420	cannabis pharmacy unless the home delivery medical cannabis pharmacy:
1421	(i) has an operating plan that includes a closed-door medical cannabis pharmacy; and
1422	(ii) obtains a license issued by the department for a closed-door medical cannabis

1423	pharmacy.
1424	(c) An entity that owns multiple home delivery medical cannabis pharmacies may open
1425	only one closed-door medical cannabis pharmacy.
1426	(d) The department may institute a fee in accordance with Section 63J-1-504 to
1427	administer this section.
1428	(2) A home delivery medical cannabis pharmacy that opens a closed-door medical cannabis
1429	pharmacy under Subsection (1) shall ensure:
1430	(a) that a pharmacy medical provider who is a licensed pharmacist:
1431	(i) is directly supervising the packaging of an order; and
1432	(ii) is present in the closed-door medical cannabis pharmacy when an order is
1433	packaged for delivery; and
1434	(b) all record keeping requirements, labeling requirements, and patient counseling
1435	requirements described in this chapter and Title 26B, Chapter 4, Part 2, Cannabinoid
1436	Research and Medical Cannabis, are satisfied before sending out an order.
1437	(3) An individual who prepares an order at a closed-door medical cannabis pharmacy under
1438	this section shall be registered as:
1439	(a) a pharmacy medical provider; or
1440	(b) a medical cannabis pharmacy agent.
1441	(4)(a) A closed-door medical cannabis pharmacy shall operate:
1442	(i) except as provided in Subsection (4)(b), in a facility that is accessible only by an
1443	individual who is a pharmacy medical provider or a medical cannabis pharmacy
1444	agent; and
1445	(ii) at a physical address in accordance with Subsection (6).
1446	(b) A closed-door medical cannabis pharmacy may authorize an individual who is at
1447	least 18 years old and is not a pharmacy medical provider or a cannabis pharmacy
1448	agent to access the closed-door medical cannabis pharmacy if the closed-door
1449	medical cannabis pharmacy:
1450	(i) tracks and monitors the individual at all times while the individual is at the
1451	closed-door medical cannabis pharmacy; and
1452	(ii) maintains a record of the individual's access, including arrival and departure.
1453	(c) A closed-door medical cannabis pharmacy shall operate in a facility that has:
1454	(i) a single, secure public entrance; and
1455	(ii) a security system with a backup power source that:
1456	(A) detects and records entry into the closed-door medical cannabis pharmacy;

1457	(B) provides notice of an unauthorized entry to law enforcement when the
1458	closed-door medical cannabis pharmacy is closed; and
1459	(C) a lock or equivalent restrictive security feature on any area where the
1460	closed-door medical cannabis pharmacy stores a cannabis product.
1461	(d) A closed-door medical cannabis pharmacy shall ensure that any cannabis or cannabis
1462	products in the closed-door medical cannabis pharmacy that are intended for home
1463	delivery are separated in a manner that is readily distinguishable from any other
1464	cannabis or cannabis product in the facility.
1465	(5) A closed-door medical cannabis pharmacy may only provide cannabis or a cannabis
1466	product to an individual through a delivery that complies with this part.
1467	(6)(a) A person may not locate a closed-door medical cannabis pharmacy:
1468	(i) within 1,000 feet of a community location; or
1469	(ii) in or within 600 feet of a district that the relevant municipality or county has
1470	zoned as primarily residential.
1471	(b) The proximity requirements described in Subsection (6)(a) shall be measured from
1472	the nearest entrance to the closed-door medical cannabis pharmacy by following the
1473	shortest route of ordinary pedestrian travel to the property boundary of the
1474	community location or residential area.
1475	(c) The licensing board may grant a waiver to reduce the proximity requirements in
1476	Subsection (6)(a) by up to 20% if the licensing board determines that it is not
1477	reasonably feasible for the applicant to site the proposed closed-door medical
1478	cannabis pharmacy without the waiver.
1479	(d) An applicant for a license under this section shall provide evidence of compliance
1480	with the proximity requirements described in Subsection (6)(a).
1481	(7) When determining where a closed-door medical cannabis pharmacy may open, the
1482	licensing board:
1483	(a) shall utilize geographic regions created by the department through rule;
1484	(b) shall prioritize allowing entities that do not have a medical cannabis pharmacy in a
1485	region to open a closed-door medical cannabis pharmacy in the region;
1486	(c) of the total amount of closed-door medical cannabis pharmacies, may allow only
1487	three closed-door medical cannabis pharmacies to operate in counties of the first and
1488	second class as described in Section 17-50-501; and
1489	(d) for determining the three closed-door medical cannabis pharmacies described in
1490	Subsection (7)(c), consider the following:

1491	(i) the history of compliance with state law and rules for all licenses issued under this
1492	chapter;
1493	(ii) the medical cannabis pharmacy's willingness to offer a variety of brands and
1494	products;
1495	(iii) the ability of the operating plan to ensure the safety and security of the
1496	community;
1497	(iv) the suitability of the proposed location and the location's ability to serve the local
1498	community; and
1499	(v) any other relevant information determined through rule.
1500	(8) A closed-door medical cannabis pharmacy may not account for more than:
1501	(a) for an entity that holds a single medical cannabis pharmacy license, the greater of:
1502	(i) 35% of the medical cannabis pharmacy's total revenue; or
1503	(ii) \$2,000,000 in total revenue; or
1504	(b) for an entity that holds more than one medical cannabis pharmacy license, the greater
1505	of:
1506	(i) 35% of the total revenue of the entity's medical cannabis pharmacy that generates
1507	the most revenue; or
1508	(ii) \$2,000,000 in total revenue.
1509	(9) Notwithstanding any other provision of this section, the [department] licensing board
1510	may issue only [three] one closed-door medical cannabis pharmacy [licenses] license
1511	before July 1, 2027.
1512	(10) In accordance with Title 63G, Chapter 3, Utah Administrative Rulemaking Act, the
1513	department shall make rules to implement this section.
1514	Section 22. Section 26B-1-435 is amended to read:
1515	26B-1-435 . Medical Cannabis Policy Advisory Board creation Membership
1516	Duties.
1517	(1) There is created within the department the Medical Cannabis Policy Advisory Board.
1518	(2)(a) The advisory board shall consist of the following members:
1519	(i) appointed by the executive director:
1520	(A) a qualified medical provider who has recommended medical cannabis to at
1521	least 100 patients before being appointed;
1522	[(B) a medical research professional;]
1523	[(C)] (B) a mental health specialist;
1524	[(D)] (C) an individual who represents an organization that advocates for medical

1525	cannabis patients;
1526	[(E)] (D) [an individual] a member of the general public who holds a medical
1527	cannabis patient card; and
1528	[(F)] (E) a member of the general public who does not hold a medical cannabis
1529	card;[-and]
1530	(ii) appointed by the commissioner of the Department of Agriculture and Food:
1531	(A) an individual who owns or operates a licensed cannabis cultivation facility, as
1532	defined in Section 4-41a-102;
1533	(B) an individual who owns or operates a licensed medical cannabis pharmacy;
1534	and
1535	(C) a law enforcement officer[-]; and
1536	(iii) a representative from the Center for Medical Cannabis Research created in
1537	Section 53B-14-1402, appointed by the Center for Medical Cannabis Research.
1538	(b) The commissioner of the Department of Agriculture and Food shall ensure that at
1539	least one individual appointed under Subsection (2)(a)(ii)(A) or (B) also owns or
1540	operates a licensed cannabis processing facility.
1541	(3)(a) Subject to Subsection (3)(b), a member of the advisory board shall serve for a four
1542	year term.
1543	(b) When appointing the initial membership of the advisory board, the executive director
1544	and the commissioner of the Department of Agriculture and Food shall coordinate to
1545	appoint four advisory board members to serve a term of two years to ensure that
1546	approximately half of the board is appointed every two years.
1547	(4)(a) If an advisory board member is no longer able to serve as a member, a new
1548	member shall be appointed in the same manner as the original appointment.
1549	(b) A member appointed in accordance with Subsection (4)(a) shall serve for the
1550	remainder of the unexpired term of the original appointment.
1551	(5)(a) A majority of the advisory board members constitutes a quorum.
1552	(b) The action of a majority of a quorum constitutes an action of the advisory board.
1553	(c) For a term lasting one year, the advisory board shall annually designate members of
1554	the advisory board to serve as chair and vice-chair.
1555	(d) When designating the chair and vice-chair, the advisory board shall ensure that at
1556	least one individual described Subsection (2)(a)(i) is appointed as chair or vice-chair.
1557	(6) An advisory board member may not receive compensation or benefits for the member's
1558	service on the advisory board but may receive per diem and reimbursement for travel

1559		expenses incurred as an advisory board member in accordance with:
1560		(a) Sections 63A-3-106 and 63A-3-107; and
1561		(b) rules made by the Division of Finance pursuant to Sections 63A-3-106 and
1562		63A-3-107.
1563	(7)	The department shall:
1564		(a) provide staff support for the advisory board; and
1565		(b) assist the advisory board in conducting meetings.
1566	(8)	The advisory board may recommend:
1567		(a) to the department or the Department of Agriculture and Food changes to current or
1568		proposed medical cannabis rules or statutes; and
1569		(b) to the appropriate legislative committee whether the advisory board supports a
1570		change to medical cannabis statutes.
1571	(9)	The advisory board shall:
1572		(a) review any draft rule that is authorized under [this chapter] Chapter 4, Part 2,
1573		Cannabinoid Research and Medical Cannabis, or Title 4, Chapter 41a, Cannabis
1574		Production Establishments and Pharmacies;
1575		(b) consult with the Department of Agriculture and Food regarding the issuance of an
1576		additional:
1577		(i) cultivation facility license under Section 4-41a-205; or
1578		(ii) pharmacy license under Section 4-41a-1005;
1579		(c) consult with the department regarding cannabis patient education;
1580		(d) consult regarding the reasonableness of any fees set by the department or the
1581		Department of Agriculture and Food that pertain to the medical cannabis program;
1582		and
1583		(e) consult regarding any issue pertaining to medical cannabis when asked by the
1584		department or the Utah Department of Agriculture and Food.
1585		Section 23. Section 26B-4-201 is amended to read:
1586		26B-4-201 . Definitions.
1587		As used in this part:
1588	(1)	"Active tetrahydrocannabinol" means THC, any THC analog, and
1589		tetrahydrocannabinolic acid.
1590	(2)	"Administration of criminal justice" means the performance of detection, apprehension,
1591		detention, pretrial release, post-trial release, prosecution, and adjudication.
1592	(3)	"Advertise" means information provided by a person in any medium:

1593	(a) to the public; and
1594	(b) that is not age restricted to an individual who is at least 21 years old.
1595	(4) "Advisory board" means the Medical Cannabis Policy Advisory Board created in
1596	Section 26B-1-435.
1597	(5) "Cannabis Research Review Board" means the Cannabis Research Review Board
1598	created in Section 26B-1-420.
1599	(6) "Cannabis" means marijuana.
1600	(7) "Cannabis processing facility" means the same as that term is defined in Section
1601	4-41a-102.
1602	(8) "Cannabis product" means a product that:
1603	(a) is intended for human use; and
1604	(b) contains cannabis or any tetrahydrocannabinol or THC analog in a total
1605	concentration of 0.3% or greater on a dry weight basis.
1606	(9) "Cannabis production establishment" means the same as that term is defined in Section
1607	4-41a-102.
1608	(10) "Cannabis production establishment agent" means the same as that term is defined in
1609	Section 4-41a-102.
1610	(11) "Cannabis production establishment agent registration card" means the same as that
1611	term is defined in Section 4-41a-102.
1612	(12) "Conditional medical cannabis card" means an electronic medical cannabis card that
1613	the department issues in accordance with Subsection 26B-4-213(1)(b) to allow an
1614	applicant for a medical cannabis card to access medical cannabis during the department's
1615	review of the application.
1616	(13) "Controlled substance database" means the controlled substance database created in
1617	Section 58-37f-201.
1618	(14) "Delivery address" means the same as that term is defined in Section 4-41a-102.
1619	(15) "Department" means the Department of Health and Human Services.
1620	(16) "Designated caregiver" means:
1621	(a) an individual:
1622	(i) whom an individual with a medical cannabis patient card or a medical cannabis
1623	guardian card designates as the patient's caregiver; and
1624	(ii) who registers with the department under Section 26B-4-214; or
1625	(b)(i) a facility that an individual designates as a designated caregiver in accordance
1626	with Subsection 26B-4-214(1)(b); or

1627	(ii) an assigned employee of the facility described in Subsection 26B-4-214(1)(b)(ii).
1628	(17) "Directions of use" means recommended routes of administration for a medical
1629	cannabis treatment and suggested usage guidelines.
1630	(18) "Dosing guidelines" means a quantity range and frequency of administration for a
1631	recommended treatment of medical cannabis.
1632	(19) "Government issued photo identification" means any of the following forms of
1633	identification:
1634	(a) a valid state-issued driver license or identification card;
1635	(b) a valid United States federal-issued photo identification, including:
1636	(i) a United States passport;
1637	(ii) a United States passport card;
1638	(iii) a United States military identification card; or
1639	(iv) a permanent resident card or alien registration receipt card; or
1640	(c) a foreign passport.
1641	(20) "Home delivery medical cannabis pharmacy" means a medical cannabis pharmacy that
1642	the department authorizes, as part of the pharmacy's license, to deliver medical cannabis
1643	shipments to a delivery address to fulfill electronic orders[-that the state central patient
1644	portal facilitates].
1645	(21) "Inventory control system" means the system described in Section 4-41a-103.
1646	(22) "Legal dosage limit" means an amount that:
1647	(a) is sufficient to provide 30 days of treatment based on the dosing guidelines that the
1648	relevant recommending medical provider or [the state central patient portal or]
1649	pharmacy medical provider, in accordance with Subsection 26B-4-230(5),
1650	recommends; and
1651	(b) may not exceed:
1652	(i) for unprocessed cannabis in a medicinal dosage form, 113 grams by weight; and
1653	(ii) for a cannabis product in a medicinal dosage form, a quantity that contains, in
1654	total, greater than 20 grams of active tetrahydrocannabinol.
1655	(23) "Legal use termination date" means a date on the label of a container of unprocessed
1656	cannabis flower:
1657	(a) that is 60 days after the date of purchase of the cannabis; and
1658	(b) after which, the cannabis is no longer in a medicinal dosage form outside of the
1659	primary residence of the relevant medical cannabis patient cardholder.
1660	(24) "Limited medical provider" means an individual who:

1661	(a) meets the recommending qualifications; and
1662	(b) has no more than 15 patients with a valid medical cannabis patient card as a result of
1663	the individual's recommendation, in accordance with Subsection 26B-4-204(1)(b).
1664	(25) "Marijuana" means the same as that term is defined in Section 58-37-2.
1665	(26) "Medical cannabis" or "medical cannabis product" means cannabis in a medicinal
1666	dosage form or a cannabis product in a medicinal dosage form.
1667	(27) "Medical cannabis card" means a medical cannabis patient card, a medical cannabis
1668	guardian card, a medical cannabis caregiver card, or a conditional medical cannabis card.
1669	(28) "Medical cannabis cardholder" means:
1670	(a) a holder of a medical cannabis card; or
1671	(b) a facility or assigned employee, described in Subsection (16)(b), only:
1672	(i) within the scope of the facility's or assigned employee's performance of the role of
1673	a medical cannabis patient cardholder's caregiver designation under Subsection
1674	26B-4-214(1)(b); and
1675	(ii) while in possession of documentation that establishes:
1676	(A) a caregiver designation described in Subsection 26B-4-214(1)(b);
1677	(B) the identity of the individual presenting the documentation; and
1678	(C) the relation of the individual presenting the documentation to the caregiver
1679	designation.
1680	(29) "Medical cannabis caregiver card" means an electronic document that a cardholder
1681	may print or store on an electronic device or a physical card or document that:
1682	(a) the department issues to an individual whom a medical cannabis patient cardholder
1683	or a medical cannabis guardian cardholder designates as a designated caregiver; and
1684	(b) is connected to the electronic verification system.
1685	(30) "Medical cannabis courier" means the same as that term is defined in Section
1686	4-41a-102.
1687	(31)(a) "Medical cannabis device" means a device that an individual uses to ingest or
1688	inhale [cannabis in a medicinal dosage form or a cannabis product in a medicinal
1689	dosage form] medical cannabis.
1690	(b) "Medical cannabis device" does not include a device that:
1691	(i) facilitates cannabis combustion; or
1692	(ii) an individual uses to ingest substances other than cannabis.
1693	(32) "Medical cannabis guardian card" means an electronic document that a cardholder may
1694	print or store on an electronic device or a physical card or document that:

1695	(a) the department issues to the parent or legal guardian of a minor with a qualifying
1696	condition; and
1697	(b) is connected to the electronic verification system.
1698	(33) "Medical cannabis patient card" means an electronic document that a cardholder may
1699	print or store on an electronic device or a physical card or document that:
1700	(a) the department issues to an individual with a qualifying condition; and
1701	(b) is connected to the electronic verification system.
1702	(34) "Medical cannabis pharmacy" means a person that:
1703	(a)(i) acquires or intends to acquire medical cannabis [or a cannabis product in a
1704	medicinal dosage form-]from a cannabis processing facility or another medical
1705	cannabis pharmacy or a medical cannabis device; or
1706	(ii) possesses medical cannabis or a medical cannabis device; and
1707	(b) sells or intends to sell medical cannabis or a medical cannabis device to a medical
1708	cannabis cardholder.
1709	(35) "Medical cannabis pharmacy agent" means an individual who holds a valid medical
1710	cannabis pharmacy agent registration card issued by the department.
1711	(36) "Medical cannabis pharmacy agent registration card" means a registration card issued
1712	by the department that authorizes an individual to act as a medical cannabis pharmacy
1713	agent.
1714	(37) "Medical cannabis shipment" means the same as that term is defined in Section
1715	4-41a-102.
1716	(38) "Medical cannabis treatment" means [eannabis in a medicinal dosage form, a cannabis
1717	product in a medicinal dosage form, or] medical cannabis or a medical cannabis device.
1718	(39)(a) "Medicinal dosage form" means:
1719	(i) for processed medical cannabis, the following with a specific and consistent
1720	cannabinoid content:
1721	(A) a tablet;
1722	(B) a capsule;
1723	(C) a concentrated liquid or viscous oil;
1724	(D) a liquid suspension that does not exceed 30 milliliters;
1725	(E) a topical preparation;
1726	(F) a transdermal preparation;
1727	(G) a sublingual preparation;
1728	(H) a gelatinous cube, gelatinous rectangular cuboid, or lozenge in a cube or

1729	rectangular cuboid shape;
1730	(I) a resin or wax;
1731	(J) an aerosol;
1732	(K) a suppository preparation; or
1733	(L) a soft or hard confection that is a uniform rectangular cuboid or uniform
1734	spherical shape, is homogeneous in color and texture, and each piece is a single
1735	serving; or
1736	(ii) for unprocessed cannabis flower, a container described in Section 4-41a-602 that:
1737	(A) contains cannabis flower in a quantity that varies by no more than 10% from
1738	the stated weight at the time of packaging;
1739	(B) at any time the medical cannabis cardholder transports or possesses the
1740	container in public, is contained within an opaque bag or box that the medical
1741	cannabis pharmacy provides; and
1742	(C) is labeled with the container's content and weight, the date of purchase, the
1743	legal use termination date, and a barcode that provides information connected
1744	to an inventory control system.
1745	(b) "Medicinal dosage form" includes a portion of unprocessed cannabis flower that:
1746	(i) the medical cannabis cardholder has recently removed from the container
1747	described in Subsection (39)(a)(ii) for use; and
1748	(ii) does not exceed the quantity described in Subsection (39)(a)(ii).
1749	(c) "Medicinal dosage form" does not include:
1750	(i) any unprocessed cannabis flower outside of the container described in Subsection
1751	(39)(a)(ii), except as provided in Subsection (39)(b);
1752	(ii) any unprocessed cannabis flower in a container described in Subsection
1753	(39)(a)(ii) after the legal use termination date;
1754	(iii) a process of vaporizing and inhaling concentrated cannabis by placing the
1755	cannabis on a nail or other metal object that is heated by a flame, including a
1756	blowtorch;
1757	(iv) a liquid suspension that is branded as a beverage;
1758	(v) a substance described in Subsection (39)(a)(i) or (ii) if the substance is not
1759	measured in grams, milligrams, or milliliters; or
1760	(vi) a substance that contains or is covered to any degree with chocolate.
1761	(40) "Nonresident patient" means an individual who:
1762	(a) is not a resident of Utah or has been a resident of Utah for less than 45 days;

1763	(b) has a currently valid medical cannabis card or the equivalent of a medical cannabis
1764	card under the laws of another state, district, territory, commonwealth, or insular
1765	possession of the United States; and
1766	(c) has been diagnosed with a qualifying condition as described in Section 26B-4-203.
1767	(41) "Pharmacy medical provider" means the medical provider required to be on site at a
1768	medical cannabis pharmacy under Section 26B-4-219.
1769	(42) "Provisional patient card" means a card that:
1770	(a) the department issues to a minor with a qualifying condition for whom:
1771	(i) a recommending medical provider has recommended a medical cannabis
1772	treatment; and
1773	(ii) the department issues a medical cannabis guardian card to the minor's parent or
1774	legal guardian; and
1775	(b) is connected to the electronic verification system.
1776	(43) "Qualified medical provider" means an individual:
1777	(a) who meets the recommending qualifications; and
1778	(b) whom the department registers to recommend treatment with cannabis in a medicinal
1779	dosage form under Section 26B-4-204.
1780	(44) "Qualified Patient Enterprise Fund" means the enterprise fund created in Section
1781	26B-1-310.
1782	(45) "Qualifying condition" means a condition described in Section 26B-4-203.
1783	(46) "Recommend" or "recommendation" means, for a recommending medical provider, the
1784	act of suggesting the use of medical cannabis treatment, which:
1785	(a) certifies the patient's eligibility for a medical cannabis card; and
1786	(b) may include, at the recommending medical provider's discretion, directions of use,
1787	with or without dosing guidelines.
1788	(47) "Recommending medical provider" means a qualified medical provider or a limited
1789	medical provider.
1790	(48) "Recommending qualifications" means that an individual:
1791	(a)(i) has the authority to write a prescription;
1792	(ii) is licensed to prescribe a controlled substance under Title 58, Chapter 37, Utah
1793	Controlled Substances Act; and
1794	(iii) possesses the authority, in accordance with the individual's scope of practice, to
1795	prescribe a Schedule II controlled substance; and
1796	(b) is licensed as:

1/9/	(1) a podiatrist under Title 58, Chapter 5a, Podiatric Physician Licensing Act;
1798	(ii) an advanced practice registered nurse under Title 58, Chapter 31b, Nurse Practice
1799	Act;
1800	(iii) a physician under Title 58, Chapter 67, Utah Medical Practice Act, or Title 58,
1801	Chapter 68, Utah Osteopathic Medical Practice Act; or
1802	(iv) a physician assistant under Title 58, Chapter 70a, Utah Physician Assistant Act.
1803	[(49) "State central patient portal" means the website the department creates, in accordance
1804	with Section 26B-4-236, to facilitate patient safety, education, and an electronic medical
1805	eannabis order.]
1806	[(50)] (49) "State electronic verification system" means the system described in Section
1807	26B-4-202.
1808	[(51)] (50) "Targeted marketing" means the promotion by a qualified medical provider,
1809	medical clinic, or medical office that employs a qualified medical provider of a medical
1810	cannabis recommendation service using any of the following methods:
1811	(a) electronic communication to an individual who is at least 21 years old and has
1812	requested to receive promotional information;
1813	(b) an in-person marketing event that is held in an area where only an individual who is
1814	at least 21 years old may access the event;
1815	(c) other marketing material that is physically or digitally displayed in the office of the
1816	medical clinic or office that employs a qualified medical provider; or
1817	(d) a leaflet that a qualified medical provider, medical clinic, or medical office that
1818	employs a qualified medical provider shares with an individual who is at least 21
1819	years old.
1820	[(52)] (51) "Tetrahydrocannabinol" or "THC" means a substance derived from cannabis or a
1821	synthetic equivalent as described in Subsection 58-37-4(2)(a)(iii)(AA).
1822	[(53)] (52) "THC analog" means the same as that term is defined in Section 4-41-102.
1823	Section 24. Section 26B-4-202 is amended to read:
1824	26B-4-202 . Electronic verification system.
1825	(1) The Department of Agriculture and Food, the department, the Department of Public
1826	Safety, and the Division of Technology Services shall:
1827	(a) enter into a memorandum of understanding in order to determine the function and
1828	operation of the state electronic verification system in accordance with Subsection
1829	(2);
1830	(b) coordinate with the Division of Purchasing, under Title 63G, Chapter 6a, Utah

1831	Procurement Code, to develop a request for proposals for a third-party provider to
1832	develop and maintain the state electronic verification system in coordination with the
1833	Division of Technology Services; and
1834	(c) select a third-party provider who:
1835	(i) meets the requirements contained in the request for proposals issued under
1836	Subsection (1)(b); and
1837	(ii) may not have any commercial or ownership interest in a cannabis production
1838	establishment or a medical cannabis pharmacy.
1839	(2) The Department of Agriculture and Food, the department, the Department of Public
1840	Safety, and the Division of Technology Services shall ensure that the state electronic
1841	verification system described in Subsection (1):
1842	(a) allows an individual to apply for a medical cannabis patient card or, if applicable, a
1843	medical cannabis guardian card, provided that the card may not become active until:
1844	(i) the relevant qualified medical provider completes the associated medical cannabis
1845	recommendation; or
1846	(ii) for a medical cannabis card related to a limited medical provider's
1847	recommendation, the medical cannabis pharmacy completes the recording
1848	described in Subsection (2)(d);
1849	(b) allows an individual to apply to renew a medical cannabis patient card or a medical
1850	cannabis guardian card in accordance with Section 26B-4-213;
1851	(c) allows a qualified medical provider, or an employee described in Subsection (3)
1852	acting on behalf of the qualified medical provider, to:
1853	(i) access dispensing and card status information regarding a patient:
1854	(A) with whom the qualified medical provider has a provider-patient relationship;
1855	and
1856	(B) for whom the qualified medical provider has recommended or is considering
1857	recommending a medical cannabis card;
1858	(ii) electronically recommend treatment with [eannabis in a medicinal dosage form or
1859	a cannabis product in a medicinal dosage form] medical cannabis and optionally
1860	recommend dosing guidelines;
1861	(iii) electronically renew a recommendation to a medical cannabis patient cardholder
1862	or medical cannabis guardian cardholder:
1863	(A) using telehealth services, for the qualified medical provider who originally
1864	recommended a medical cannabis treatment during a face-to-face visit with the

1865	patient; or
1866	(B) during a face-to-face visit with the patient, for a qualified medical provider
1867	who did not originally recommend the medical cannabis treatment during a
1868	face-to-face visit; and
1869	(iv) submit an initial application, renewal application, or application payment on
1870	behalf of an individual applying for any of the following:
1871	(A) a medical cannabis patient card;
1872	(B) a medical cannabis guardian card; or
1873	(C) a medical cannabis caregiver card;
1874	(d) allows a medical cannabis pharmacy medical provider or medical cannabis pharmacy
1875	agent, in accordance with Subsection 4-41a-1101(10)(a), to:
1876	(i) access the electronic verification system to review the history within the system of
1877	a patient with whom the provider or agent is interacting, limited to read-only
1878	access for medical cannabis pharmacy agents unless the medical cannabis
1879	pharmacy's pharmacist in charge authorizes add and edit access;
1880	(ii) record a patient's recommendation from a limited medical provider, including any
1881	directions of use, dosing guidelines, or caregiver indications from the limited
1882	medical provider;
1883	(iii) record a limited medical provider's renewal of the provider's previous
1884	recommendation; and
1885	(iv) submit an initial application, renewal application, or application payment on
1886	behalf of an individual applying for any of the following:
1887	(A) a medical cannabis patient card;
1888	(B) a medical cannabis guardian card; or
1889	(C) a medical cannabis caregiver card;
1890	(e) connects with:
1891	(i) an inventory control system that a medical cannabis pharmacy uses to track in real
1892	time and archive purchases of any [eannabis in a medicinal dosage form, cannabis
1893	product in a medicinal dosage form,] medical cannabis or a medical cannabis
1894	device, including:
1895	(A) the time and date of each purchase;
1896	(B) the quantity and type of [eannabis, eannabis product,] medical cannabis or
1897	medical cannabis device purchased;
1898	(C) any cannabis production establishment, any medical cannabis pharmacy, or

1899	any medical cannabis courier associated with the [eannabis, eannabis product,]
1900	medical cannabis or medical cannabis device; and
1901	(D) the personally identifiable information of the medical cannabis cardholder
1902	who made the purchase; and
1903	(ii) any commercially available inventory control system that a cannabis production
1904	establishment utilizes in accordance with Section 4-41a-103 to use data that the
1905	Department of Agriculture and Food requires by rule, in accordance with Title
1906	63G, Chapter 3, Utah Administrative Rulemaking Act, from the inventory
1907	tracking system that a licensee uses to track and confirm compliance;
1908	(f) provides access to:
1909	(i) the department to the extent necessary to carry out the department's functions and
1910	responsibilities under this part;
1911	(ii) the Department of Agriculture and Food to the extent necessary to carry out the
1912	functions and responsibilities of the Department of Agriculture and Food under
1913	Title 4, Chapter 41a, Cannabis Production Establishments and Pharmacies; and
1914	(iii) the Division of Professional Licensing to the extent necessary to carry out the
1915	functions and responsibilities related to the participation of the following in the
1916	recommendation and dispensing of medical cannabis:
1917	(A) a podiatrist licensed under Title 58, Chapter 5a, Podiatric Physician Licensing
1918	Act;
1919	(B) a pharmacist licensed under Title 58, Chapter 17b, Pharmacy Practice Act;
1920	(C) an advanced practice registered nurse licensed under Title 58, Chapter 31b,
1921	Nurse Practice Act;
1922	(D) a physician licensed under Title 58, Chapter 67, Utah Medical Practice Act, or
1923	Title 58, Chapter 68, Utah Osteopathic Medical Practice Act; or
1924	(E) a physician assistant licensed under Title 58, Chapter 70a, Utah Physician
1925	Assistant Act;
1926	[(g) provides access to and interaction with the state central patient portal;]
1927	[(h)] (g) communicates dispensing information from a record that a medical cannabis
1928	pharmacy submits to the state electronic verification system under Subsection
1929	4-41a-1102(3)(a)(ii) to the controlled substance database;
1930	[(i)] (h) provides access to state or local law enforcement only to verify the validity of an
1931	individual's medical cannabis card for the administration of criminal justice and
1932	through a database used by law enforcement; and

1933	[(j)] (i) creates a record each time a person accesses the system that identifies the person
1934	who accesses the system and the individual whose records the person accesses.
1935	(3)(a) An employee of a qualified medical provider may access the electronic
1936	verification system for a purpose described in Subsection (2)(c) on behalf of the
1937	qualified medical provider if:
1938	(i) the qualified medical provider has designated the employee as an individual
1939	authorized to access the electronic verification system on behalf of the qualified
1940	medical provider;
1941	(ii) the qualified medical provider provides written notice to the department of the
1942	employee's identity and the designation described in Subsection (3)(a)(i); and
1943	(iii) the department grants to the employee access to the electronic verification
1944	system.
1945	(b) An employee of a business that employs a qualified medical provider may access the
1946	electronic verification system for a purpose described in Subsection (2)(c) on behalf
1947	of the qualified medical provider if:
1948	(i) the qualified medical provider has designated the employee as an individual
1949	authorized to access the electronic verification system on behalf of the qualified
1950	medical provider;
1951	(ii) the qualified medical provider and the employing business jointly provide written
1952	notice to the department of the employee's identity and the designation described
1953	in Subsection (3)(b)(i); and
1954	(iii) the department grants to the employee access to the electronic verification
1955	system.
1956	(4)(a) As used in this Subsection (4), "prescribing provider" means:
1957	(i) a podiatrist licensed under Title 58, Chapter 5a, Podiatric Physician Licensing Act
1958	(ii) an advanced practice registered nurse licensed under Title 58, Chapter 31b, Nurse
1959	Practice Act;
1960	(iii) a physician licensed under Title 58, Chapter 67, Utah Medical Practice Act, or
1961	Title 58, Chapter 68, Utah Osteopathic Medical Practice Act; or
1962	(iv) a physician assistant licensed under Title 58, Chapter 70a, Utah Physician
1963	Assistant Act.
1964	(b) A prescribing provider may access information in the electronic verification system
1965	regarding a patient the prescribing provider treats.
1966	(5) The department may release limited data that the system collects for the purpose of:

1967	(a) conducting medical and other department approved research;
1968	(b) providing the report required by Section 26B-4-222; and
1969	(c) other official department purposes.
1970	(6) The department shall make rules, in accordance with Title 63G, Chapter 3, Utah
1971	Administrative Rulemaking Act, to establish:
1972	(a) the limitations on access to the data in the state electronic verification system as
1973	described in this section; and
1974	(b) standards and procedures to ensure accurate identification of an individual requesting
1975	information or receiving information in this section.
1976	(7) Any person who negligently or recklessly releases any information in the state
1977	electronic verification system in violation of this section is guilty of a class C
1978	misdemeanor.
1979	(8) Any person who obtains or attempts to obtain information from the state electronic
1980	verification system by misrepresentation or fraud is guilty of a third degree felony.
1981	(9)(a) Except as provided in Subsections (9)(c) and (9)(e), a person may not knowingly
1982	and intentionally use, release, publish, or otherwise make available to any other
1983	person information obtained from the state electronic verification system for any
1984	purpose other than a purpose specified in this section.
1985	(b) Each separate violation of this Subsection (9) is:
1986	(i) a third degree felony; and
1987	(ii) subject to a civil penalty not to exceed \$5,000.
1988	(c) A law enforcement officer who uses the database used by law enforcement to access
1989	information in the electronic verification system for a reason that is not the
1990	administration of criminal justice is guilty of a class B misdemeanor.
1991	(d) The department shall determine a civil violation of this Subsection (9) in accordance
1992	with Title 63G, Chapter 4, Administrative Procedures Act.
1993	(e) Civil penalties assessed under this Subsection (9) shall be deposited into the General
1994	Fund.
1995	(f) This Subsection (9) does not prohibit a person who obtains information from the state
1996	electronic verification system under Subsection (2)(a), (c), or (f) from:
1997	(i) including the information in the person's medical chart or file for access by a
1998	person authorized to review the medical chart or file;
1999	(ii) providing the information to a person in accordance with the requirements of the
2000	Health Insurance Portability and Accountability Act of 1996; or

2001	(iii) discussing or sharing that information about the patient with the patient.
2002	Section 25. Section 26B-4-214 is amended to read:
2003	26B-4-214 . Medical cannabis caregiver card Registration Renewal
2004	Revocation.
2005	(1)(a) A cardholder described in Section 26B-4-213 may designate[, through the state
2006	central patient portal,] up to two individuals, or an individual and a facility in
2007	accordance with Subsection (1)(b), to serve as a designated caregiver for the
2008	cardholder.
2009	(b)(i) A cardholder described in Section 26B-4-213 may designate one of the
2010	following types of facilities as one of the caregivers described in Subsection (1)(a):
2011	(A) for a patient or resident, an assisted living facility, as that term is defined in
2012	Section 26B-2-201;
2013	(B) for a patient or resident, a nursing care facility, as that term is defined in
2014	Section 26B-2-201; or
2015	(C) for a patient, a general acute hospital, as that term is defined in Section
2016	26B-2-201.
2017	(ii) A facility may:
2018	(A) assign one or more employees to assist patients with medical cannabis
2019	treatment under the caregiver designation described in this Subsection (1)(b);
2020	and
2021	(B) receive a medical cannabis shipment from a medical cannabis pharmacy or a
2022	medical cannabis courier on behalf of the medical cannabis cardholder within
2023	the facility who designated the facility as a caregiver.
2024	(iii) The department shall make rules to regulate the practice of facilities and facility
2025	employees serving as designated caregivers under this Subsection (1)(b).
2026	(c) A parent or legal guardian described in Subsection 26B-4-213(2)(d), in consultation
2027	with the minor and the minor's qualified medical provider, may designate[, through
2028	the state central patient portal,] up to two individuals to serve as a designated
2029	caregiver for the minor, if the department determines that the parent or legal guardian
2030	is not eligible for a medical cannabis guardian card under Section 26B-4-213.
2031	(d)(i) Upon the entry of a caregiver designation under Subsection (1) by a patient
2032	with a terminal illness described in Section 26B-4-203, the department shall issue
2033	to the designated caregiver an electronic conditional medical cannabis caregiver
2034	card, in accordance with this Subsection (1)(d).

2035	(ii) A conditional medical cannabis caregiver card is valid for the lesser of:
2036	(A) 60 days; or
2037	(B) the day on which the department completes the department's review and issues
2038	a medical cannabis caregiver card under Subsection (1)(a), denies the patient's
2039	medical cannabis caregiver card application, or revokes the conditional
2040	medical cannabis caregiver card under <u>Section</u> 26B-4-246.
2041	(iii) The department may issue a conditional medical cannabis card to an individual
2042	applying for a medical cannabis patient card for which approval of the
2043	Compassionate Use Board is not required.
2044	(iv) An individual described in Subsection (1)(b)(iii) has the rights, restrictions, and
2045	obligations under law applicable to a holder of the medical cannabis card for
2046	which the individual applies and for which the department issues the conditional
2047	medical cannabis card.
2048	(2) An individual that the department registers as a designated caregiver under this section
2049	and a facility described in Subsection (1)(b):
2050	(a) for an individual designated caregiver, may carry a valid medical cannabis caregiver
2051	card;
2052	(b) in accordance with this part, may purchase, possess, transport, or assist the patient in
2053	the use of [cannabis in a medicinal dosage form, a cannabis product in a medicinal
2054	dosage form,] medical cannabis or a medical cannabis device on behalf of the
2055	designating medical cannabis cardholder;
2056	(c) may not charge a fee to an individual to act as the individual's designated caregiver
2057	or for a service that the designated caregiver provides in relation to the role as a
2058	designated caregiver; and
2059	(d) may accept reimbursement from the designating medical cannabis cardholder for
2060	direct costs the designated caregiver incurs for assisting with the designating
2061	cardholder's medicinal use of cannabis.
2062	(3)(a) The department shall:
2063	(i) within 15 days after the day on which an individual submits an application in
2064	compliance with this section, issue a medical cannabis card to the applicant if the
2065	applicant:
2066	(A) is designated as a caregiver under Subsection (1);
2067	(B) is eligible for a medical cannabis caregiver card under Subsection (4); and
2068	(C) complies with this section; and

2069			(ii) notify the Department of Public Safety of each individual that the department
2070			registers as a designated caregiver.
2071		(b)	The department shall ensure that a medical cannabis caregiver card contains the
2072			information described in Subsections (5)(b) and (3)(c)(i).
2073		(c)	If a cardholder described in Section 26B-4-213 designates an individual as a
2074			caregiver who already holds a medical cannabis caregiver card, the individual with
2075			the medical cannabis caregiver card:
2076			(i) shall report to the department the information required of applicants under
2077			Subsection (5)(b) regarding the new designation;
2078			(ii) if the individual makes the report described in Subsection (3)(c)(i), is not required
2079			to file an application for another medical cannabis caregiver card;
2080			(iii) may receive an additional medical cannabis caregiver card in relation to each
2081			additional medical cannabis patient who designates the caregiver; and
2082			(iv) is not subject to an additional background check.
2083	(4)	An	individual is eligible for a medical cannabis caregiver card if the individual:
2084		(a)	is at least 21 years old;
2085		(b)	is a Utah resident;
2086		(c)	pays to the department a fee in an amount that, subject to Subsection 26B-1-310(5),
2087			the department sets in accordance with Section 63J-1-504, plus the cost of the
2088			criminal background check described in Section 26B-4-215; and
2089		(d)	signs an acknowledgment stating that the applicant received the information
2090			described in Subsection 26B-4-213(9)[-].
2091	(5)	An	eligible applicant for a medical cannabis caregiver card shall:
2092		(a)	submit an application for a medical cannabis caregiver card to the department
2093			through an electronic application connected to the state electronic verification
2094			system; and
2095		(b)	submit the following information in the application described in Subsection (5)(a):
2096			(i) the applicant's name, gender, age, and address;
2097			(ii) the name, gender, age, and address of the cardholder described in Section
2098			26B-4-213 who designated the applicant;
2099			(iii) if a medical cannabis guardian cardholder designated the caregiver, the name,
2100			gender, and age of the minor receiving a medical cannabis treatment in relation to
2101			the medical cannabis guardian cardholder; and
2102			(iv) any additional information that the department requests to assist in matching the

2103	application with the designating medical cannabis patient.
2104	(6) Except as provided in Subsection (6)(b), a medical cannabis caregiver card that the
2105	department issues under this section is valid for the lesser of:
2106	(a) an amount of time that the cardholder described in Section 26B-4-213 who
2107	designated the caregiver determines; or
2108	(b) the amount of time remaining before the card of the cardholder described in Section
2109	26B-4-213 expires.
2110	(7)(a) If a designated caregiver meets the requirements of Subsection (4), the designated
2111	caregiver's medical cannabis caregiver card renews automatically at the time the
2112	cardholder described in Section 26B-4-213 who designated the caregiver:
2113	(i) renews the cardholder's card; and
2114	(ii) renews the caregiver's designation, in accordance with Subsection (7)(b).
2115	(b) The department shall provide a method in the card renewal process to allow a
2116	cardholder described in Section 26B-4-213 who has designated a caregiver to:
2117	(i) signify that the cardholder renews the caregiver's designation;
2118	(ii) remove a caregiver's designation; or
2119	(iii) designate a new caregiver.
2120	(8) The department shall record the issuance or revocation of a medical cannabis card under
2121	this section in the controlled substance database.
2122	Section 26. Section 26B-4-222 is amended to read:
2123	26B-4-222 . Report.
2124	(1) By the November interim meeting each year, the department shall report to the Health
2125	and Human Services Interim Committee on:
2126	(a) the number of applications and renewal applications filed for medical cannabis cards;
2127	(b) the number of qualifying patients and designated caregivers;
2128	(c) the nature of the debilitating medical conditions of the qualifying patients;
2129	(d) the age and county of residence of cardholders;
2130	(e) the number of medical cannabis cards revoked;
2131	(f) the number of practitioners providing recommendations for qualifying patients;
2132	(g) the number of license applications and renewal license applications received;
2133	(h) the number of licenses the department has issued in each county;
2134	(i) the number of licenses the department has revoked;
2135	(j) the quantity of medical cannabis shipments[that the state central patient portal
2136	facilitates]:

2137	(k) the number of overall purchases of medical cannabis [and medical cannabis products]
2138	from each medical cannabis pharmacy;
2139	(1) the expenses incurred and revenues generated from the medical cannabis program;
2140	and
2141	(m) an analysis of product availability in medical cannabis pharmacies in consultation
2142	with the Department of Agriculture and Food.
2143	(2) The report shall include information provided by the Center for Medical Cannabis
2144	Research described in Section 53B-17-1402.
2145	(3) The department may not include personally identifying information in the report
2146	described in this section.
2147	(4) The department shall report to the working group described in Section 36-12-8.2 as
2148	requested by the working group.
2149	Section 27. Section 58-37-3.6 is amended to read:
2150	58-37-3.6. Exemption for possession or distribution of a cannabinoid product,
2151	expanded cannabinoid product, or transportable industrial hemp concentrate.
2152	(1) As used in this section:
2153	(a) "Cannabinoid product" means a product intended for human ingestion that:
2154	(i) contains an extract or concentrate that is obtained from cannabis; and
2155	[(ii) is prepared in a medicinal dosage form; and]
2156	[(iii)] (iii) contains at least 10 units of cannabidiol for every one unit of
2157	tetrahydrocannabinol.
2158	(b) "Cannabis" means any part of the plant cannabis sativa, whether growing or not.
2159	[(e) "Drug paraphernalia" means the same as that term is defined in Section 58-37a-3.]
2160	[(d)] (c) "Expanded cannabinoid product" means a product intended for human ingestion
2161	that:
2162	(i) contains an extract or concentrate that is obtained from cannabis; and
2163	[(ii) is prepared in a medicinal dosage form; and]
2164	[(iii)] (ii) contains less than 10 units of cannabidiol for every one unit of
2165	tetrahydrocannabinol.
2166	[(e) "Hemp cannabinoid product" means a product that:]
2167	[(i) contains or is represented to contain one or more naturally occurring
2168	cannabinoids;]
2169	[(ii) contains less than the cannabinoid product THC level, by dry weight;]
2170	[(iii) contains a combined amount of total THC and any THC analog that does not

2171	exceed 10% of the total cannabinoid content;]
2172	[(iv) does not exceed a total of THC and any THC analog that is greater than five
2173	milligrams per serving and 150 milligrams per package; and]
2174	[(v) unless the product is in an oil based suspension, has a serving size that is an
2175	integer.]
2176	[(f)] (d) "Transportable industrial hemp concentrate" means any amount of a natural
2177	cannabinoid in a purified state that:
2178	(i) is the product of any chemical or physical process applied to naturally occurring
2179	biomass that concentrates or isolates the cannabinoids contained in the biomass;
2180	(ii) is derived from a cannabis plant that, based on sampling that was collected no
2181	more than 30 days before the day on which the cannabis plant was harvested,
2182	contains a combined concentration of total THC and any THC analog of less than
2183	0.3% on a dry weight basis; and
2184	(iii) has a THC and THC analog concentration total less than 20% when concentrated
2185	from the cannabis plant to the purified state.
2186	[(g) "Medicinal dosage form" means:]
2187	[(i) a tablet;]
2188	[(ii) a capsule;]
2189	[(iii) a concentrated oil;]
2190	[(iv) a liquid suspension;]
2191	[(v) a transdermal preparation; or]
2192	[(vi) a sublingual preparation.]
2193	[(h)] (e) "Tetrahydrocannabinol" means a substance derived from cannabis that meets the
2194	description in Subsection 58-37-4(2)(a)(iii)(AA).
2195	(2) Notwithstanding any other provision of this chapter an individual who possesses or
2196	distributes a cannabinoid product or an expanded cannabinoid product is not subject to
2197	the penalties described in this title for the possession or distribution of marijuana or
2198	tetrahydrocannabinol to the extent that the individual's possession or distribution of the
2199	cannabinoid product or expanded cannabinoid product complies with [Title 26B,
2200	Chapter 4, Part 2, Cannabinoid Research and Medical Cannabis] Section 26B-4-212.
2201	(3) Notwithstanding any other provision of this chapter, a person who possesses and
2202	distributes transportable industrial hemp concentrate is not subject to the penalties
2203	described in this chapter for the possession or distribution of transportable industrial
2204	hemp concentrate if the transportable industrial hemp concentrate is handled in

2205		accordance with the rules established under Subsection 4-41-103.1(1)(e) or is destroyed.
2206		Section 28. Section 58-85-102 is amended to read:
2207		58-85-102 . Definitions.
2208		As used in this chapter:
2209	(1)	"Eligible patient" means an individual who has been diagnosed with a terminal illness
2210		by a physician.
2211	(2)	"Insurer" means the same as that term is defined in Section 31A-1-301.
2212	(3)	"Investigational device" means a device that:
2213		(a) meets the definition of "investigational device" in 21 C.F.R. Sec. 812.3; and
2214		(b) has successfully completed the United States Food and Drug Administration Phase $\boldsymbol{1}$
2215		testing for an investigational device described in 21 C.F.R. Part 812.
2216	(4)	"Investigational drug" means a drug that:
2217		(a) meets the definition of "investigational new drug" in 21 C.F.R. Sec. 312.3; and
2218		(b) has successfully completed the United States Food and Drug Administration Phase $\boldsymbol{1}$
2219		testing for an investigational new drug described in 21 C.F.R. Part 312.
2220	(5)	"Medicinal dosage form" [means the same as that term is defined in Section 58-37-3.6.]
2221		means:
2222		(a) a tablet;
2223		(b) a capsule;
2224		(c) a concentrated oil;
2225		(d) a liquid suspension;
2226		(e) a transdermal preparation; or
2227		(f) a sublingual preparation.
2228	(6)	"Physician" means an individual who is licensed under:
2229		(a) Title 58, Chapter 67, Utah Medical Practice Act; or
2230		(b) Title 58, Chapter 68, Utah Osteopathic Medical Practice Act.
2231	(7)	"Terminal illness" means a condition of a patient that:
2232		(a) as determined by a physician:
2233		(i) is likely to pose a greater risk to the patient than the risk posed to the patient by
2234		treatment with an investigational drug or investigational device; and
2235		(ii) will inevitably lead to the patient's death; and
2236		(b) presents the patient, after the patient has explored conventional therapy options, with
2237		no treatment option that is satisfactory or comparable to treatment with an
2238		investigational drug or device.

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2239	Section 29. Section 63N-3-1301 is amended to read:
2240	63N-3-1301 . Definitions.
2241	As used in this part:
2242	(1) "Cannabinoid processor license" means the same as that term is defined in Section
2243	4-41-102.
2244	(2) "Cannabinoid product" means the same as that term is defined in Section 4-41-102.
2245	(3) "Industrial hemp_product" means the same as that term is defined in Section 4-41-102.
2246	(4) "Industrial hemp producer registration" means the same as that term is defined in
2247	Section 4-41-102.
2248	Section 30. Section 77-39-101 is amended to read:
2249	77-39-101. Investigation of sales of alcohol, tobacco products, electronic
2250	cigarette products, nicotine products, and cannabinoid products to underage individuals.
2251	(1) As used in this section:
2252	(a) "Cannabinoid product" means the same as that term is defined in Section 4-41-102.
2253	(b) "Electronic cigarette product" means the same as that term is defined in Section
2254	76-10-101.
2255	(c) "Nicotine product" means the same as that term is defined in Section 76-10-101.
2256	(d) "Peace officer" means the same as the term is described in Section 53-13-109.
2257	(e) "Tobacco product" means the same as that term is defined in Section 76-10-101.
2258	(2)(a) A peace officer may investigate the possible violation of:
2259	(i) Section 32B-4-403 by requesting an individual under 21 years old to enter into
2260	and attempt to purchase or make a purchase of alcohol from a retail establishment;
2261	(ii) Section 76-10-114 by requesting an individual under 21 years old to enter into
2262	and attempt to purchase or make a purchase from a retail establishment of:
2263	(A) a tobacco product;
2264	(B) an electronic cigarette product; or
2265	(C) a nicotine product; or
2266	(iii) Subsection $[4-41-105(2)(d)]$ $4-41-105(2)(a)(iv)$ by requesting an individual under
2267	21 years old to enter into and attempt to purchase or make a purchase of a
2268	cannabinoid product that contains THC or a THC analog from a retail
2269	establishment.
2270	(b) A peace officer who is present at the site of a proposed purchase shall direct,
2271	supervise, and monitor the individual requested to make the purchase.

(c) Immediately following a purchase or attempted purchase or as soon as practical the

2273	supervising peace officer shall inform the cashier and the proprietor or manager of
2274	the retail establishment that the attempted purchaser was under the legal age to
2275	purchase:
2276	(i) alcohol;
2277	(ii)(A) a tobacco product;
2278	(B) an electronic cigarette product; or
2279	(C) a nicotine product; or
2280	(iii) a cannabinoid product that contains THC or a THC analog.
2281	(d) If a citation or information is issued, the citation or information shall be issued
2282	within seven days after the day on which the purchase occurs.
2283	(3)(a) If an individual under 18 years old is requested to attempt a purchase, a written
2284	consent of that individual's parent or guardian shall be obtained before the individual
2285	participates in any attempted purchase.
2286	(b) An individual requested by the peace officer to attempt a purchase may:
2287	(i) be a trained volunteer; or
2288	(ii) receive payment, but may not be paid based on the number of successful
2289	purchases of alcohol, tobacco products, electronic cigarette products, nicotine
2290	products, or cannabinoid products that contain THC or a THC analog.
2291	(4) The individual requested by the peace officer to attempt a purchase and anyone
2292	accompanying the individual attempting a purchase may use false identification in
2293	attempting the purchase if:
2294	(a) the Department of Public Safety created in Section 53-1-103 provides the false
2295	identification;
2296	(b) the false identification:
2297	(i) accurately represents the individual's age; and
2298	(ii) displays a current photo of the individual; and
2299	(c) the peace officer maintains possession of the false identification at all times outside
2300	the attempt to purchase.
2301	(5) An individual requested to attempt to purchase or make a purchase pursuant to this
2302	section is immune from prosecution, suit, or civil liability for the purchase of, attempted
2303	purchase of, or possession of alcohol, a tobacco product, an electronic cigarette product,
2304	a nicotine product, or a cannabinoid product that contains THC or a THC analog if a
2305	peace officer directs, supervises, and monitors the individual.
2306	(6)(a) Except as provided in Subsection (6)(b), a purchase attempted under this section

2307	shall be conducted within a 12-month period:
2308	(i) on a random basis at any one retail establishment location, not more often than
2309	four times for the attempted purchase of alcohol;
2310	(ii) a minimum of two times at a retail establishment that sells tobacco products,
2311	electronic cigarette products, or nicotine products for the attempted purchase of a
2312	tobacco product, an electronic cigarette product, or a nicotine product; and
2313	(iii) a minimum of one time at a retail establishment that sells a cannabinoid product
2314	that contains THC or a THC analog.
2315	(b) This section does not prohibit an investigation or an attempt to purchase alcohol, a
2316	tobacco product, an electronic cigarette product, or a nicotine product under this
2317	section if:
2318	(i) there is reasonable suspicion to believe the retail establishment has sold alcohol, a
2319	tobacco product, an electronic cigarette product, a nicotine product, or a
2320	cannabinoid product that contains THC or a THC analog to an individual under
2321	the age established by Section 32B-4-403, Section 76-10-114, or Subsection
2322	4-41-105(2)(d); and
2323	(ii) the supervising peace officer makes a written record of the grounds for the
2324	reasonable suspicion.
2325	(7)(a) The peace officer exercising direction, supervision, and monitoring of the
2326	attempted purchase shall make a report of the attempted purchase, whether or not a
2327	purchase was made.
2328	(b) The report required by this Subsection (7) shall include:
2329	(i) the name of the supervising peace officer;
2330	(ii) the name of the individual attempting the purchase;
2331	(iii) a photograph of the individual attempting the purchase showing how that
2332	individual appeared at the time of the attempted purchase;
2333	(iv) the name and description of the cashier or proprietor from whom the individual
2334	attempted the purchase;
2335	(v) the name and address of the retail establishment; and
2336	(vi) the date and time of the attempted purchase.
2337	Section 31. Repealer.
2338	This bill repeals:
2339	Section 26B-4-236, State central patient portal Department duties.
2340	Section 32. Effective Date.

2341 This bill takes effect on May 7, 2025.