26	4-41a-406, as last amended by Laws of Utah 2023, Chapter 327
27	4-41a-1001, as last amended by Laws of Utah 2023, Chapter 317 and renumbered and
28	amended by Laws of Utah 2023, Chapters 273, 307 and last amended by
29	Coordination Clause, Laws of Utah 2023, Chapter 307
30	10-9a-528, as last amended by Laws of Utah 2023, Chapters 273, 327 and last amended
31	by Coordination Clause, Laws of Utah 2023, Chapter 327
32	17-27a-525, as last amended by Laws of Utah 2023, Chapters 273, 327 and last
33	amended by Coordination Clause, Laws of Utah 2023, Chapter 327
34	26B-1-435, as enacted by Laws of Utah 2023, Chapter 273
35	26B-4-219, as last amended by Laws of Utah 2023, Chapters 273, 317 and renumbered
36	and amended by Laws of Utah 2023, Chapter 307 and last amended by
37	Coordination Clause, Laws of Utah 2023, Chapter 307
38	26B-4-231, as last amended by Laws of Utah 2023, Chapter 317 and renumbered and
39	amended by Laws of Utah 2023, Chapters 273, 307 and last amended by
40	Coordination Clause, Laws of Utah 2023, Chapter 307
41	ENACTS:
<b>4</b> 2	4-41a-1206, Utah Code Annotated 1953
13	REPEALS:
14 15	26B-1-435.1, as enacted by Laws of Utah 2023, Chapter 273
+3 46	Be it enacted by the Legislature of the state of Utah:
<b>1</b> 7	Section 1. Section 4-41a-102 is amended to read:
18	4-41a-102. Definitions.
19	As used in this chapter:
50	(1) "Adulterant" means any poisonous or deleterious substance in a quantity that may
51	be injurious to health, including:
52	(a) pesticides;
53	(b) heavy metals;
54	(c) solvents;
55	
	(d) microbial life;
56	<ul><li>(d) microbial life;</li><li>(e) artificially derived cannabinoid;</li></ul>

57	(f) toxins; or
58	(g) foreign matter.
59	(2) "Advisory board" means the Medical Cannabis Policy Advisory Board created in
60	Section 26B-1-435.
61	(3) (a) "Artificially derived cannabinoid" means a chemical substance that is created by
62	a chemical reaction that changes the molecular structure of any chemical substance derived
63	from the cannabis plant.
64	(b) "Artificially derived cannabinoid" does not include:
65	(i) a naturally occurring chemical substance that is separated from the cannabis plant
66	by a chemical or mechanical extraction process; or
67	(ii) a cannabinoid that is produced by decarboxylation from a naturally occurring
68	cannabinoid acid without the use of a chemical catalyst.
69	(4) "Cannabis Research Review Board" means the Cannabis Research Review Board
70	created in Section 26B-1-420.
71	(5) "Cannabis" means the same as that term is defined in Section 26B-4-201.
72	(6) "Cannabis concentrate" means:
73	(a) the product of any chemical or physical process applied to naturally occurring
74	biomass that concentrates or isolates the cannabinoids contained in the biomass; and
75	(b) any amount of a natural cannabinoid or artificially derived cannabinoid in an
76	artificially derived cannabinoid's purified state.
77	(7) "Cannabis cultivation byproduct" means any portion of a cannabis plant that is not
78	intended to be sold as a cannabis plant product.
79	(8) "Cannabis cultivation facility" means a person that:
80	(a) possesses cannabis;
81	(b) grows or intends to grow cannabis; and
82	(c) sells or intends to sell cannabis to a cannabis cultivation facility, a cannabis
83	processing facility, or a medical cannabis research licensee.
84	(9) "Cannabis cultivation facility agent" means an individual who[:]
85	holds a valid cannabis production establishment agent registration card with a cannabis
86	cultivation facility designation.
87	(10) "Cannabis derivative product" means a product made using cannabis concentrate.

88	(11) "Cannabis plant product" means any portion of a cannabis plant intended to be
89	sold in a form that is recognizable as a portion of a cannabis plant.
90	(12) "Cannabis processing facility" means a person that:
91	(a) acquires or intends to acquire cannabis from a cannabis production establishment;
92	(b) possesses cannabis with the intent to manufacture a cannabis product;
93	(c) manufactures or intends to manufacture a cannabis product from unprocessed
94	cannabis or a cannabis extract; and
95	(d) sells or intends to sell a cannabis product to a medical cannabis pharmacy or a
96	medical cannabis research licensee.
97	(13) "Cannabis processing facility agent" means an individual who[:]
98	holds a valid cannabis production establishment agent registration card with a cannabis
99	processing facility designation.
100	(14) "Cannabis product" means the same as that term is defined in Section 26B-4-201.
101	(15) "Cannabis production establishment" means a cannabis cultivation facility, a
102	cannabis processing facility, or an independent cannabis testing laboratory.
103	(16) "Cannabis production establishment agent" means a cannabis cultivation facility
104	agent, a cannabis processing facility agent, or an independent cannabis testing laboratory agent.
105	(17) "Cannabis production establishment agent registration card" means a registration
106	card that the department issues that:
107	(a) authorizes an individual to act as a cannabis production establishment agent; and
108	(b) designates the type of cannabis production establishment for which an individual is
109	authorized to act as an agent.
110	(18) "Closed-door medical cannabis pharmacy" means a facility operated by a home
111	delivery medical cannabis pharmacy for delivering cannabis or a medical cannabis product.
112	[(18)] (19) "Community location" means a public or private elementary or secondary
113	school, a church, a public library, a public playground, or a public park.
114	[(19)] (20) "Cultivation space" means, quantified in square feet, the horizontal area in
115	which a cannabis cultivation facility cultivates cannabis, including each level of horizontal area
116	if the cannabis cultivation facility hangs, suspends, stacks, or otherwise positions plants above
117	other plants in multiple levels.
118	[ <del>(20)</del> ] <u>(21)</u> "Delivery address" means:

119	(a) for a medical cannabis cardholder who is not a facility, the medical cannabis
120	cardholder's home address; or
121	(b) for a medical cannabis cardholder that is a facility, the facility's address.
122	[(21)] (22) "Department" means the Department of Agriculture and Food.
123	[(22)] (23) "Family member" means a parent, step-parent, spouse, child, sibling,
124	step-sibling, uncle, aunt, nephew, niece, first cousin, mother-in-law, father-in-law,
125	brother-in-law, sister-in-law, son-in-law, daughter-in-law, grandparent, or grandchild.
126	[(23)] (24) "Home delivery medical cannabis pharmacy" means a medical cannabis
127	pharmacy that the department authorizes, as part of the pharmacy's license, to deliver medical
128	cannabis shipments to a delivery address to fulfill electronic orders that the state central patient
129	portal facilitates.
130	$\left[\frac{(24)}{(25)}\right]$ (a) "Independent cannabis testing laboratory" means a person that:
131	(i) conducts a chemical or other analysis of cannabis or a cannabis product; or
132	(ii) acquires, possesses, and transports cannabis or a cannabis product with the intent to
133	conduct a chemical or other analysis of the cannabis or cannabis product.
134	(b) "Independent cannabis testing laboratory" includes a laboratory that the department
135	or a research university operates in accordance with Subsection 4-41a-201(14).
136	[(25)] (26) "Independent cannabis testing laboratory agent" means an individual who[:]
137	holds a valid cannabis production establishment agent registration card with an
138	independent cannabis testing laboratory designation.
139	[(26)] (27) "Inventory control system" means a system described in Section 4-41a-103.
140	[(27)] (28) "Licensing board" or "board" means the Cannabis Production Establishment
141	Licensing Advisory Board created in Section 4-41a-201.1.
142	[(28)] (29) "Medical cannabis" means the same as that term is defined in Section
143	26B-4-201.
144	[(29)] (30) "Medical cannabis card" means the same as that term is defined in Section
145	26B-4-201.
146	[(30)] (31) "Medical cannabis courier" means a courier that:
147	(a) the department licenses in accordance with Section 4-41a-1201; and
148	(b) contracts with a home delivery medical cannabis pharmacy to deliver medical
149	cannabis shipments to fulfill electronic orders that the state central patient portal facilitates.

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150	[(31)] (32) "Medical cannabis courier agent" means an individual who:
151	(a) is an employee of a medical cannabis courier; and
152	(b) who holds a valid medical cannabis courier agent registration card.
153	[(32)] (33) "Medical cannabis pharmacy" means the same as that term is defined in
154	Section 26B-4-201.
155	[(33)] (34) "Medical cannabis pharmacy agent" means the same as that term is defined
156	in Section 26B-4-201.
157	[(34)] (35) "Medical cannabis research license" means a license that the department
158	issues to a research university for the purpose of obtaining and possessing medical cannabis for
159	academic research.
160	[(35)] (36) "Medical cannabis research licensee" means a research university that the
161	department licenses to obtain and possess medical cannabis for academic research, in
162	accordance with Section 4-41a-901.
163	[(36)] (37) "Medical cannabis shipment" means a shipment of medical cannabis or a
164	medical cannabis product that a home delivery medical cannabis pharmacy or a medical
165	cannabis courier delivers to a delivery address to fulfill an electronic medical cannabis order
166	that the state central patient portal facilitates.
167	[(37)] (38) "Medical cannabis treatment" means the same as that term is defined in
168	Section 26B-4-201.
169	[(38)] (39) "Medicinal dosage form" means the same as that term is defined in Section
170	26B-4-201.
171	(40) "Pharmacy ownership limit" means an amount equal to 30% of the total number of
172	medical cannabis pharmacy licenses issued by the department rounded down to the nearest
173	whole number.
174	[(39)] (41) "Pharmacy medical provider" means the same as that term is defined in
175	Section 26B-4-201.
176	[(40)] (42) "Qualified medical provider" means the same as that term is defined in
177	Section 26B-4-201.
178	[(41)] (43) "Qualified Production Enterprise Fund" means the fund created in Section
179	4-41a-104.
180	[42] (44) "Recommending medical provider" means the same as that term is defined

181	in Section 26B-4-201.
182	[ <del>(43)</del> ] (45) "Research university" means the same as that term is defined in Section
183	53B-7-702 and a private, nonprofit college or university in the state that:
184	(a) is accredited by the Northwest Commission on Colleges and Universities;
185	(b) grants doctoral degrees; and
186	(c) has a laboratory containing or a program researching a schedule I controlled
187	substance described in Section 58-37-4.
188	[44)] (46) "State electronic verification system" means the system described in Section
189	26B-4-202.
190	[45] Tetrahydrocannabinol" or "THC" means the same as that term is defined in
191	Section 4-41-102.
192	[48] "THC analog" means the same as that term is defined in Section 4-41-102.
193	[(47)] (49) "Total composite tetrahydrocannabinol" means all detectable forms of
194	tetrahydrocannabinol.
195	[(48)] (50) "Total tetrahydrocannabinol" or "total THC" means the same as that term is
196	defined in Section 4-41-102.
197	Section 2. Section <b>4-41a-406</b> is amended to read:
198	4-41a-406. Local control.
199	(1) As used in this section:
200	(a) "Cannabis production establishment" means the same as that term is defined in
201	Section 4-41a-102 and includes a closed-door medical cannabis pharmacy.
202	(b) "Land use decision" means the same as that term is defined in Sections 10-9a-103
203	and 17-27a-103.
204	[(b)] (c) "Land use permit" means the same as that term is defined in Sections
205	10-9a-103 and 17-27a-103.
206	[(c)] (d) "Land use regulation" means the same as that term is defined in Sections
207	10-9a-103 and 17-27a-103.
208	(2) (a) If a municipality's or county's zoning ordinances provide for an industrial zone,
209	the operation of a cannabis production establishment shall be a permitted industrial use in any
210	industrial zone unless the municipality or county has designated by ordinance, before an
211	individual submits a land use permit application for a cannabis production establishment, at

212	least one industrial zone in which the operation of a cannabis production establishment is a
213	permitted use.
214	(b) If a municipality's or county's zoning ordinances provide for an agricultural zone,
215	the operation of a cannabis production establishment shall be a permitted agricultural use in
216	any agricultural zone unless the municipality or county has designated by ordinance, before an
217	individual submits a land use permit application for a cannabis production establishment, at
218	least one agricultural zone in which the operation of a cannabis production establishment is a
219	permitted use.
220	(c) The operation of a cannabis production establishment shall be a permitted use on
221	land that the municipality or county has not zoned.
222	(3) A municipality or county may not:
223	(a) on the sole basis that the applicant, or cannabis production establishment violates
224	federal law regarding the legal status of cannabis, deny or revoke:
225	(i) a land use permit to operate a cannabis production facility; or
226	(ii) a business license to operate a cannabis production facility;
227	(b) require a certain distance between a cannabis production establishment and:
228	(i) another cannabis production establishment;
229	(ii) a medical cannabis pharmacy;
230	(iii) a retail tobacco specialty business, as that term is defined in Section 26B-7-501; or
231	(iv) an outlet, as that term is defined in Section 32B-1-202; or
232	(c) in accordance with Subsections 10-9a-509(1) and 17-27a-508(1), enforce a land use
233	regulation against a cannabis production establishment that was not in effect on the day on
234	which the cannabis production establishment submitted a complete land use application.
235	(4) An applicant for a land use permit to operate a cannabis production establishment
236	shall comply with the land use requirements and application process described in:
237	(a) Title 10, Chapter 9a, Municipal Land Use, Development, and Management Act,
238	including Section 10-9a-528; and
239	(b) Title 17, Chapter 27a, County Land Use, Development, and Management Act,
240	including Section 17-27a-525.

Section 3. Section **4-41a-1001** is amended to read:

4-41a-1001. Medical cannabis pharmacy -- License -- Eligibility.

241

242

243	(1) A person may not:
244	(a) operate as a medical cannabis pharmacy without a license that the department issues
245	under this part[-];
246	(b) obtain a medical cannabis pharmacy license if obtaining the license would cause the
247	person to exceed the pharmacy ownership limit;
248	(c) obtain a partial ownership share of a medical cannabis pharmacy if obtaining the
249	partial ownership share would cause the person to exceed the pharmacy ownership limit; or
250	(d) enter into any contract or agreement that allows the person to directly or indirectly
251	control the operations of a medical cannabis pharmacy if the person's control of the medical
252	cannabis pharmacy would cause the person to effectively exceed the pharmacy ownership limit.
253	(2) (a) (i) Subject to Subsections (4) and (5) and to Section 4-41a-1005, the department
254	shall issue a license to operate a medical cannabis pharmacy in accordance with Title 63G,
255	Chapter 6a, Utah Procurement Code.
256	(ii) The department may not issue a license to operate a medical cannabis pharmacy to
257	an applicant who is not eligible for a license under this section.
258	(b) An applicant is eligible for a license under this section if the applicant submits to
259	the department:
260	(i) subject to Subsection (2)(c), a proposed name and address where the applicant will
261	operate the medical cannabis pharmacy;
262	(ii) the name and address of an individual who:
263	(A) for a publicly traded company, has a financial or voting interest of 10% or greater
264	in the proposed medical cannabis pharmacy;
265	(B) for a privately held company, a financial or voting interest in the proposed medical
266	cannabis pharmacy; or
267	(C) has the power to direct or cause the management or control of a proposed medical
268	cannabis pharmacy;
269	(iii) for each application that the applicant submits to the department, a statement from
270	the applicant that the applicant will obtain and maintain:
271	(A) a performance bond in the amount of \$100,000 issued by a surety authorized to
272	transact surety business in the state; or

(B) a liquid cash account in the amount of \$100,000 with a financial institution;

274 (iv) an operating plan that:

- 275 (A) complies with Section 4-41a-1004;
  - (B) includes operating procedures to comply with the operating requirements for a medical cannabis pharmacy described in this part and with a relevant municipal or county law that is consistent with Section 4-41a-1106; and
    - (C) the department approves;
  - (v) an application fee in an amount that, subject to Subsection 4-41a-104(5), the department sets in accordance with Section 63J-1-504; and
  - (vi) a description of any investigation or adverse action taken by any licensing jurisdiction, government agency, law enforcement agency, or court in any state for any violation or detrimental conduct in relation to any of the applicant's cannabis-related operations or businesses.
    - (c) (i) A person may not locate a medical cannabis pharmacy:
    - (A) within 200 feet of a community location; or
  - (B) in or within 600 feet of a district that the relevant municipality or county has zoned as primarily residential.
  - (ii) The proximity requirements described in Subsection (2)(c)(i) shall be measured from the nearest entrance to the medical cannabis pharmacy establishment by following the shortest route of ordinary pedestrian travel to the property boundary of the community location or residential area.
  - (iii) The department may grant a waiver to reduce the proximity requirements in Subsection (2)(c)(i) by up to 20% if the department determines that it is not reasonably feasible for the applicant to site the proposed medical cannabis pharmacy without the waiver.
  - (iv) An applicant for a license under this section shall provide evidence of compliance with the proximity requirements described in Subsection (2)(c)(i).
  - (d) The department may not issue a license to an eligible applicant that the department has selected to receive a license until the selected eligible applicant complies with the bond or liquid cash requirement described in Subsection (2)(b)(iii).
  - (e) If the department receives more than one application for a medical cannabis pharmacy within the same city or town, the department shall consult with the local land use authority before approving any of the applications pertaining to that city or town.

305 (3) If the department selects an applicant for a medical cannabis pharmacy license 306 under this section, the department shall: 307 (a) charge the applicant an initial license fee in an amount that, subject to Subsection 308 4-41a-104(5), the department sets in accordance with Section 63J-1-504; 309 (b) notify the Department of Public Safety of the license approval and the names of 310 each individual described in Subsection (2)(b)(ii); and 311 (c) charge the licensee a fee in an amount that, subject to Subsection 4-41a-104(5), the 312 department sets in accordance with Section 63J-1-504, for any change in location, ownership, 313 or company structure. 314 (4) The department may not issue a license to operate a medical cannabis pharmacy to 315 an applicant if an individual described in Subsection (2)(b)(ii): 316 (a) has been convicted under state or federal law of: 317 (i) a felony: or 318 (ii) after December 3, 2018, a misdemeanor for drug distribution; 319 (b) is younger than 21 years old; or 320 (c) after September 23, 2019, until January 1, 2023, is actively serving as a legislator. 321 (5) (a) If an applicant for a medical cannabis pharmacy license under this section holds another license under this chapter, the department may not give preference to the applicant 322 323 based on the applicant's status as a holder of the license. 324 (b) If an applicant for a medical cannabis pharmacy license under this section holds a 325 license to operate a cannabis cultivation facility under this section, the department may give 326 consideration to the applicant's status as a holder of the license if: 327 (i) the applicant demonstrates that a decrease in costs to patients is more likely to result 328 from the applicant's vertical integration than from a more competitive marketplace; and 329 (ii) the department finds multiple other factors, in addition to the existing license, that 330 support granting the new license. 331 (6) (a) The department may revoke a license under this part: 332 (i) if the medical cannabis pharmacy does not begin operations within one year after 333 the day on which the department issues an announcement of the department's intent to award a 334 license to the medical cannabis pharmacy;

(ii) after the third the same violation of this chapter in any of the licensee's licensed

cannabis production establishments or medical cannabis pharmacies;

- (iii) if an individual described in Subsection (2)(b)(ii) is convicted, while the license is active, under state or federal law of:
  - (A) a felony; or

- (B) after December 3, 2018, a misdemeanor for drug distribution;
- (iv) if the licensee fails to provide the information described in Subsection (2)(b)(vi) at the time of application, or fails to supplement the information described in Subsection (2)(b)(vi) with any investigation or adverse action that occurs after the submission of the application within 14 calendar days after the licensee receives notice of the investigation or adverse action;
- (v) if the medical cannabis pharmacy demonstrates a willful or reckless disregard for the requirements of this chapter or the rules the department makes in accordance with this chapter; or
- (vi) if, after a change of ownership described in Subsection (11)(c), the department determines that the medical cannabis pharmacy no longer meets the minimum standards for licensure and operation of the medical cannabis pharmacy described in this chapter.
- (b) The department shall rescind a notice of an intent to issue a license under this part to an applicant or revoke a license issued under this part if the associated medical cannabis pharmacy does not begin operation on or before June 1, 2021.
- (7) (a) A person who receives a medical cannabis pharmacy license under this chapter, if the municipality or county where the licensed medical cannabis pharmacy will be located requires a local land use permit, shall submit to the department a copy of the licensee's approved application for the land use permit within 120 days after the day on which the department issues the license.
- (b) If a licensee fails to submit to the department a copy the licensee's approved land use permit application in accordance with Subsection (7)(a), the department may revoke the licensee's license.
- (8) The department shall deposit the proceeds of a fee imposed by this section into the Qualified Production Enterprise Fund.
- 365 (9) The department shall begin accepting applications under this part on or before 366 March 1, 2020.

367	(10) (a) The department's authority to issue a license under this section is plenary and is
368	not subject to review.
369	(b) Notwithstanding Subsection (2), the decision of the department to award a license
370	to an applicant is not subject to:
371	(i) Title 63G, Chapter 6a, Part 16, Protests; or
372	(ii) Title 63G, Chapter 6a, Part 17, Procurement Appeals Board.
373	(11) (a) A medical cannabis pharmacy license is not transferrable or assignable.
374	(b) A medical cannabis pharmacy shall report in writing to the department no later than
375	10 business days before the date of any change of ownership of the medical cannabis
376	pharmacy.
377	(c) If the ownership of a medical cannabis pharmacy changes by 50% or more:
378	(i) concurrent with the report described in Subsection (11)(b), the medical cannabis
379	pharmacy shall submit a new application described in Subsection (2)(b), subject to Subsection
380	(2)(c);
381	(ii) within 30 days of the submission of the application, the department shall:
382	(A) conduct an application review; and
383	(B) award a license to the medical cannabis pharmacy for the remainder of the term of
384	the medical cannabis pharmacy's license before the ownership change if the medical cannabis
385	pharmacy meets the minimum standards for licensure and operation of the medical cannabis
386	pharmacy described in this chapter; and
387	(iii) if the department approves the license application, notwithstanding Subsection (3),
388	the medical cannabis pharmacy shall pay a license fee that the department sets in accordance
389	with Section 63J-1-504 in an amount that covers the board's cost of conducting the application
390	review.
391	Section 4. Section 4-41a-1206 is enacted to read:
392	4-41a-1206. Closed-door medical cannabis pharmacy.
393	(1) (a) Subject to Subsections (1)(b) and (c), a home delivery medical cannabis
394	pharmacy may open a single closed-door medical cannabis pharmacy.
395	(b) A home delivery medical cannabis pharmacy may not open a closed-door medical
396	cannabis pharmacy unless the home delivery medical cannabis pharmacy:
397	(i) has an operating plan that includes a closed-door medical cannabis pharmacy; and

398	(ii) obtains a license issued by the department for a closed-door medical cannabis
399	pharmacy.
400	(c) An entity that owns multiple home delivery medical cannabis pharmacies may open
401	only one closed-door medical cannabis pharmacy.
402	(d) The department may institute a fee in accordance with Section 63J-1-504 to
403	administer this section.
404	(2) A home delivery medical cannabis pharmacy that opens a closed-door medical
405	cannabis pharmacy under Subsection (1) shall ensure:
406	(a) that a pharmacy medical provider who is a licensed pharmacist:
407	(i) is directly supervising the packaging of an order; and
408	(ii) is present in the closed-door medical cannabis pharmacy when an order is packaged
409	for delivery; and
410	(b) all record keeping requirements, labeling requirements, and patient counseling
411	requirements described in this chapter and Title 26B, Chapter 4, Part 2, Cannabinoid Research
412	and Medical Cannabis, are satisfied before sending out an order.
413	(3) An individual who prepares an order at a closed-door medical cannabis pharmacy
414	under this section shall be registered as:
415	(a) a pharmacy medical provider; or
416	(b) a medical cannabis pharmacy agent.
417	(4) (a) A closed-door medical cannabis pharmacy shall operate:
418	(i) except as provided in Subsection (4)(b), in a facility that is accessible only by an
419	individual who is a pharmacy medical provider or a medical cannabis pharmacy agent; and
420	(ii) at a physical address in accordance with Subsection (6).
421	(b) A closed-door medical cannabis pharmacy may authorize an individual who is at
422	least 18 years old and is not a pharmacy medical provider or a cannabis pharmacy agent to
423	access the closed-door medical cannabis pharmacy if the closed-door medical cannabis
424	pharmacy:
425	(i) tracks and monitors the individual at all times while the individual is at the
426	closed-door medical cannabis pharmacy; and
427	(ii) maintains a record of the individual's access, including arrival and departure.
428	(c) A closed-door medical cannabis pharmacy shall operate in a facility that has:

429	(i) a single, secure public entrance; and
430	(ii) a security system with a backup power source that:
431	(A) detects and records entry into the closed-door medical cannabis pharmacy;
432	(B) provides notice of an unauthorized entry to law enforcement when the closed-door
433	medical cannabis pharmacy is closed; and
434	(C) a lock or equivalent restrictive security feature on any area where the closed-door
435	medical cannabis pharmacy stores a cannabis product.
436	(d) A closed-door medical cannabis pharmacy shall ensure that any cannabis or
437	cannabis products in the closed-door medical cannabis pharmacy that are intended for home
438	delivery are separated in a manner that is readily distinguishable from any other cannabis or
439	cannabis product in the facility.
440	(5) A closed-door medical cannabis pharmacy may only provide cannabis or a cannabis
441	product to an individual through a delivery that complies with this part.
442	(6) (a) A person may not locate a closed-door medical cannabis pharmacy:
443	(i) within 1,000 feet of a community location; or
444	(ii) in or within 600 feet of a district that the relevant municipality or county has zoned
445	as primarily residential.
446	(b) The proximity requirements described in Subsection (6)(a) shall be measured from
447	the nearest entrance to the closed-door medical cannabis pharmacy by following the shortest
448	route of ordinary pedestrian travel to the property boundary of the community location or
449	residential area.
450	(c) The licensing board may grant a waiver to reduce the proximity requirements in
451	Subsection (6)(a) by up to 20% if the licensing board determines that it is not reasonably
452	feasible for the applicant to site the proposed closed-door medical cannabis pharmacy without
453	the waiver.
454	(d) An applicant for a license under this section shall provide evidence of compliance
455	with the proximity requirements described in Subsection (6)(a).
456	(7) When determining where a closed-door medical cannabis pharmacy may open, the
457	licensing board:
458	(a) shall utilize geographic regions created by the department through rule;
459	(b) shall prioritize allowing entities that do not have a medical cannabis pharmacy in a

160	region to open a closed-door medical cannabis pharmacy in the region;
461	(c) if the total amount of closed-door medical cannabis pharmacies, may allow only
462	three closed-door medical cannabis pharmacies to operate in counties of the first and second
463	class as described in Section 17-50-501; and
464	(d) for determining the three closed-door medical cannabis pharmacies described in
465	Subsection (7)(c), consider the following:
466	(i) the history of compliance with state law and rules for all licenses issued under this
467	chapter;
468	(ii) the medical cannabis pharmacy's willingness to offer a variety of brands and
169	products;
470	(iii) the ability of the operating plan to ensure the safety and security of the community;
471	(iv) the suitability of the proposed location and the location's ability to serve the local
472	community; and
473	(v) any other relevant information determined through rule.
174	(8) A closed-door medical cannabis pharmacy may not account for more than:
175	(a) for an entity that holds a single medical cannabis pharmacy license, the greater of:
476	(i) 35% of the medical cannabis pharmacy's total revenue; or
177	(ii) \$2,000,000 in total revenue; or
478	(b) for an entity that holds more than one medical cannabis pharmacy license, the
179	greater of:
480	(i) 35% of the total revenue of the entity's medical cannabis pharmacy that generates
481	the most revenue; or
482	(ii) \$2,000,000 in total revenue.
483	(9) In accordance with Title 63G, Chapter 3, Utah Administrative Rulemaking Act, the
184	department shall make rules to implement this section.
485	Section 5. Section 10-9a-528 is amended to read:
486	10-9a-528. Cannabis production establishments, medical cannabis pharmacies,
187	and industrial hemp producer licensee.
488	(1) As used in this section:
189	(a) "Cannabis production establishment" means the same as that term is defined in
190	Section 4-41a-102 and includes a closed-door medical cannabis pharmacy.

491	(b) "Closed-door medical cannabis pharmacy" means the same as that term is defined
492	in Section 4-41a-102.
493	[(b)] (c) "Industrial hemp producer licensee" means the same as the term "licensee" is
494	defined in Section 4-41-102.
495	[(c)] (d) "Medical cannabis pharmacy" means the same as that term is defined in
496	Section 26B-4-201.
497	(2) (a) (i) A municipality may not regulate a cannabis production establishment or a
498	medical cannabis pharmacy in conflict with:
499	(A) Title 4, Chapter 41a, Cannabis Production Establishments and Pharmacies, and
500	applicable jurisprudence; and
501	(B) this chapter.
502	(ii) A municipality may not regulate an industrial hemp producer licensee in conflict
503	with:
504	(A) Title 4, Chapter 41, Hemp and Cannabinoid Act, and applicable jurisprudence; and
505	(B) this chapter.
506	(b) The Department of Agriculture and Food has plenary authority to license programs
507	or entities that operate a cannabis production establishment or a medical cannabis pharmacy.
508	(3) (a) Within the time period described in Subsection (3)(b), a municipality shall
509	prepare and adopt a land use regulation, development agreement, or land use decision in
510	accordance with this title and:
511	(i) regarding a cannabis production establishment, Section 4-41a-406; or
512	(ii) regarding a medical cannabis pharmacy, Section [4-41a-110] 4-41a-1105.
513	(b) A municipality shall take the action described in Subsection (3)(a):
514	(i) before January 1, 2021, within 45 days after the day on which the municipality
515	receives a petition for the action; and
516	(ii) after January 1, 2021, in accordance with Subsection 10-9a-509.5(2).
517	Section 6. Section 17-27a-525 is amended to read:
518	17-27a-525. Cannabis production establishments and medical cannabis
519	pharmacies.
520	(1) As used in this section:
521	(a) "Cannabis production establishment" means the same as that term is defined in

522	Section 4-41a-102 and includes a closed-door medical cannabis pharmacy.
523	(b) "Closed-door medical cannabis pharmacy" means the same as that term is defined
524	in Section 4-41a-102.
525	[(b)] (c) "Industrial hemp producer licensee" means the same as the term "licensee" is
526	defined in Section 4-41-102.
527	[(c)] (d) "Medical cannabis pharmacy" means the same as that term is defined in
528	Section 26B-4-201.
529	(2) (a) (i) A county may not regulate a cannabis production establishment or a medical
530	cannabis pharmacy in conflict with:
531	(A) Title 4, Chapter 41a, Cannabis Production Establishments and Pharmacies, and
532	applicable jurisprudence; and
533	(B) this chapter.
534	(ii) A county may not regulate an industrial hemp producer licensee in conflict with:
535	(A) Title 4, Chapter 41, Hemp and Cannabinoid Act, and applicable jurisprudence; and
536	(B) this chapter.
537	(b) The Department of Agriculture and Food has plenary authority to license programs
538	or entities that operate a cannabis production establishment or a medical cannabis pharmacy.
539	(3) (a) Within the time period described in Subsection (3)(b), a county shall prepare
540	and adopt a land use regulation, development agreement, or land use decision in accordance
541	with this title and:
542	(i) regarding a cannabis production establishment, Section 4-41a-406; or
543	(ii) regarding a medical cannabis pharmacy, Section [4-41a-110] 4-41a-1105.
544	(b) A county shall take the action described in Subsection (3)(a):
545	(i) before January 1, 2021, within 45 days after the day on which the county receives a
546	petition for the action; and
547	(ii) after January 1, 2021, in accordance with Subsection 17-27a-509.5(2).
548	Section 7. Section <b>26B-1-435</b> is amended to read:
549	26B-1-435. Medical Cannabis Policy Advisory Board creation Membership
550	Duties.
551	(1) There is created within the department the Medical Cannabis Policy Advisory
552	Board.

553	(2) (a) The advisory board shall consist of the following members:
554	(i) appointed by the executive director:
555	(A) a qualified medical provider who has recommended medical cannabis to at least
556	100 patients [who have a medical cannabis patient card at the time of appointment] before
557	being appointed;
558	(B) a medical research professional;
559	(C) a mental health specialist;
560	(D) an individual who represents an organization that advocates for medical cannabis
561	patients;
562	(E) an individual who holds a medical cannabis patient card; and
563	(F) a member of the general public who does not hold a medical cannabis card; and
564	(ii) appointed by the commissioner of the Department of Agriculture and Food:
565	(A) an individual who owns or operates a licensed cannabis cultivation facility;
566	(B) an individual who owns or operates a licensed medical cannabis pharmacy; and
567	(C) a law enforcement officer.
568	(b) The commissioner of the Department of Agriculture and Food shall ensure that at
569	least one individual appointed under Subsection (2)(a)(ii)(A) or (B) also owns or operates a
570	licensed cannabis processing facility.
571	(3) (a) Subject to Subsection (3)(b), a member of the advisory board shall serve for a
572	four year term.
573	(b) When appointing the initial membership of the advisory board, the executive
574	director and the commissioner of the Department of Agriculture and Food shall coordinate to
575	appoint four advisory board members to serve a term of two years to ensure that approximately
576	half of the board is appointed every two years.
577	(4) (a) If an advisory board member is no longer able to serve as a member, a new
578	member shall be appointed in the same manner as the original appointment.
579	(b) A member appointed in accordance with Subsection (4)(a) shall serve for the
580	remainder of the unexpired term of the original appointment.
581	(5) (a) A majority of the advisory board members constitutes a quorum.
582	(b) The action of a majority of a quorum constitutes an action of the advisory board.
583	(c) [The] For a term lasting one year, the advisory board shall annually designate [one

584	of the advisory board's members of the advisory board to serve as chair [for a
585	one-year period.] and vice-chair.
586	(d) When designating the chair and vice-chair, the advisory board shall ensure that at
587	least one individual described Subsection (2)(a)(i) is appointed as chair or vice-chair.
588	(6) An advisory board member may not receive compensation or benefits for the
589	member's service on the advisory board but may receive per diem and reimbursement for travel
590	expenses incurred as an advisory board member in accordance with:
591	(a) Sections 63A-3-106 and 63A-3-107; and
592	(b) rules made by the Division of Finance pursuant to Sections 63A-3-106 and
593	63A-3-107.
594	(7) The department shall:
595	(a) provide staff support for the advisory board; and
596	(b) assist the advisory board in conducting meetings.
597	(8) The advisory board may recommend:
598	(a) to the department or the Department of Agriculture and Food changes to current or
599	proposed medical cannabis rules or statutes;
600	(b) to the appropriate legislative committee whether the advisory board supports a
601	change to medical cannabis statutes.
602	(9) The advisory board shall:
603	(a) review any draft rule that is authorized under this chapter or Title 4, Chapter 41a,
604	Cannabis Production Establishments and Pharmacies;
605	(b) consult with the Department of Agriculture and Food regarding the issuance of an
606	additional:
607	(i) cultivation facility license under Section 4-41a-205; or
608	(ii) pharmacy license under Section 4-41a-1005;
609	(c) consult with the department regarding cannabis patient education;
610	(d) consult regarding the reasonableness of any fees set by the department or the
611	Department of Agriculture and Food that pertain to the medical cannabis program; and
612	(e) consult regarding any issue pertaining to medical cannabis when asked by the
613	department or the Utah Department of Agriculture and Food.
614	Section 8 Section 26B-4-219 is amended to read:

615	26B-4-219. Pharmacy medical providers Registration Continuing education.
616	(1) (a) A medical cannabis pharmacy:
617	(i) shall employ a pharmacist who is licensed under Title 58, Chapter 17b, Pharmacy
618	Practice Act, as a pharmacy medical provider;
619	(ii) may employ a physician who has the authority to write a prescription and is
620	licensed under Title 58, Chapter 67, Utah Medical Practice Act, or Title 58, Chapter 68, Utah
621	Osteopathic Medical Practice Act, as a pharmacy medical provider;
622	(iii) shall ensure that a pharmacy medical provider described in Subsection (1)(a)(i)
623	works onsite during all business hours; and
624	(iv) shall designate one pharmacy medical provider described in Subsection (1)(a)(i) as
625	the pharmacist-in-charge to oversee the operation of and generally supervise the medical
626	cannabis pharmacy.
627	(b) The pharmacist-in-charge shall determine which cannabis and cannabis products
628	the medical cannabis pharmacy maintains in the medical cannabis pharmacy's inventory.
629	[(b)] (c) An individual may not serve as a pharmacy medical provider unless the
630	department registers the individual as a pharmacy medical provider in accordance with
631	Subsection (2).
632	(2) (a) The department shall, within 15 days after the day on which the department
633	receives an application from a medical cannabis pharmacy on behalf of a prospective pharmacy
634	medical provider, register and issue a pharmacy medical provider registration card to the
635	prospective pharmacy medical provider if the medical cannabis pharmacy:
636	(i) provides to the department:
637	(A) the prospective pharmacy medical provider's name and address;
638	(B) the name and location of the licensed medical cannabis pharmacy where the
639	prospective pharmacy medical provider seeks to act as a pharmacy medical provider;
640	(C) a report detailing the completion of the continuing education requirement described
641	in Subsection (3); and
642	(D) evidence that the prospective pharmacy medical provider is a pharmacist who is
643	licensed under Title 58, Chapter 17b, Pharmacy Practice Act, or a physician who has the
644	authority to write a prescription and is licensed under Title 58, Chapter 67, Utah Medical
645	Practice Act, or Title 58, Chapter 68, Utah Osteopathic Medical Practice Act; and

646 (ii) pays a fee to the department in an amount that, subject to Subsection 26B-1-310(5), 647 the department sets in accordance with Section 63J-1-504. 648 (b) The department may not register a recommending medical provider as a pharmacy 649 medical provider. 650 (3) (a) A pharmacy medical provider shall complete the continuing education described 651 in this Subsection (3) in the following amounts: 652 (i) as a condition precedent to registration, four hours; and (ii) as a condition precedent to renewal of the registration, four hours every two years. 653 654 (b) In accordance with Subsection (3)(a), the pharmacy medical provider shall: 655 (i) complete continuing education: 656 (A) regarding the topics described in Subsection (3)(d); and 657 (B) offered by the department under Subsection (3)(c) or an accredited or approved 658 continuing education provider that the department recognizes as offering continuing education 659 appropriate for the medical cannabis pharmacy practice; and 660 (ii) make a continuing education report to the department in accordance with a process 661 that the department establishes by rule, in accordance with Title 63G, Chapter 3, Utah 662 Administrative Rulemaking Act, and in collaboration with the Division of Professional 663 Licensing and: 664 (A) for a pharmacy medical provider who is licensed under Title 58, Chapter 17b, 665 Pharmacy Practice Act, the Board of Pharmacy; 666 (B) for a pharmacy medical provider licensed under Title 58, Chapter 67, Utah Medical Practice Act, the Physicians Licensing Board; and 667 668 (C) for a pharmacy medical provider licensed under Title 58, Chapter 68, Utah 669 Osteopathic Medical Practice Act, the Osteopathic Physician and Surgeon's Licensing Board. 670 (c) The department may, in consultation with the Division of Professional Licensing, 671 develop the continuing education described in this Subsection (3). 672 (d) The continuing education described in this Subsection (3) may discuss: 673 (i) the provisions of this part; 674 (ii) general information about medical cannabis under federal and state law; 675 (iii) the latest scientific research on the endocannabinoid system and medical cannabis, 676 including risks and benefits;

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applicable rules.

677 (iv) recommendations for medical cannabis as it relates to the continuing care of a 678 patient in pain management, risk management, potential addiction, and palliative care; or 679 (v) best practices for recommending the form and dosage of [a] medical cannabis 680 [product] based on the qualifying condition underlying a medical cannabis recommendation. 681 (4) (a) A pharmacy medical provider registration card expires two years after the day 682 on which the department issues or renews the card. 683 (b) A pharmacy medical provider may renew the provider's registration card if the 684 provider: 685 (i) is eligible for a pharmacy medical provider registration card under this section; 686 (ii) certifies to the department in a renewal application that the information in 687 Subsection (2)(a) is accurate or updates the information; 688 (iii) submits a report detailing the completion of the continuing education requirement 689 described in Subsection (3); and 690 (iv) pays to the department a renewal fee in an amount that: 691 (A) subject to Subsection 26B-1-310(5), the department sets in accordance with 692 Section 63J-1-504; and 693 (B) may not exceed the cost of the relatively lower administrative burden of renewal in 694 comparison to the original application process. 695 (5) (a) Except as provided in Subsection (5)(b), a person may not advertise that the 696 person or another person dispenses medical cannabis. 697 (b) Notwithstanding Subsection (5)(a) and Section 4-41a-109, a registered pharmacy 698 medical provider may advertise the following: 699 (i) a green cross; 700 (ii) that the person is registered as a pharmacy medical provider and dispenses medical 701 cannabis; or 702 (iii) a scientific study regarding medical cannabis use. 703 (6) (a) The department may revoke a pharmacy medical provider's registration for a 704 violation of this chapter.

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(b) The department may inspect patient records held by a medical cannabis pharmacy

to ensure a pharmacy medical provider is practicing in accordance with this chapter and

Section 9. Section **26B-4-231** is amended to read:

#### 26B-4-231. Partial filling -- Pharmacy medical provider directions of use.

- (1) As used in this section, "partially fill" means to provide less than the full amount of cannabis or cannabis product that the recommending medical provider recommends, if the recommending medical provider recommended specific dosing guidelines.
- (2) A pharmacy medical provider may partially fill a recommendation for a medical cannabis treatment at the request of the recommending medical provider who issued the medical cannabis treatment recommendation or the medical cannabis cardholder.
- (3) The department shall make rules, in collaboration with the Division of Professional Licensing and the Board of Pharmacy and in accordance with Title 63G, Chapter 3, Utah Administrative Rulemaking Act, specifying how to record the date, quantity supplied, and quantity remaining of a partially filled medical cannabis treatment recommendation.
- (4) A pharmacy medical provider who is a pharmacist may, upon the request of a medical cannabis cardholder, determine different dosing guidelines, subject to the dosing limits in Subsection 4-41a-1102(2), to fill the quantity remaining of a partially filled medical cannabis treatment recommendation if:
- (a) the pharmacy medical provider determined dosing guidelines for the partial fill under Subsection 4-41a-1102(5) or (6); and
  - (b) the medical cannabis cardholder reports that:
- (i) the partial fill did not substantially affect the qualifying condition underlying the medical cannabis recommendation; or
- (ii) the patient experienced an adverse reaction to the partial fill or was otherwise unable to successfully use the partial fill.
- (5) If a recommending medical provider recommends treatment with medical cannabis but wishes for the pharmacy medical provider to determine directions of use and dosing guidelines:
- (a) the recommending medical provider shall provide to the pharmacy medical provider, either through the state electronic verification system or through a medical cannabis pharmacy's recording of a recommendation under the order of a limited medical provider, any of the following information that the recommending medical provider feels would be needed to provide appropriate directions of use and dosing guidelines:

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739	(i) information regarding the qualifying condition underlying the recommendation;
740	(ii) information regarding prior treatment attempts with medical cannabis; and
741	(iii) portions of the patient's current medication list; and
742	(b) before the relevant medical cannabis cardholder may obtain medical cannabis, the
743	pharmacy medical provider shall:
744	(i) review pertinent medical records, including the recommending medical provider
745	documentation described in Subsection (5)(a); and
746	(ii) [unless the pertinent medical records show directions of use and dosing guidelines
747	from a state central patient portal medical provider in accordance with Subsection (6),] after
748	completing the review described in Subsection (5)(b)(i) and consulting with the recommending
749	medical provider as needed, determine the best course of treatment through consultation with
750	the cardholder regarding:
751	(A) the patient's qualifying condition underlying the recommendation from the
752	recommending medical provider;
753	(B) indications for available treatments;
754	(C) directions of use and dosing guidelines; and
755	(D) potential adverse reactions.
756	Section 10. Repealer.
757	This bill repeals:
758	Section 26B-1-435.1, Medical Cannabis Policy Advisory Board duties.
759	Section 11. Effective date.
760	This bill takes effect on May 1, 2024.