Minnesota’s Medical Cannabis Therapeutic Research Act

This information brief explains the Medical Cannabis Therapeutic Research Act, Minnesota Statutes, sections 152.22 to 152.37, enacted in 2014 and amended through the 2017 first special session. The act was not amended in 2018. The act established a patient registry program that allows qualifying patients to use and possess cannabis for medical purposes. A brief history of medical cannabis legislation in Minnesota is also provided.

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Overview of the Law

The Medical Cannabis Therapeutic Research Act was enacted in May 2014. The law established a patient registry program, administered by the Minnesota Department of Health (MDH), which allows qualifying patients to possess and use cannabis for medical use.

The law allows two manufacturers to be registered in the state. Each manufacturer must have one manufacturing facility and four distribution sites throughout the state.

The manufacturers may only distribute medical cannabis in pill, liquid, or topical form, and patients may only possess medical cannabis in those limited forms.

Qualifying medical conditions include:

1. Cancer*
2. Glaucoma
3. HIV/AIDS
4. Tourette’s syndrome
5. Amyotrophic lateral sclerosis (ALS)
6. Seizures
7. Severe and persistent muscle spasms
8. Inflammatory bowel disease, including Crohn’s disease
9. Terminal illness with life expectancy of under one year*
10. Intractable pain
11. Post-traumatic stress disorder
12. Autism spectrum disorder
13. Obstructive sleep apnea
14. Any other condition or its treatment approved by the commissioner (subject to legislative oversight)

* Illness or treatment must produce one or more of the following: (1) severe or chronic pain; (2) nausea or severe vomiting; or (3) cachexia or severe wasting.

The general design of the registry program is as follows:

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1 Laws 2014, ch. 311; codified as Minn. Stat. §§ 152.22 to 152.37.
General Design of the Registry Program

1. Patient diagnosed with qualifying medical condition

2. Patient sends annual application to MDH

3. MDH issues registry verification to patient, health care practitioner, and manufacturer

4. Health care practitioner continues treatment of patient for qualifying condition

5. Patient obtains medical cannabis

6. Manufacturer distributes medical cannabis to patient

7. Practitioner and manufacturer report to MDH

8. MDH submits research report to legislature and major scientific journals
General Design of the Registry Program

The general design of the registry program is explained briefly below, but many aspects are more fully detailed in subsequent sections of this information brief.

Patient diagnosed with qualifying medical condition

Prior to applying to participate in the registry program, a patient must be diagnosed by a health care practitioner with one or more of the qualifying medical conditions.

Patient sends annual application to MDH

Once the patient receives certification of a diagnosis with a qualifying medical condition from a health care practitioner, the patient must apply to the Minnesota Department of Health (MDH) to be a part of the registry program. The patient must submit this application, along with an application fee, on an annual basis.

MDH issues a registry verification to patient, health care practitioner, and manufacturer

Once the patient has been accepted into the registry program, MDH issues a registry verification listing the patient’s information, along with the information of the patient’s registered designated caregiver or parent or legal guardian, if applicable. The registry verification is issued to the patient, the patient’s listed health care practitioner, and the manufacturer as proof of the patient’s participation in the registry program.

Health care practitioner continues treatment of qualifying condition

As part of the health care practitioner’s duties, the practitioner must continue to treat the qualifying medical condition of the patient.

Manufacturer distributes medical cannabis to patient

A manufacturer may only distribute medical cannabis to a person listed on the patient’s registry verification. Final approval for distribution must be made by a licensed pharmacist after a consultation with the patient.

Patient obtains medical cannabis from manufacturer

A patient may only obtain medical cannabis from a registered manufacturer. If a patient has a registered designated caregiver or parent or legal guardian listed on the registry verification, that person may also obtain the medical cannabis from the manufacturer on the patient’s behalf.

Reports to MDH

The health care practitioner is required to report the patient’s health records to MDH through the registry program. The manufacturer is also required to submit a report to MDH containing various information.
MDH submits reports to legislature and major medical journals

MDH is required to conduct research on the information in the registry program and submit reports to certain legislative committees as well as major medical journals.

The Patient Registry Program

MDH Program Administration

MDH and its commissioner are responsible for administering the patient registry program. The Office of Medical Cannabis within MDH administers the program and ensures patients, health care practitioners, and manufacturers comply with state statutes and rules governing the program.

Range of compounds\(^2\)

MDH must annually review existing medical and scientific literature on the recommended range of dosages and chemical compounds for each qualifying medical condition and must publicly report that review. The most recent review was published in May 2018 and is posted on the MDH website.\(^3\)

Rulemaking authority\(^4\)

MDH adopted the initial medical cannabis rules using the expedited rulemaking process under Minnesota Statutes, section 14.389. MDH was required to have in place rules necessary for the manufacturers to begin distributing medical cannabis to patients by July 1, 2015. The initial rules were adopted January 20, 2015, and are found in Minnesota Rules, chapter 4770. MDH must use the standard rulemaking process for any subsequent rulemaking.

Adverse incidents\(^5\)

MDH adopted rules to require law enforcement and emergency medical personnel to report incidents when individuals not authorized to use medical cannabis are found in possession of medical cannabis. The department also adopted rules requiring law enforcement, health care professionals, registered patients, caregivers, and manufacturers to report serious adverse incidents involving medical cannabis, including incidents involving an overdose of medical cannabis.

\(^2\) Minn. Stat. § 152.25, subd. 2.

\(^3\) http://www.health.state.mn.us/topics/cannabis/practitioners/dosagesandcompositions2018.pdf.

\(^4\) Minn. Stat. §§ 152.26; 152.261.

\(^5\) Minn. Stat. § 152.261; Minn. Rules, parts 4770.4004 and 4770.4010.
Adding additional allowable forms and qualifying medical conditions

The commissioner may add to the list of qualifying medical conditions and also add to the list of allowable forms of medical cannabis. The commissioner is prohibited, however, from adding smoking as an allowable form. To add an additional form or condition, the commissioner must notify the chairs and ranking minority members of the legislative committees having jurisdiction over health and human services as to the reasons for the addition. This notice must include any public comments the commissioner has received and any guidance the commissioner has received from the task force on medical cannabis research. The notification must be given by January 15 of the year the commissioner wishes to make the change. The change becomes effective August 1 of that year unless the legislature provides otherwise by law.

Financial audit and examination

MDH may inspect the manufacturer’s financial documents through an annual certified financial audit or through an examination of its business affairs. (For more on manufacturer financial audits, see page 14).

Reports

The commissioner is required to regularly update the Task Force on Medical Cannabis Therapeutic Research and the chairs and ranking minority members of certain legislative committees regarding any changes in federal law or regulation of medical cannabis. The commissioner may also submit medical research collected through the registry program to federal agencies with regulatory authority over medical cannabis in order to demonstrate the effectiveness of medical cannabis for treating qualifying conditions. The commissioner must also submit findings from the registry program to both the legislature and major scientific journals.

Patients

Participation in the registry program

To participate in the registry program, a patient must first consult with a health care practitioner to determine whether the patient has one or more of the qualifying medical conditions. Qualifying medical conditions include:

1. Cancer
2. Glaucoma
3. HIV/AIDS

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6 Minn. Stat. § 152.27, subd. 2, para. (b).
7 Minn. Stat. § 152.37.
8 Minn. Stat. § 152.25, subd. 4.
9 Minn. Stat. § 152.22, subd. 14; additional conditions added by Commissioner of Health.
4. Tourette’s syndrome
5. Amyotrophic lateral sclerosis (ALS)
6. Seizures
7. Severe and persistent muscle spasms
8. Inflammatory bowel disease, including Crohn’s disease
9. Terminal illness with life expectancy of under one year*
10. Intractable pain
11. Post-traumatic stress disorder
12. Autism spectrum disorder
13. Obstructive sleep apnea
14. Any other condition or its treatment approved by the commissioner (subject to legislative oversight)

* Illness or treatment must produce one or more of the following: (1) severe or chronic pain; (2) nausea or severe vomiting; or (3) cachexia or severe wasting.

Laws 2014, chapter 311, section 20, required the Commissioner of Health to consider adding intractable pain to the list of qualifying medical conditions before considering adding any other condition to the list. The commissioner added intractable pain to the list of qualifying medical conditions using the procedure in Minnesota Statutes, section 152.27, subdivision 2, paragraph (b). Patients with intractable pain were eligible to enroll in the registry program beginning July 1, 2016, and to receive medical cannabis beginning August 1, 2016. Intractable pain is defined in Minnesota Statutes, section 152.125, subdivision 1. In the following years, the commissioner added post-traumatic stress disorder, autism spectrum disorder, and obstructive sleep apnea to the list of qualifying medical conditions.

Following diagnosis of a qualifying medical condition, the patient must submit an application and application fee to MDH to enroll in the registry program. The application must include a health care practitioner’s certification of diagnosis and other forms required by MDH. The application must also include the name, mailing address, and date of birth of the patient, the designated caregiver if the patient is unable to self-administer medication, and the patient’s parent or legal guardian if the parent or legal guardian will act as caregiver.

As part of the yearly application, the patient is required to pay an application fee of $200. If the patient attests to receiving Social Security disability or Supplemental Security Insurance payments, or being enrolled in Medical Assistance or MinnesotaCare, the patient’s yearly fee is $50.

The commissioner must approve or deny an application within 30 days of receiving the application and fee. Once the application is approved by MDH, the patient receives a registry verification.

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10 Minn. Stat. § 152.27, subd. 3.
11 Minn. Stat. § 152.35.
Reasons to deny participation in the registry program\textsuperscript{12}

The commissioner is authorized to deny a patient entry into the registry program only if the patient:

- does not have a certification from a health care practitioner of diagnosis with a qualifying medical condition;
- does not provide the required information or signed disclosures;
- has previously been removed from the registry program for a violation of patient duties or the commission of a crime related to medical cannabis; or
- provides false information.

If a patient is denied entry, the commissioner must give the patient a written reason for the denial. A denial is considered a final decision of the commissioner and is subject to judicial review under the Administrative Procedure Act, pursuant to chapter 14.

Responsibilities during participation\textsuperscript{13}

To maintain enrollment in the registry program, the patient must resubmit a copy of the certification of diagnosis to MDH on a yearly basis and pay a yearly application fee. Patients must also continue to receive regularly scheduled treatment for that qualifying medical condition and report changes in that condition to their health care practitioner throughout enrollment in the registry program.

Registered designated caregivers\textsuperscript{14}

A patient is permitted to have a registered designated caregiver if the patient’s health care practitioner certifies that the patient has a developmental or physical disability that prevents the patient from either self-administering the medication or acquiring the medication from a distribution facility. The registered designated caregiver must agree, in writing, to act as the patient’s caregiver. As conditions of registration, the caregiver must:

- be at least 21 years of age;
- agree to only possess medical cannabis for purposes of assisting the patient; and
- agree to not be a caregiver for more than one patient, unless the patients reside in the same residence.

\textsuperscript{12} Minn. Stat. § 152.27, subd. 6.
\textsuperscript{13} Minn. Stat. § 152.30.
\textsuperscript{14} Minn. Stat. § 152.27, subd. 4.
Registered designated caregivers are subject to a criminal background check. If the caregiver has a disqualifying felony offense, the commissioner is prohibited from registering that caregiver. Registered designated caregivers are also subject to criminal sanctions for diversion of medical cannabis in the same way as patients. (For more information on that criminal sanction, see page 10).

**Parents or legal guardians**

A parent or legal guardian, if listed on the registry verification as a patient’s caregiver, may act as a patient’s caregiver without having to register as a designated caregiver. Parents and legal guardians are also subject to criminal sanctions for diversion of medical cannabis in the same way as patients. (For more information on that criminal sanction, see page 10).

**Civil and criminal protections**

*Presumption.* Once a patient enrolls in the registry program, there is a presumption that the patient is engaging in the authorized use of medical cannabis. This presumption may be rebutted by evidence that the patient’s use of medical cannabis was not for the purpose of treating the patient’s qualifying medical condition or associated symptoms.

*Exemption from criminal sanctions for use or possession.* Patients who use or possess medical cannabis, and registered designated caregivers and parents and legal guardians who possess medical cannabis, are exempt from criminal sanctions for use or possession of a controlled substance.

*Forfeiture.* Medical cannabis and associated property are not subject to forfeiture under Minnesota law.

*Search warrant needed to access registry.* Law enforcement personnel must have a valid search warrant to access the patient registry.

*Use of registry verification or application to support a search.* A person’s possession of a registry verification or registry program application does not constitute probable cause or reasonable suspicion, and cannot be used to support a search of the person or property.

*Circumstances in which penalties still apply.* Although a patient is exempt from criminal sanctions for possession under Minnesota law, the patient is not exempt from penalties for:

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15 Disqualifying felony offenses are defined as violations of any state or federal controlled substance law that would be a felony in Minnesota, regardless of the sentence imposed, unless the commissioner determines that the conviction was for either the use or assistance with use of medical cannabis. Minn. Stat. § 152.22, subd. 3.

16 Minn. Stat. § 152.27, subd. 5.

17 See generally Minn. Stat. § 152.32, subd. 2.
(1) undertaking any task under the influence of medical cannabis that would constitute negligence or professional malpractice;

(2) possessing or using medical cannabis:
   a. on a school bus or van;
   b. on the grounds of any preschool, primary school, or secondary school;
   c. in any correctional facility; or
   d. on the grounds of any child care facility or home daycare;

(3) vaporizing medical cannabis:
   a. on any form of public transportation;
   b. where the vapor may be inhaled by a nonpatient minor child, or
   c. in a public place, including any indoor or outdoor area used by or open to the general public or a place of employment;\textsuperscript{18} and

(4) operating, navigating, or being in actual physical control of any motor vehicle, aircraft, train, or motorboat, or working on transportation property, equipment, or facilities while under the influence of medical cannabis.\textsuperscript{19}

Criminal sanctions

\textit{Diversion of medical cannabis.} A patient, registered designated caregiver, or, if listed on the registry verification, a parent or legal guardian of a patient who intentionally sells or otherwise transfers medical cannabis to a person other than a patient, registered designated caregiver, or, if listed on the registry verification, a parent or legal guardian, is guilty of a felony. This crime is punishable by imprisonment for not more than two years or payment of a fine of not more than $3,000, or both.\textsuperscript{20}

\textit{False statements.} A person who intentionally makes a false statement to law enforcement about any fact or circumstance relating to the use of medical cannabis in order to avoid arrest or prosecution is guilty of a misdemeanor. This crime is punishable by imprisonment for up to 90 days, payment of a fine of not more than $1,000, or both, in addition to any other applicable penalty under the law. A patient or a registered designated caregiver convicted of this crime is disqualified from any further participation in the registry program.\textsuperscript{21}

Patient discrimination prohibited\textsuperscript{22}

A patient is protected from discrimination in a variety of circumstances.

\textsuperscript{18} See Minn. Stat. § 144.413, subd. 1b, for a definition of place of employment.\textsuperscript{19}

Minn. Stat. § 152.23.

\textsuperscript{20} See generally Minn. Stat. § 152.33, subd. 2.

\textsuperscript{21} Minn. Stat. § 152.33, subd. 3.

\textsuperscript{22} Minn. Stat. § 152.32, subd. 3.
Enrollment in school. A school cannot refuse to enroll a person or otherwise penalize a person solely because the person is enrolled in the medical cannabis registry program. This prohibition does not apply if enrolling the person would cause the school to violate federal law or cause the school to lose a monetary or licensing-related benefit under federal law.

Leasing. A landlord cannot refuse to lease to a person or otherwise penalize a person solely because the person is enrolled in the medical cannabis registry program. This prohibition does not apply if leasing to the person would cause the landlord to violate federal law or cause the landlord to lose a monetary or licensing-related benefit under federal law.

Medical care. A patient’s use of medical cannabis under the registry program is considered the authorized use of medication for purposes of medical care, including organ transplants. This use of medical cannabis is not use of an illicit substance and does not disqualify a patient from needed medical care.

Employment. An employer is prohibited from discriminating against a person in hiring, termination, or any term or condition of employment, or otherwise penalize the employee based on:

- the person’s status as a patient in the registry program; or
- a patient’s positive drug test for cannabis components or metabolites, unless the patient used, possessed, or was impaired by medical cannabis while on the employer’s premises or during the hours of employment.

An employer is not required to take actions, however, that would violate federal law or cause the employer to lose a federal monetary or licensing-related benefit. If an employee is required to take a drug test for the employer pursuant to section 181.953, the employee may present verification of enrollment in the patient registry as part of the employee’s explanation of a positive urine test under section 181.953, subdivision 6.

Custody/Visitation. A person cannot be denied custody of a minor child or visitation rights with a minor child solely based on a person’s status as a patient enrolled in the registry program. The law also provides that there is no presumption of neglect or child endangerment for conduct allowed under the registry program, unless the person’s behavior creates an unreasonable danger to the safety of the child as established by clear and convincing evidence.

Federally approved clinical trials

The Commissioner of Health must provide information to all patients about the existence of any federally approved clinical trials for the treatment of that patient’s qualifying condition with medical cannabis. The commissioner may prohibit enrollment of a patient in the registry program if that patient is simultaneously enrolled in a federally approved clinical trial for the treatment of the patient’s qualifying condition with medical cannabis.

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23 Minn. Stat. § 152.24.
Manufacturers

Registration

Initial registration and reregistration\textsuperscript{24}

On December 1, 2014, MDH registered two medical cannabis manufacturers: LeafLine Labs and Minnesota Medical Solutions. Manufacturers are subject to re-registration every two years. As a condition of initial registration, each manufacturer agreed to begin distribution of medical cannabis to patients by July 1, 2015, and comply with other requirements under the law.

MDH is required to consider the following factors when determining which manufacturers to register or reregister:

- Technical expertise in cultivation of medical cannabis and conversion into allowable forms
- The qualifications of the manufacturer’s employees
- The long-term financial stability of the manufacturer
- The ability to provide appropriate security measures on the premises of the manufacturer
- Whether the manufacturer has demonstrated an ability to meet the medical cannabis production needs required by the registry program
- The manufacturer’s projection and ongoing assessment of patient fees

The commissioner may accept additional applications for registration if one of the registered manufacturers ceases to be registered.

Enforcement actions against manufacturer registrations\textsuperscript{25}

The commissioner has authority to revoke or not renew a manufacturer’s registration, or to deny consent for a manufacturer to transfer its registration to another entity. In particular, the commissioner is authorized to not renew a manufacturer’s registration if certain individuals affiliated with the manufacturer intentionally divert medical cannabis to a person other than allowed by law. Before revoking, not renewing, or denying consent to transfer a registration, the commissioner must provide written notice to the affected manufacturer, and the manufacturer may request a contested case hearing. In addition, the commissioner may temporarily suspend a manufacturer’s registration for up to 90 days in certain circumstances. If the commissioner takes any enforcement action against a manufacturer that affects the ability of patients, designated caregivers, and parents and legal guardians to obtain medical cannabis from that manufacturer,

\textsuperscript{24} Minn. Stat. § 152.25, subd. 1.

\textsuperscript{25} Minn. Stat. § 152.25, subds. 1, 1a, 1b,
the commissioner must give notice of the enforcement action and alternatives for obtaining medical cannabis to affected patients, designated caregivers, and parents and legal guardians.

**Regulation**

**Fees**\(^{26}\)

The Commissioner of Health collects from manufacturers an annual fee for the cost incurred by MDH for the regulation and inspection of the manufacturer for that year. The yearly fee is established by the Commissioner of Health. Each manufacturer is allowed to charge patients enrolled in the program a “reasonable fee” for operating costs of the manufacturer. Manufacturers are allowed, but not required, to establish a sliding scale of patient fees based on a patient’s household income. Manufacturers may also accept private donations in order to reduce patient fees.

**Operating documents**\(^{27}\)

A manufacturer’s operating documents must include procedures for oversight, procedures to ensure accurate recordkeeping, and procedures for appropriate security measures to deter theft and unauthorized entrance into areas that contain medical cannabis.

**Location of facilities**\(^{28}\)

Each manufacturer is required to have four distribution facilities and one production facility (the production facility may be at the same location as a distribution facility). A manufacturer cannot operate more than four or fewer than four distribution facilities, and the distribution facilities must be located throughout the state based on geographical need in order to improve patient access. No facility may be within 1,000 feet of a public or private school that was in existence prior to the manufacturer’s registration with MDH. Distribution facilities are currently located in Bloomington, Eagan, Hibbing, Minneapolis, Moorhead, Rochester, St. Cloud, and St. Paul.

**Employees**\(^{29}\)

A manufacturer is prohibited from employing any person under the age of 21 or any person who has been convicted of a disqualifying felony offense. However, a manufacturer may employ a person who has been convicted of a disqualifying felony offense if the Commissioner of Health

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\(^{26}\) Minn. Stat. § 152.35.

\(^{27}\) Minn. Stat. § 152.29, subd. 1, para. (c).

\(^{28}\) Minn. Stat. § 152.29, subd. 1, paras. (a) and (j).

\(^{29}\) Minn. Stat. § 152.29, subd. 1, para. (l).

\(^{30}\) A disqualifying felony offense is defined as a violation of any state or federal controlled substance crime that would be a felony under Minnesota law, whether or not the offense was committed in Minnesota and regardless of the sentence imposed. Minn. Stat. § 152.22, subd. 3.
determines the conviction was for the use of or assistance with the use of medical cannabis. All potential employees must undergo a criminal history background check through the Bureau of Criminal Apprehension prior to working with the manufacturer.

Due to distribution requirements, manufacturers must also employ at least one pharmacist licensed in Minnesota. The pharmacist employees must be the only employees who give final approval for distribution of medical cannabis and must consult with the patient before distributing the medical cannabis.31

Any employee of the manufacturer involved in delivering medical cannabis or medical cannabis products from one location to another must carry identification showing that the person is an employee of the manufacturer.32

Security33

Manufacturers must have certain security measures at all distribution sites as well as the production site. These security measures include:

- a fully operational security alarm system;
- facility access controls;
- perimeter intrusion detection systems; and
- a personnel identification system.

Contract with an independent laboratory34

Each manufacturer must contract with an independent laboratory approved by the Commissioner of Health to test the manufacturer’s medical cannabis for content, contamination, and consistency. The cost of this contract must be paid by the manufacturer and is subject to any additional requirements set by the Commissioner of Health.

Inspections35

Manufacturers are subject to reasonable inspections by the Commissioner of Health. Each manufacturer must keep detailed financial records in a manner approved by the commissioner and make these records available for the commissioner's review. In addition, the manufacturers must submit to the commissioner the results of an annual financial audit conducted by an independent certified public accountant, paid for by the manufacturer. The commissioner may

31 Minn. Stat. § 152.29, subd. 3, paras. (a) and (c).
32 Minn. Stat. § 152.29, subd. 3, para. (d).
33 Minn. Stat. § 152.29, subd. 1, para. (d).
34 Minn. Stat. § 152.29, subd. 1, para. (b).
35 Minn. Stat. §§ 152.29, subd. 1, para. (g); 152.37.
require a second financial audit by a certified public accountant chosen by the commissioner, also at the expense of the manufacturer.

The commissioner or the commissioner’s designee may examine the business affairs of the manufacturer, including a review of the manufacturer’s financing, budgets, revenues, sales, and pricing. The commissioner may retain outside professionals, such as attorneys and certified public accountants, to conduct or assist with this examination, but may not retain the same certified public accountant as used in the annual audit. If the commissioner conducts this examination, the commissioner must complete a report and provide a copy to the manufacturer and post a copy on the department’s website. All data collected during this examination, except for the public report, are private data on individuals or nonpublic data.

**Monthly report to MDH**

Each manufacturer must submit a monthly report to MDH. The report must include the following information for each patient served in the prior month:

- the amount and dosages of medical cannabis distributed
- the chemical composition of the medical cannabis
- the tracking number assigned to any medical cannabis distributed

These reports are not public. MDH compares data in the reports to data in the medical cannabis registry to ensure that the medical cannabis registry has a record of all medical cannabis transactions. MDH has also used these reports to fill in incomplete records in the registry.

**Production**

**Requirements**

Each manufacturer must produce a reliable and ongoing supply of medical cannabis to patients and process the medical cannabis into an allowed form prior to its distribution. Production of medical cannabis must be done in one location and must be in an enclosed and locked facility.

**Allowable forms**

Medical cannabis may only be distributed as a pill or liquid, including oil, or as a topical formulation, such as a patch, lotion, cream, gel, and ointment. The first two allowable forms are specified in statute. The commissioner added topical formulations as an allowable form, and patients were authorized to use topical formulations as of August 1, 2017.

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36 Minn. Stat. § 152.29, subd. 4.
37 Minn. Stat. § 152.29, subd. 2.
38 Minn. Stat. § 152.22, subd. 6.
The Commissioner of Health may allow other forms, except smoking. Any addition by the commissioner must follow the procedure in Minnesota Statutes, section 152.27, subdivision 2, paragraph (b), and is subject to legislative oversight.

**Deadlines for distribution**\(^{39}\)

Each manufacturer had to begin distribution to patients from at least one distribution site by July 1, 2015. Distribution had to occur from all four of the manufacturer’s distribution sites by July 1, 2016.

**Distribution**

**What may be distributed**\(^{40}\)

A manufacturer may only distribute medical cannabis as a pill, liquid, or as a topical formulation. Manufacturers are allowed, but not required, to distribute medical cannabis products, such as delivery devices and educational material.

All medical cannabis must be assigned a tracking number and be packaged in compliance with the United States Poison Prevention Packaging Act.\(^{41}\) All medical cannabis must also be labeled with the following information:

- All active ingredients
- Individually identifying information, including:
  - the patient’s name and date of birth
  - if applicable, the name and date of birth of the patient’s registered designated caregiver or parent or legal guardian
  - the patient’s registry identification number
  - the chemical composition
  - the dosage

**People allowed to receive medical cannabis**\(^{42}\)

A manufacturer may distribute medical cannabis only to a person listed on the patient’s registry verification that the manufacturer received from MDH. The manufacturer may not distribute any medical cannabis until the registry verification has been received. The registry verification

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\(^{39}\) Minn. Stat. § 152.29, subd. 1, para. (a).

\(^{40}\) Minn. Stat. §§ 152.22, subd. 6; 152.29, subd. 3.

\(^{41}\) 15 U.S.C. §§ 1471-1477. (The United States Poison Prevention Packaging Act, P.L. 97-601, was enacted to protect children from unintended ingestion of medicines and common household products.)

\(^{42}\) Minn. Stat. § 152.29, subd. 3.
includes patient information and may also include a registered designated caregiver or a parent or
guardian of the patient. If a person is listed on the registry verification, the manufacturer may
distribute the medical cannabis after verifying the person’s identity by photographic
identification, unless the individual distributing the medical cannabis personally knows the
recipient. 43

Who may distribute the medical cannabis 44

Only employees of the manufacturer who are licensed pharmacists in Minnesota may give final
approval for distribution of medical cannabis. Distribution may only occur after a pharmacist has
consulted with the patient to determine the proper dosage and range of chemical compositions
for that individual patient. The pharmacist may consult with the patient via videoconference, as
long as the consultation meets certain requirements.

Amount of medical cannabis that can be distributed 45

A maximum of a 30-day supply of the dosage determined for the individual patient may be
distributed at one time.

Other

Relationship with health care practitioners 46

A manufacturer must not share office space with a health care practitioner. A manufacturer is
also prohibited from referring patients to a health care practitioner or having any financial
relationship with a health care practitioner.

Marketing restrictions 47

Manufacturers must comply with reasonable restrictions set by the Commissioner of Health
relating to signage, marketing, display, and advertising of medical cannabis.

Transportation 48

A manufacturer may staff a motor vehicle with one employee to transport medical cannabis to a
certified laboratory to be tested or to a facility for disposal. A manufacturer must staff a motor

43 Minn. Stat. § 152.11, subd. 2d.
44 Minn. Stat. § 152.29, subd. 3, para. (a).
45 Minn. Stat. § 152.29, subd. 3, para. (c), cl. (6).
46 Minn. Stat. § 152.29, subd. 1, para. (c).
47 Minn. Stat. § 152.29, subd. 1, para. (k).
48 Minn. Stat. § 152.29, subd. 3a.
vehicle with at least two employees when transporting medical cannabis for any other purpose or to any other destination.

Criminal and civil liability

The law establishes criminal penalties that apply to manufacturers or employees of manufacturers in addition to any other applicable penalty in law. Any manufacturer or agent of a manufacturer who intentionally transfers medical cannabis to a person other than one listed on a registry verification or submits false records or documentation required by MDH to register as a manufacturer is guilty of a felony punishable by up to two years of imprisonment, a fine of not more than $3,000, or both. If certain individuals affiliated with a manufacturer intentionally divert medical cannabis outside the state to a person other than allowed by law, the commissioner may fine the manufacturer $250,000 and may begin proceedings to revoke the manufacturer’s registration. A manufacturer may also be fined up to $1,000, in addition to any other applicable penalty in law, for any violation of laws or regulations relating to the registry program where no penalty is specified.

Criminal protections

Employees of a manufacturer or an independent laboratory that tests medical cannabis are exempted from criminal liability under Minnesota law for the possession, dosage determination, and sale of medical cannabis as permitted under the registry program.

Data practices

Data submitted to a medical cannabis manufacturer, and data submitted by a medical cannabis manufacturer to the Commissioner of Health, are classified as private data on individuals or nonpublic data. This data may be used to comply with requirements in chapter 13 and to comply with requests from the legislative auditor or state auditor.

Health Care Practitioners

A health care practitioner, for purposes of the registry program, is defined as a Minnesota-licensed doctor of medicine, a Minnesota-licensed physician assistant acting within the scope of practice, or a Minnesota-licensed advanced practice registered nurse, with the primary responsibility for care and treatment of the patient’s underlying qualifying medical condition.

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49 Minn. Stat. § 152.33, subds. 1, 4, and 6.
50 Minn. Stat. § 152.32, subd. 2.
51 Minn. Stat. § 152.31.
52 Minn. Stat. § 152.22, subd. 4.


**Participation**

**MDH training/notification**

The Commissioner of Health must notify all eligible health care practitioners in the state about the registry program. This notice must include an explanation of the purposes and requirements of the program. If a health care practitioner meets the requirements and requests to participate in the program, the commissioner must allow that participation. However, no health care practitioner is required to participate in the program. In addition to notification, the commissioner also must provide practitioners with explanatory information and assistance in understanding the therapeutic uses of medical cannabis under the program requirements. The commissioner must provide patient applications to participating health care practitioners, who then provide those applications to patients.

**Certifications**

In order for a patient to participate in the registry program, a health care practitioner must provide a certification of diagnosis with at least one qualifying medical condition. The patient’s application must include this certification in order to participate in the registry program, and the certification must have been given by the practitioner within the 90 days prior to the patient’s application. Practitioners must use the certification form developed by the Commissioner of Health.

In certain circumstances, the practitioner may also provide a certification of a patient’s disability as part of the patient’s certification of diagnosis. The law allows for patients in the registry program to have a registered designated caregiver if the patient is either unable to self-administer medication or is unable to acquire medical cannabis from a distribution facility due to a developmental or physical disability.

**Responsibilities during participation**

The law requires that if a health care practitioner agrees to participate in the registry program, the practitioner must continue to treat the patient for the patient’s qualifying medical condition. Throughout that ongoing treatment, the practitioner must report the health records of the patient to the commissioner. The reporting of health records must be made in a manner set by the commissioner and is subject to data privacy provisions. Each year, the practitioner also must determine if the patient continues to suffer from a qualifying medical condition and, if so, issue a new certification of that diagnosis.

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53 Minn. Stat. § 152.27, subd. 2, para. (a).
54 Minn. Stat. § 152.28, subd. 1, para. (c).
55 Minn. Stat. § 152.28, subd. 1, para. (a), cls. (1) and (2).
56 Minn. Stat. § 152.28, subd. 1, paras. (a), cl. (5), and (b).
Advice to patients and others\textsuperscript{37}

A health care practitioner working with a patient in the program must provide the patient, registered designated caregiver, and parent or legal guardian with information on nonprofit patient support groups or organizations. The practitioner is also required to provide explanatory information provided by MDH disclosing:

- the experimental nature of therapeutic use of medical cannabis;
- the possible risks, benefits, and side effects of the proposed treatment;
- the application for participation in the program;
- other materials from the commissioner; and
- the Tennessee warning.\textsuperscript{38}

Medical Assistance/MinnesotaCare reimbursement\textsuperscript{39}

Medical Assistance (MA) and MinnesotaCare are not required to reimburse an enrollee or a provider for “costs associated with the medical use of cannabis.” Medical cannabis is not listed on the drug formulary for MA and MinnesotaCare, so those programs do not cover medical cannabis.\textsuperscript{60} MA and MinnesotaCare are, however, still required to reimburse for services related to the treatment of the patient’s qualifying medical condition if that service is covered under applicable statutes.

Legal Issues

Health records\textsuperscript{61}

All data collected on patients and reported to the patient registry are health records under the Health Records Act and are classified as private data on individuals. The data may, however, be used or reported in an aggregated, nonidentifiable form as part of the scientific, peer-reviewed publication of research required under the law or in the creation of summary data.

\textsuperscript{37} Minn. Stat. § 158.28, subd. 1, para. (a), cls. (3) and (4).

\textsuperscript{38} See Minn. Stat. § 15.04, subd. 2 (explaining the Tennessee warning).

\textsuperscript{39} Minn. Stat. § 152.23, para. (b).

\textsuperscript{60} See Minn. Stat. § 256B.0625, subd. 13d; Minnesota Health Care Programs Provider Manual, http://www.dhs.state.mn.us.

\textsuperscript{61} Minn. Stat. §§ 152.28, subd. 2; 152.31.
Civil/disciplinary protections^62^  

The law prohibits the Board of Medical Practice, the Board of Nursing, or any other professional licensing board from subjecting a health care practitioner to any civil or disciplinary penalties solely for participation in the registry program. This protection also extends to pharmacists under the Board of Pharmacy. The protection does not prevent a professional licensing board from taking action in response to violations of any other section of law. The law also does not provide any civil protections for health care practitioners for claims of malpractice, negligence, or any other civil claim.

Criminal liability^63^  

Although the law creates exemptions from criminal liability for certain actions by patients, caregivers, and manufacturers, it does not exempt health care practitioners from criminal liability. Under the registry program, a health care practitioner does not possess or distribute medical cannabis and is therefore not exempted from criminal controlled substance possession laws.

A health care practitioner is subject to a misdemeanor penalty, punishable by up to 90 days in jail or payment of a fine up to $1,000, or both, for the following actions:

- knowingly providing patients with referrals to a specific manufacturer or a specific designated caregiver
- advertising as a manufacturer
- issuing a certification that a patient has a qualifying medical condition while holding a financial interest in a manufacturer

A case decided by the federal Court of Appeals for the Ninth Circuit addressed whether a health care practitioner may be criminally liable for aiding and abetting a federal crime for the physician’s “recommendation” to a patient to use marijuana for medicinal purposes. In Conant v. Walters, the court held that a doctor’s “recommendation” alone did not amount to aiding and abetting.\(^64\) The case was based on California law that required a doctor to “recommend” a patient’s use of medical marijuana. Minnesota law differs from California law in that respect, as a practitioner in Minnesota is providing a “certification of diagnosis” and not a “recommendation.” It is also important to note that the Ninth Circuit Court of Appeals does not have jurisdiction over Minnesota and therefore this decision is not binding on Minnesota courts.

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\(^{62}\) Minn. Stat. § 152.32, subd. 2, para. (c).

\(^{63}\) Minn. Stat. § 152.33, subd. 5.

\(^{64}\) Conant v. Walters, 309 F.3d 629 (9th Cir. 2002).
Advertising restrictions

Health care practitioners are prohibited from publishing advertisements that:

- contain false or misleading statements about medical cannabis or the registry program;
- use colloquial terms to refer to medical cannabis;
- state or imply that a health care practitioner is endorsed by MDH;
- include images of cannabis or of cannabis-smoking paraphernalia; or
- contain medical symbols that may be confused with symbols of medical associations.

A health care practitioner who violates these advertising restrictions cannot certify patients to participate in the registry.

Other

Prescription Monitoring Program

Medical cannabis is not eligible to be entered into the Prescription Monitoring Program (PMP). Under Minnesota and federal law, cannabis is a Schedule I controlled substance, and therefore medical cannabis is not dispensed under a prescription drug order, as required by statute to be entered in the PMP.

Discrimination for purposes of medical care prohibited

Discrimination against patients for the purpose of medical care is prohibited. The law states that a patient's use of medical cannabis is considered the equivalent to the authorized use of any other medication and does not constitute the use of an illicit substance or otherwise disqualify a patient from needed medical care, including organ transplants.

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65 Minn. Stat. § 152.28, subd. 3
66 Minn. Stat. § 152.126.
67 The Prescription Monitoring Program (PMP) is codified in Minnesota Statutes, section 152.126. The PMP allows health care practitioners with prescribing authority to check the database for a patient's history of controlled substance prescriptions. The information in the PMP is generally inputted by the pharmacist who delivers the controlled substance. Among the included substances in the PMP are all substances classified as a Schedule II through V.
68 Minn. Stat. § 152.32, subd. 3, para. (b).
Health care facilities and home care providers

Under the law, health care facilities and home care providers may adopt reasonable restrictions on the use of medical cannabis by a patient who resides at or is actively seeking care or treatment at the facility or from the provider. For purposes of this provision, health care facilities include those licensed under chapter 144A (nursing homes and hospice facilities), boarding care homes licensed under section 144.50, assisted living facilities, and facilities owned, controlled, managed, or under common control with hospitals licensed under chapter 144. Restrictions may include that the facility or provider will not store or maintain the patient's medical cannabis supply, that the facility or provider is not responsible for providing the medical cannabis for patients, and that medical cannabis may only be used in specified places within the facility. The facilities and providers are not required to adopt any restrictions and are prohibited from unreasonably limiting a patient's access to or use of medical cannabis.

Employees of a health care facility, emergency medical services personnel, and home care providers are not subject to a violation under this chapter for possessing medical cannabis during the course of their duties and may distribute medical cannabis to a registered patient who resides at or is seeking active care and treatment at the facility or from the provider. Under this section, employees acting within the course of their duties are not required to register as a designated caregiver.

Operation of the Program

Appropriations

MDH receives appropriations from the general fund and the state government special revenue fund to fund the activities of its Office of Medical Cannabis. Appropriations from the state government special revenue fund are from annual registration fees collected from manufacturers and annual fees collected from patients for enrollment in the patient registry.

### Appropriations to MDH for the Office of Medical Cannabis

<table>
<thead>
<tr>
<th>Fiscal Year</th>
<th>General Fund</th>
<th>State Government Special Revenue Fund</th>
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<tr>
<td>FY 2015</td>
<td>$2,795,000</td>
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<tr>
<td>FY 2016</td>
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<tr>
<td>FY 2019</td>
<td>$708,000</td>
<td>$729,000</td>
</tr>
</tbody>
</table>

69 Minn. Stat. §§ 144A.4791, subd. 14; 152.34.
**Task Force on Medical Cannabis Therapeutic Research**70

The Task Force on Medical Cannabis Therapeutic Research was established to conduct an impact assessment of the registry program on Minnesota. Initially the task force was also involved in certain deadline extensions for the program. The 23-member task force consists of representatives from:

- the House of Representatives and the Senate;
- consumers or patients enrolled in the registry program;
- health care providers;
- law enforcement and prosecutors;
- substance use disorder treatment providers; and
- the commissioners of health, human services, and public safety.

All members, except the members from the House of Representatives and the Senate, are appointed by the governor. Two members of the House of Representatives and two members of the Senate are also appointed, with one member of each body serving as a co-chair. The co-chairs are appointed by the Senate majority leader and the Speaker of the House. The second member from each body is appointed by the minority leader of that body. All members serve at the pleasure of their appointing authority. The Commissioner of Health provides administrative and technical support to the task force.

**Deadline extensions**71

The task force could have extended the deadline to register manufacturers and the distribution deadline by six months if requested by the Commissioner of Health. MDH did register two manufacturers by the December 1, 2014, deadline, and the manufacturers began distributing medical cannabis on the July 1, 2015, deadline, so no extensions were needed.

**Cost assessments**

Beginning with a report on January 15, 2015, and continuing annually until January 15, 2019, the commissioners of the state executive agencies impacted by the medical cannabis therapeutic research study must report to the co-chairs of the task force the costs incurred by each agency in implementing the study. Agencies are required to report actual costs incurred compared to estimated costs.

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70 Minn. Stat. § 152.36.
71 Minn. Stat. § 152.25, subd. 3.
Impact assessments

The task force must complete an impact assessment and report it to the legislature every two years beginning in 2017. The impact assessment must be conducted by holding hearings to evaluate the impact of medical cannabis use and evaluate Minnesota’s activities involving medical cannabis. The impact assessment must include analysis of:

- the program design and implementation;
- the impact on the health care provider community;
- patient experiences;
- the impact on the incidence of substance abuse;
- access to and quality of medical cannabis and medical cannabis products;
- the impact on law enforcement and prosecutions;
- public awareness and perception; and
- any unintended consequences.

The task force issued its first impact assessment report February 1, 2017. In that report, the task force provided an overview of Minnesota’s medical cannabis program’s design and implementation; participation in the program by patients, health care practitioners, and registered designated caregivers; and rules adopted by the department. The report also described the department’s periodic surveys of patients and health care practitioners regarding benefits and negative effects of medical cannabis use and included survey results for the July to September 2015 time period. No substance abuse impacts related to the medical cannabis program or serious adverse incidents related to overdoses or diversions have been reported to the department. Additionally, the report summarized testing requirements for medical cannabis and factors that affect access to medical cannabis. Finally, the report provided information about patient perceptions of the extent to which medical cannabis products for sale in Minnesota meet patient needs.

Additional reports to the legislature

The task force had to report to the legislature by February 1, 2015, on the design and implementation of the registry program. In addition, it must make reports based on the biennial cost assessments from the state agencies.

At any time, the task force may recommend to the legislature whether to add or remove conditions from the list of qualifying medical conditions.

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72 This report may be found on the Minnesota Department of Health’s website at http://www.health.state.mn.us/topics/cannabis/taskforce/reportfinal061417.pdf
Legislative History of Medical Cannabis in Minnesota

In 1980, the THC Therapeutic Research Act was adopted and signed into law. The purpose of the act was to research whether cannabis could alleviate the effects of chemotherapy during the treatment of cancer. The act required the Commissioner of the Department of Health to appoint a principal investigator. The principal investigator was required to obtain cannabis only from the National Institute on Drug Abuse and comply with federal laws and regulations while conducting the research program. In 1980, $100,000 was appropriated by the legislature to the Commissioner of Health to administer the act, but the appropriation was vetoed by Governor Al Quie.

In 2001, Representative Phyllis Kahn introduced House File 2164, known as the Compassionate Use Act. That act would have allowed for the medical use of cannabis after a patient had been diagnosed by a physician as having a debilitating medical condition. The House bill, and its companion bill in the Senate, were both introduced but not heard in committee.

In 2007, Representative Thomas Huntley introduced House File 655 and Senator Steve Murphy introduced Senate File 345. Both bills would have allowed the use of medical cannabis for treatment of a debilitating medical condition. The Senate file passed the Senate floor and was referred to the House where it was given a second reading, but not passed.

In 2009, the first medical cannabis law that would have allowed patient possession of medical cannabis passed both bodies of the legislature. The act allowed patients to possess and use cannabis if diagnosed with a terminal illness that was accompanied by a variety of symptoms. The act passed both the House and the Senate and was vetoed by Governor Tim Pawlenty on May 22, 2009.

In 2013, Representative Carly Melin and Senator Scott Dibble introduced House File 1818 and Senate File 1641, respectively, both allowing for the use and possession of medical cannabis by patients with a specified list of conditions. House File 1818 was referred to committee but did not pass the House floor. Senate File 1641 passed the Senate on May 6, 2014, and was referred to the House for consideration, but was not heard in committee.

On April 24, 2014, Senate File 2470, originally a bill relating to education, passed the Senate and was referred to the House for consideration. The bill was heard in the Rules and Administration Committee where an amendment was offered and adopted that allowed for the medical use of cannabis through a clinical trial model. The bill was then heard in the Ways and Means Committee where another amendment was offered and adopted, altering the program to a registry program.

73 Minn. Stat. § 152.21, subd. 1.
74 Minn. Stat. § 152.21, subd. 4.
75 Minn. Stat. § 152.21, subd. 5.
77 Laws 2009, ch. 166; Senate File 97, House File 292.
The bill was sent to the House floor where it was passed with additional amendments. Because the bill originated in the Senate and already passed the Senate, the Senate was able to either concur on the bill as amended or refuse to concur. The Senate refused to concur and the bill was heard in conference committee and passed by both bodies as amended in conference committee. Governor Mark Dayton signed the bill into law on May 29, 2014.\textsuperscript{78}

Laws 2015, chapter 74, amended various sections of the medical cannabis act by:

- modifying the definition of medical cannabis to include possession by a manufacturer or laboratory of any part of the cannabis plant prior to processing the plant into an approved liquid or pill form;
- establishing time limits for the Commissioner of Health to either approve or deny a patient's application for the registry program; and
- adding facilities owned, controlled, managed, or under common control of a hospital to those facilities that may adopt reasonable restrictions on the use of medical cannabis by patients who reside at or are actively receiving care or treatment at the facility.

A provision was also added to allow employees of a health care facility, in the course of their duties, to possess medical cannabis for a registered patient without registering with the commissioner as a designated caregiver.

Laws 2016, chapter 179, amended various sections of the medical cannabis act by:

- expanding the definition of qualifying medical condition to include inflammatory bowel disease;
- requiring the Commissioner of Health to regularly update legislators about certain topics;
- specifying that only manufacturer employees licensed as pharmacists may give final approval for distribution of medical cannabis;
- allowing patient consultations via videoconference to determine dosages;
- allowing the transportation of medical cannabis by only one manufacturer employee in certain circumstances; and
- directing the Commissioner of Health to provide administrative and technical support to the Task Force on Medical Cannabis Therapeutic Research.

A separate provision was added to statutes governing home care providers, allowing home care providers to adopt reasonable restrictions on the use of medical cannabis by patients in the registry program who receive care from home care providers, and to protect home care provider employees from being subject to violations of controlled substance laws for carrying out employment duties and caring for patients in the registry program.

\textsuperscript{78} Laws 2014, ch. 311.
Laws 2017, first special session, chapter 6, modified the Commissioner of Health's authority and tools for regulating registered medical cannabis manufacturers and health care practitioners. Regarding manufacturers, the commissioner is authorized to:

- accept additional registration applications from manufacturers if a manufacturer registered before December 1, 2014, ceases to be registered;
- not renew a manufacturer's registration, if an officer, director, or controlling person of a manufacturer diverts medical cannabis to a person other than allowed by law; and
- impose a civil penalty and revoke a manufacturer's registration, if an officer, director, or controlling person of a manufacturer diverts medical cannabis to a person other than allowed by law and transports medical cannabis outside the state.

The law also established procedures for the commissioner to follow to revoke, not renew, deny consent to transfer, or temporarily suspend a registration of a medical cannabis manufacturer. In addition, the law prohibited health care practitioners from including certain terms, images, and symbols in their advertising. A health care practitioner who violates these advertising restrictions cannot certify patients as having qualifying medical conditions, for purposes of participating in the medical cannabis registry.

For more information about health issues, visit the health and human services area of our website, www.house.mn/hrd/.
152.22 DEFINITIONS.
   Subdivision 1. Applicability. For purposes of sections 152.22 to 152.37, the terms
   defined in this section have the meanings given them.

   Subd. 2. Commissioner. "Commissioner" means the commissioner of health.

   Subd. 3. Disqualifying felony offense. "Disqualifying felony offense" means a
   violation of a state or federal controlled substance law that is a felony under Minnesota
   law, or would be a felony if committed in Minnesota, regardless of the sentence imposed,
   unless the commissioner determines that the person's conviction was for the medical use
   of cannabis or assisting with the medical use of cannabis.

   Subd. 4. Health care practitioner. "Health care practitioner" means a Minnesota
   licensed doctor of medicine, a Minnesota licensed physician assistant acting within the
   scope of authorized practice, or a Minnesota licensed advanced practice registered nurse
   who has the primary responsibility for the care and treatment of the qualifying medical
   condition of a person diagnosed with a qualifying medical condition.

   Subd. 5. Health records. "Health records" means health records as defined in
   section 144.291, subdivision 2, paragraph (c).

   Subd. 6. Medical cannabis. (a) "Medical cannabis" means any species of the genus
   cannabis plant, or any mixture or preparation of them, including whole plant extracts and
   resins, and is delivered in the form of:
      (1) liquid, including, but not limited to, oil;
      (2) pill;
      (3) vaporized delivery method with use of liquid or oil but which does not require the
          use of dried leaves or plant form; or
      (4) any other method, excluding smoking, approved by the commissioner.

      (b) This definition includes any part of the genus cannabis plant prior to being
          processed into a form allowed under paragraph (a), that is possessed by a person while
          that person is engaged in employment duties necessary to carry out a requirement under
          sections 152.22 to 152.37 for a registered manufacturer or a laboratory under contract
          with a registered manufacturer.

   Subd. 7. Medical cannabis manufacturer. "Medical cannabis manufacturer" or
   "manufacturer" means an entity registered by the commissioner to cultivate, acquire, manufacture,
   possess, prepare, transfer, transport, supply, or dispense medical cannabis, delivery devices, or
   related supplies and educational materials.

   Subd. 8. Medical cannabis product. "Medical cannabis product" means any
   delivery device or related supplies and educational materials used in the administration of
   medical cannabis for a patient with a qualifying medical condition enrolled in the registry
   program.

   Subd. 9. Patient. "Patient" means a Minnesota resident who has been diagnosed
   with a qualifying medical condition by a health care practitioner and who has otherwise
   met any other requirements for patients under sections 152.22 to 152.37 to participate in
   the registry program under sections 152.22 to 152.37.

   Subd. 10. Patient registry number. "Patient registry number" means a unique
   identification number assigned by the commissioner to a patient enrolled in the registry
   program.

   Subd. 11. Registered designated caregiver. "Registered designated caregiver"
   means a person who:
      (1) is at least 21 years old;
      (2) does not have a conviction for a disqualifying felony offense;

   (3) has been approved by the commissioner to assist a patient who has been identified
       by a health care practitioner as developmentally or physically disabled and therefore
       unable to self-administer medication or acquire medical cannabis from a distribution
       facility due to the disability; and
(4) is authorized by the commissioner to assist the patient with the use of medical cannabis.

Subd. 12. Registry program. "Registry program" means the patient registry established in sections 152.22 to 152.37.

Subd. 13. Registry verification. "Registry verification" means the verification provided by the commissioner that a patient is enrolled in the registry program and that includes the patient's name, registry number, and qualifying medical condition and, if applicable, the name of the patient's registered designated caregiver or parent or legal guardian.

Subd. 14. Qualifying medical condition. "Qualifying medical condition" means a diagnosis of any of the following conditions:

(1) cancer, if the underlying condition or treatment produces one or more of the following:

(i) severe or chronic pain;

(ii) nausea or severe vomiting; or

(iii) cachexia or severe wasting;

(2) glaucoma;

(3) human immunodeficiency virus or acquired immune deficiency syndrome;

(4) Tourette's syndrome;

(5) amyotrophic lateral sclerosis;

(6) seizures, including those characteristic of epilepsy;

(7) severe and persistent muscle spasms, including those characteristic of multiple sclerosis;

(8) inflammatory bowel disease, including Crohn's disease;

(9) terminal illness, with a probable life expectancy of under one year, if the illness or its treatment produces one or more of the following:

(i) severe or chronic pain;

(ii) nausea or severe vomiting; or

(iii) cachexia or severe wasting; or

(10) any other medical condition or its treatment approved by the commissioner.

History: 2014 c 311 s 2; 2015 c 74 s 2; 2016 c 179 s 27

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