PHYSICIAN’S GUIDE TO MAINE LAW

As a service to members of the Maine Medical Association, this book is intended to alert Maine physicians to laws affecting aspects of their practice. It represents an attempt to highlight those laws and rules of particular interest to Maine physicians from Maine statutory law, the ever-increasing regulations promulgated by various state agencies, applicable federal law, and the Current Opinions of the American Medical Association’s Council on Ethical and Judicial Affairs. In some instances, excerpts from the statute or regulation have been quoted. In other cases, the relevant material has been paraphrased or electronic links to sources or more information have been provided.

State laws passed through May 2018 have been included and continuing updates will be made; however, since amendments to state law occur frequently, and excerpts can be misinterpreted when viewed in isolation, the full text of a relevant statute or rule should be consulted. Members are advised to consult with an MMA attorney or attorney knowledgeable in health care law for answers to specific questions.

The Association hopes that this booklet will be a ready and valuable reference for your practice. Robert Schlager, MD
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Notes of Importance:

- The information provided in this resource manual is subject to change without notice. Statutes and regulations change on a frequent basis. Every effort will be made to keep this Guide current, but if you need to know absolutely about a section’s current status, please contact the Maine Medical Association for more information.

- The information provided is for reference purposes only and not intended to provide legal advice. If you need legal advice, please contact the MMA or a licensed attorney.

- Websites and other web-links are provided as a courtesy to our members. The links are subject to change or cancellation without notice. MMA does not take a particular position on any of these websites or their content and does not have a financial incentive to support any of the listed sites. They are provided for reference purposes only.

- The abbreviation MRSA in citations refers to the Maine Revised Statues. The number appearing before MRSA is the title of the statute; the number after MSRA refers to the specific section. You can find the full text of the statutes at: http://www.mainelegislature.org/legis/statutes. The abbreviation USC refers to federal statues, or the United States Code, found at: http://uscode.house.gov. CFR refers to the Code of Federal Regulations, found at: http://www.ecfr.gov

- In 2018 the Legislature passed P.L. 2017 c. 407, An Act to Implement the Recommendations of the Task Force to Address the Opioid Crisis in the State Regarding Respectful Language. This law calls for the replacement of stigmatizing language in many parts of Maine statutes with more respectful language; for example, the terms “alcohol and drug abuse” are replaced by “substance use.” This Guide has not yet incorporated those changes.
(1) **List of Maine State Agency Rules Links**

All of the below listed Agency rules are adopted from the State of Maine website under [Maine State Agency Rules](#). These rules and links are subject to change without notice. The select links below are for the benefit of the users of this manual who may need to refer to a particular agency rule for further information or guidance. Please note that selecting a specific chapter will download a Word document from the agency websites.

**a. Rule Chapters for the Department of Health and Human Services**

- [10-144](#) Department of Health and Human Services - General
- [10-146](#) Office of Data Research and Vital Statistics
- [10-148](#) Community Services Programs *(formerly Office of Child and Family Services; see also 14-472)*
- [10-149](#) Office of Aging and Disability Services, part 1
- [14-118](#) Community Services Programs / Office of Substance Abuse
- [14-191](#) Mental Health and Mental Retardation - General
- [14-193](#) Office of Adult Mental Health
- [14-197](#) Office of Aging and Disability Services, part 2
- [14-472](#) Office of Child and Family Services *(the part that was formerly Bureau of Children with Special Needs; see also 10-148)*

**b. Rule Chapters for the Department of Professional & Financial Regulation**

- [02-031](#) Bureau of Insurance
- [02-297](#) Board of Chiropractic Licensure
- [02-313](#) Board of Dental Examiners
- [02-343](#) Board of Respiratory Care Practitioners
- [02-373](#) Board of Licensure in Medicine
- [02-380](#) State Board of Nursing
- [02-382](#) State Board of Optometry
- [02-383](#) Board of Osteopathic Licensure
- [02-392](#) Maine Board of Pharmacy
- [02-393](#) Board of Examiners in Physical Therapy
- [02-396](#) Board of Licensure of Podiatric Medicine
- [02-415](#) State Board of Examiners of Psychologists
- [02-465](#) Radiologic Technology Board of Examiners
- [02-502](#) Board of Complementary Health Care Providers *(formerly Acupuncture Licensing Board)*
- [02-643](#) Board of Speech, Audiology and Hearing

**c. Independent Agencies**

- [90-351](#) Workers' Compensation Board
- [90-590](#) Maine Health Data Organization
- [94-457](#) Finance Authority of Maine (FAME)
- [94-630](#) Maine Biomedical Research Board
d. **Department of Public Safety**

16-163 Office of Emergency Medical Services
16-230 Maine Drug Enforcement Agency

**e. Department of Environmental Protection**

Ch. 900 Biomedical Waste Management Rules

f. **Department of the Secretary of State, Bureau of Motor Vehicles**

Ch. 3 Physical, Emotional and Mental Competence to Operate a Motor Vehicle

(2) **Abandonment**

See AMA Opinions (AMA Code of Medical Ethics and Policy Finder)


**Disciplinary Sanctions**

The board (of licensure in Medicine) may suspend or revoke a license pursuant to Title 5, section 10004. The following are grounds for an action to refuse to issue, modify, restrict, suspend, revoke or refuse to renew the license of an individual licensed under this chapter:

32 MRSA §3282-A (2)

**Fraud, deceit or misrepresentation**

The practice of fraud, deceit or misrepresentation in obtaining a license under this chapter or in connection with service rendered within the scope of the license issued.

32 MRSA §3282-A (2)(A)

**Misuse of alcohol, drugs or other substances**

Misuse of alcohol, drugs or other substances that has resulted or may result in the licensee performing services in a manner that endangers the health or safety of patients.

32 MRSA §3282-A (2)(B)

**Diagnosis of mental or physical condition**

A professional diagnosis of a mental or physical condition that has resulted or may result in the licensee performing services in a manner that endangers the health or safety of patients.

32 MRSA §3282-A (2)(C)

**Aiding in the unlicensed practice of medicine**

Aiding or abetting the practice of medicine by an individual who is not licensed under this chapter and who claims to be legally licensed.

32 MRSA §3282-A (2)(D)

**Incompetence**

A licensee is considered incompetent in the practice if the licensee has:

A. Engaged in conduct that evidences a lack of ability or fitness to discharge the duty owed by the licensee to a client or patient or the general public; or

B. Engaged in conduct that evidences a lack of knowledge or inability to apply principles or skills to carry out the practice for which the licensee is licensed;

32 MRSA §3282-A (2)(E)

**Unprofessional conduct**

A licensee is considered to have engaged in unprofessional conduct if the licensee violates a standard of professional behavior that has been established in the practice for which the licensee is licensed.

32 MRSA §3282-A (2)(F)
Conviction of certain crimes
Conviction of a crime that involves dishonesty or false statement or relates directly to the practice for which the licensee is licensed, or conviction of a crime for which incarceration for one year or more may be imposed.  

32 MRSA §3282-A (2)(G)

Rule violation
A violation of this chapter or a rule adopted by the board.  

32 MRSA §3282-A (2)(H)

False advertising
Engaging in false, misleading or deceptive advertising.  

32 MRSA §3282-A (2)(I)

Prescribing of controlled substances for unauthorized purposes
Prescribing narcotic or hypnotic or other drugs listed as controlled substances by the Drug Enforcement Administration for other than accepted therapeutic purposes.  

32 MRSA §3282-A (2)(J)

Failure to report impairment of physician
Failure to report to the secretary of the board a physician licensed under this chapter for addiction to alcohol or drugs or for mental illness in accordance with Title 24, section 2505 (obligation to report unsafe conduct), except when the impaired physician is or has been a patient of the licensee.  

32 MRSA §3282-A (2)(K)

Failure to provide informed consent for breast cancer treatment
Failure to comply with the requirements of Title 24, section 2905-A (obligation to provide information, orally and in writing, regarding breast cancer treatment options).  

32 MRSA §3282-A (2)(L)

Disciplinary actions by another state
Revocation, suspension or restriction of a license to practice medicine or other disciplinary action; denial of an application for a license; or surrender of a license to practice medicine following the institution of disciplinary action by another state or a territory of the United States or a foreign country if the conduct resulting in the disciplinary or other action involving the license would, if committed in this State, constitute grounds for discipline under the laws or rules of this State.  

32 MRSA §3282-A (2)(M)

Practicing beyond scope of license
Engaging in any activity requiring a license under the governing law of the board that is beyond the scope of acts authorized by the license held.  

32 MRSA §3282-A (2)(N)

Practicing after license suspension
Continuing to act in a capacity requiring a license under the governing law of the board after expiration, suspension or revocation of that license.  

32 MRSA §3282-A (2)(O)

Noncompliance
Noncompliance with an order or consent agreement of the board.  

32 MRSA §3282-A (2)(P)

Failure to produce documents
Failure to produce upon request of the board any documents in the licensee's possession or under the licensee's control concerning a pending complaint or proceeding or any matter under investigation by the board, unless otherwise prohibited by state or federal law.  

32 MRSA §3282-A (2)(Q)

Failure to respond to complaint
Failure to timely respond to a complaint notification sent by the board.  

32 MRSA §3282-A (2)(R)
Failure to consult prescription monitoring information
Failure to comply with the requirements of Title 22, section 7253 (prescribers and dispensers required to check prescription monitoring information.

32 MRSA §3282-A (2)(S)

(3) Abortion and Miscarriage
General
Policy
It is the public policy of the State that the State not restrict a woman's exercise of her private decision to terminate a pregnancy before viability except as provided in section 1597-A (consent to a minor’s decision to have and abortion). After viability an abortion may be performed only when it is necessary to preserve the life or health of the mother. It is also the public policy of the State that all abortions may be performed only by a physician.

22 MRSA §1598 (1)

Definitions
A. Abortion: means the intentional interruption of a pregnancy by the application of external agents, whether chemical or physical or by the ingestion of chemical agents with an intention other than to produce a live birth or to remove a dead fetus.
B. Viability: means the state of fetal development when the life of the fetus may be continued indefinitely outside the womb by natural or artificial life-supportive systems.

22 MRSA §1598 (2)

Persons who may perform abortions; penalties
A. Only a person licensed under Title 32, chapter 36 (doctor of osteopathic medicine) or chapter 48 (doctor of medicine), to practice medicine in Maine as a medical or osteopathic physician, may perform an abortion on another person.
B. Any person not so licensed who knowingly performs an abortion on another person or any person who knowingly assists a nonlicensed person to perform an abortion on another person is guilty of a Class C crime.

22 MRSA §1598 (3)

Abortions after viability; criminal liability
A person who performs an abortion after viability is guilty of a Class D crime if:
A. He knowingly disregarded the viability of the fetus; and
B. He knew that the abortion was not necessary for the preservation of the life or health of the mother.

22 MRSA §1598 (4)

Immunity and employment protection
No physician, nurse or other person who refuses to perform or assist in the performance of an abortion, and no hospital or health care facility that refuses to permit the performance of an abortion upon its premises, shall be liable to any person, firm, association or corporation for damages allegedly arising from the refusal, nor shall such refusal constitute a basis for any civil liability to any physician, nurse or other person, hospital or health care facility nor a basis for any disciplinary or other recriminatory action against them or any of them by the State or any person.

No physician, nurse or other person, who refuses to perform or assist in the performance of an abortion, shall, because of that refusal, be dismissed, suspended, demoted or otherwise prejudiced or damaged by a hospital, health care facility, firm, association, professional association, corporation or educational institution with which he or she is affiliated or requests to be affiliated or by which he or she is employed, nor shall such refusal constitute grounds for loss of any privileges or immunities to which such physician, nurse or other person would otherwise be entitled nor shall submission to an abortion or the granting of consent therefore be a condition precedent to the receipt of any public benefits.
 Discrimination for refusal
No person, hospital, health care facility, firm, association, corporation or educational institution, directly or indirectly, by himself or another, shall discriminate against any physician, nurse or other person by refusing or withholding employment from or denying admittance, when such physician, nurse or other person refuses to perform, or assist in the performance of an abortion, nor shall such refusal constitute grounds for loss of any privileges or immunities to which such physician, nurse or other person would otherwise be entitled.

Informed consent to abortion
A. **Consent by the woman.** A physician may not perform an abortion unless, prior to the performance, the attending physician certifies in writing that the woman gave her informed written consent, freely and without coercion.
B. **Informed consent.** To ensure that the consent for an abortion is truly informed consent, the attending physician shall inform the woman, in a manner that in the physician's professional judgment is not misleading and that will be understood by the patient, of at least the following:
   1. According to the physician's best judgment she is pregnant;
   2. The number of weeks elapsed from the probable time of the conception;
   3. The particular risks associated with her own pregnancy and the abortion technique to be performed; and
   4. At the woman’s request, alternatives to abortion such as childbirth and adoption and information concerning public and private agencies that will provide the woman with economic and other assistance to carry the fetus to term, including, if the woman so requests, a list of these agencies and the services available from each.

Failure to preserve life of live born person
Whenever an abortion procedure results in a live birth, failure to take all reasonable steps, in keeping with good medical practice, to preserve the life and health of the live born person shall subject the responsible party or parties to Maine law governing homicide, manslaughter and civil liability for wrongful death and medical malpractice.

Definitions:
A. **Live Birth/Live Born:** as used in this chapter, shall mean a product of conception after complete expulsion or extraction from its mother, irrespective of the duration of pregnancy, which breathes or shows any other evidence of life such as beating of the heart, pulsation of the umbilical cord or definite movement of voluntary muscles, whether or not the umbilical cord has been cut or the placenta is attached. Each product of such a birth is considered live born and fully recognized as a human person under Maine law.
Abortion and miscarriage data
Definitions:

A. Abortion: means the intentional interruption of a pregnancy by the application of external agents, whether chemical or physical, or the ingestion of chemical agents with an intention other than to produce a live birth or to remove a dead fetus, regardless of the length of gestation.

B. Miscarriage: means an interruption of a pregnancy other than as provided in paragraph A of a fetus of less than 20 weeks gestation.

22 MRSA §1596 (1)

Abortion reports
A report of each abortion performed shall be made to the Department of Human Services on forms prescribed by the department. These report forms shall not identify the patient by name or otherwise and shall contain only the information requested on the United States Standard Report of Induced Termination of Pregnancy, published by the National Center for Health Statistics, dated January 1978, or any more recent revision of a standard report form.

The form containing that information and data shall be prepared and signed by the attending physician and transmitted to the department not later than 10 days following the end of the month in which the abortion is performed.

A physician who reports data on an abortion pursuant to this section shall be immune from any criminal liability for that abortion under section 1598.

22 MRSA §1596 (2)

Miscarriage reports
A report of each miscarriage shall be made by the physician in attendance at or after the occurrence of the miscarriage to the Department of Human Services on forms prescribed by the department. These report forms shall contain all of the applicable information required on the certificate of fetal death in current use.

The report form shall be prepared and signed by the attending physician and transmitted to the department not later than 10 days following the end of the month in which the miscarriage occurs.

The identity of any patient or physician reporting pursuant to this section is confidential and the department shall take the steps which are necessary to insure the confidentiality of the identity of patients or physicians reporting pursuant to this section.

22 MRSA §1596 (3)

Consent to a minor's decision to have an abortion
Definitions:

A. Abortion: means the intentional interruption of a pregnancy by the application of external agents, whether chemical or physical, or the ingestion of chemical agents with an intention other than to produce a live birth or to remove a dead fetus.

B. Counselor: means a person who is:
   1. A psychiatrist;
   2. A psychologist licensed under Title 32, chapter 56;
   3. A social worker licensed under Title 32, chapter 83;
   4. An ordained member of the clergy;
   5. A physician's assistant registered by the Board of Licensure in Medicine, Title 32, chapter 48;
   6. A nurse practitioner registered by the Board of Licensure in Medicine, Title 32, chapter 48;
7. A certified guidance counselor;
8. A registered professional nurse licensed under Title 32, chapter 31; or

C. **Minor:** means a person who is less than 18 years of age.

**22 MRSA §1597-A (1)**

**Prohibitions; exceptions**
Except as otherwise provided by law, no person may knowingly perform an abortion upon a pregnant minor unless:

A. The attending physician has received and will make part of the medical record the informed written consent of the minor and one parent, guardian or adult family member;
B. The attending physician has secured the informed written consent of the minor as prescribed in subsection 3 and the minor, under all the surrounding circumstances, is mentally and physically competent to give consent;
C. The minor has received the information and counseling required under subsection 4, has secured written verification of receiving the information and counseling and the attending physician has received and will make part of the medical record the informed written consent of the minor and the written verification of receiving information and counseling required under subsection 4; or
D. The Probate Court or District Court issues an order under subsection 6 on petition of the minor or the next friend of the minor for purposes of filing a petition for the minor, granting:
   1. To the minor majority rights for the sole purpose of consenting to the abortion and the attending physician has received the informed written consent of the minor; or
   2. To the minor consent to the abortion, when the court has given its informed written consent and the minor is having the abortion willingly, in compliance with subsection 7.

**22 MRSA §1597-A (2)**

**Informed consent; disallowance of recovery**
No physician may perform an abortion upon a minor unless, prior to performing the abortion, the attending physician received the informed written consent of the minor.

A. To ensure that the consent for an abortion is informed consent, the attending physician shall:
   1. Inform the minor in a manner which, in the physician's professional judgment, is not misleading and which will be understood by the patient, of at least the following:
      a. According to the physician's best judgment the minor is pregnant;
      b. The number of weeks of duration of the pregnancy; and
      c. The particular risks associated with the minor's pregnancy, the abortion technique that may be performed and the risks involved for both;
   2. Provide the information and counseling described in subsection 4 or refer the minor to a counselor who will provide the information and counseling described in subsection 4; and
   3. Determines whether the minor is, under all the surrounding circumstances, mentally and physically competent to give consent.

B. No recovery may be allowed against any physician upon the grounds that the abortion was rendered without the informed consent of the minor when:
   1. The physician, in obtaining the minor's consent, acted in accordance with the standards of practice among members of the same health care profession with similar training and experience situated in the same or similar communities; or
   2. The physician has received and acted in good faith on the informed written consent to the abortion given by the minor to a counselor.

**22 MRSA §1597-A (3)**

**Information and counseling for minors**
The provision of information and counseling by any physician or counselor for any pregnant minor for decision making regarding pregnancy shall be in accordance with this subsection.

A. Any physician or counselor providing pregnancy information and counseling under this subsection shall, in a manner that will be understood by the minor:
1. Explain that the information being given to the minor is being given objectively and is not intended to coerce, persuade or induce the minor to choose either to have an abortion or to carry the pregnancy to term;

2. Explain that the minor may withdraw a decision to have an abortion at any time before the abortion is performed or may reconsider a decision not to have an abortion at any time within the time period during which an abortion may legally be performed;

3. Clearly and fully explore with the minor the alternative choices available for managing the pregnancy, including:
   a. Carrying the pregnancy to term and keeping the child;
   b. Carrying the pregnancy to term and placing the child with a relative or with another family through foster care or adoption;
   c. The elements of prenatal and postnatal care; and
   d. Having an abortion;

4. Explain that public and private agencies are available to provide birth control information and that a list of these agencies and the services available from each will be provided if the minor requests;

5. Discuss the possibility of involving the minor's parents, guardian or other adult family members in the minor's decision making concerning the pregnancy and explore whether the minor believes that involvement would be in the minor's best interests; and

6. Provide adequate opportunity for the minor to ask any questions concerning the pregnancy, abortion, child care and adoption, and provide the information the minor seeks or, if the person cannot provide the information, indicate where the minor can receive the information.

B. After the person provides the information and counseling to a minor as required by this subsection, that person shall have the minor sign and date a form stating that:

   1. The minor has received information on prenatal care and alternatives to abortion and that there are agencies that will provide assistance;
   2. The minor has received an explanation that the minor may withdraw an abortion decision or reconsider a decision to carry a pregnancy to term;
   3. The alternatives available for managing the pregnancy have been clearly and fully explored with the minor;
   4. The minor has received an explanation about agencies available to provide birth control information;
   5. The minor has discussed with the person providing the information and counseling the possibility of involving the minor's parents, guardian or other adult family members in the minor's decision making about the pregnancy;
   6. The reasons for not involving the minor's parents, guardian or other adult family members are put in writing on the form by the minor or the person providing the information and counseling; and
   7. The minor has been given an adequate opportunity to ask questions.

The person providing the information and counseling shall also sign and date the form, and include that person's address and telephone number. The person shall keep a copy for that person's files and shall give the form to the minor or, if the minor requests and if the person providing the information is not the attending physician, transmit the form to the minor's attending physician.

22 MRSA §1597-A (4)

Presumption of validity of informed written consent; rebuttal

An informed consent which is evidenced in writing containing information and statements provided in subsection 4 and which is signed by the minor shall be presumed to be a valid informed consent. This presumption may be subject to rebuttal only upon proof that the informed consent was obtained through fraud, deception or misrepresentation of material fact.
Court order concerning consent to abortion

The court may issue an order for the purpose of consenting to the abortion by the minor under the following circumstances and procedures.

A. The minor or next friend of the minor for the purposes of filing a petition may make an application to the Probate Court or District Court which shall assist the minor or next friend in preparing the petition. The minor or the next friend of the minor shall file a petition setting forth:
   1. The initials of the minor;
   2. The age of the minor;
   3. That the minor has been fully informed of the risks and consequences of the abortion;
   4. That the minor is of sound mind and has sufficient intellectual capacity to consent to the abortion;
   5. That, if the court does not grant the minor majority rights for the purpose of consent to the abortion, the court should find that the abortion is in the best interest of the minor and give judicial consent to the abortion;
   6. That, if the minor does not have private counsel, that the court may appoint counsel.

The minor or the next friend shall sign the petition.

B. The petition is a confidential record and the court files on the petition shall be impounded.

C. A hearing on the merits of the petition shall be held as soon as possible within 5 days of the filing of the petition. If any party is unable to afford counsel, the court shall appoint counsel at least 24 hours before the time of the hearing. At the hearing, the court shall hear evidence relating to:
   1. The emotional development, maturity, intellect and understanding of the minor;
   2. The nature, possible consequences and alternatives to the abortion; and
   3. Any other evidence that the court may find useful in determining whether the minor should be granted majority rights for the purpose of consenting to the abortion or whether the abortion is in the best interest of the minor.

The hearing on the petition shall be held as soon as possible within 5 days of the filing of the petition. The court shall conduct the hearing in private with only the minor, interested parties as determined by the court and necessary court officers or personnel present. The record of the hearing is not a public record.

D. In the decree, the court shall for good cause:
   1. Grant the petition for majority rights for the sole purpose of consenting to the abortion;
   2. Find the abortion to be in the best interest of the minor and give judicial consent to the abortion, setting forth the grounds for the finding; or
   3. Deny the petition only if the court finds that the minor is not mature enough to make her own decision and that the abortion is not in her best interest.

E. If the petition is allowed, the informed consent of the minor, pursuant to a court grant of majority rights or the judicial consent, shall bar an action by the parent or guardian of the minor on the grounds of battery of the minor by those performing the abortion. The immunity granted shall only extend to the performance of the abortion and any necessary accompanying services which are performed in a competent manner.

F. The minor may appeal an order issued in accordance with this section to the Superior Court. The notice of appeal shall be filed within 24 hours from the date of issuance of the order. Any record on appeal shall be completed and the appeal shall be perfected within 5 days from the filing of notice to appeal. The Supreme Judicial Court shall, by court rule, provide for expedited appellate review of cases appealed under this section.
Abortion performed against the minor's will
No abortion may be performed on any minor against her will, except that an abortion may be performed against the will of a minor pursuant to a court order described in subsection 6 that the abortion is necessary to preserve the life of the minor.  

22 MRSA §1597-A (7)

Violations; penalties
A. A person may not knowingly perform or aid in the performance of an abortion in violation of this section. A person who violates this paragraph commits a Class D crime.
B. An attending physician or counselor may not knowingly fail to perform any action required by this section. A person who violates this paragraph commits a civil violation for which a fine of not more than $1,000 may be adjudged for each violation.

22 MRSA §1597-A (8)

Nonseverability
In the event that any portion of this section is held invalid, it is the intent of the Legislature that this entire section shall be invalid.

22 MRSA §1597-A (9)

(4) Abuse, Adult
Select Definitions
A. Abuse: means the infliction of injury, unreasonable confinement, intimidation or cruel punishment that causes or is likely to cause physical harm or pain or mental anguish; sexual abuse or sexual exploitation; financial exploitation; or the intentional, knowing or reckless deprivation of essential needs. “Abuse” includes acts and omissions.
B. Adult: means any person who has attained 18 years of age or who is a legally emancipated minor.
C. Dependent adult: means an adult who has a physical or mental condition that substantially impairs the adult's ability to adequately provide for that adult's daily needs. “Dependent adult” includes, but is not limited to, any of the following:
   1. A resident of a nursing home licensed or required to be licensed under section 1817;
   2. A resident of a facility providing assisted living services licensed or required to be licensed pursuant to section 7801;
   3. A person considered a dependent person under Title 17-A, section 555; or
   4. A person, regardless of where that person resides, who is wholly or partially dependent upon one or more other persons for care or support, either emotional or physical, because the person suffers from a significant limitation in mobility, vision, hearing or emotional or mental functioning.
D. Incapacitated adult: means any adult who is impaired by reason of mental illness, mental deficiency, physical illness or disability to the extent that that individual lacks sufficient understanding or capacity to make or communicate responsible decisions concerning that individual's person, or to the extent the adult cannot effectively manage or apply that individual's estate to necessary ends
E. Exploitation: means the illegal or improper use of an incapacitated or dependent adult or that adult's resources for another's profit or advantage.
F. Financial exploitation: means the use of deception, intimidation, undue influence, force or other unlawful means to obtain control over the property of a dependent adult for another's profit or advantage.
G. Neglect: means a threat to an adult's health or welfare by physical or mental injury or impairment, deprivation of essential needs or lack of protection from these.
H. Serious harm: means:
   1. Serious physical injury or impairment;
2. Serious mental injury or impairment that now or in the future is likely to be evidenced by serious mental, behavioral or personality disorder, including, but not limited to, severe anxiety, depression or withdrawal, untoward aggressive behavior or similar serious dysfunctional behavior;
3. Sexual abuse or sexual exploitation; or
4. Serious waste or dissipation of resources.

Spiritual treatment

Treatment not considered abuse, neglect or exploitation
An incapacitated or dependent adult shall not be considered to be abused, neglected or exploited solely because treatment is by spiritual means by an accredited practitioner of a recognized religious organization.

Treatment to be considered if requested
When medical treatment is authorized, under this chapter, treatment by spiritual means by an accredited practitioner of a recognized religious organization may also be considered if requested by the incapacitated or disabled adult or his caretaker.

Persons mandated to report suspected abuse, neglect or exploitation

Persons required to report
The following persons immediately shall report or cause a report to be made to the department when the person suspects that an adult has been abused, neglected or exploited and has reasonable cause to suspect that the adult is incapacitated or dependent:

A. While acting in a professional capacity:
   1. An allopathic or osteopathic physician;
   2. A medical resident or intern;
   3. A medical examiner;
   4. A physician's assistant;
   5. A dentist, dental hygienist or dental assistant;
   6. A chiropractor;
   7. A podiatrist;
   8. A registered or licensed practical nurse;
   9. A certified nursing assistant;
   10. A social worker;
   11. A psychologist;
   12. A pharmacist;
   13. A physical therapist;
   14. A speech therapist;
   15. An occupational therapist;
   16. A mental health professional;
   17. A law enforcement official, corrections officer or other person holding a certification from the Maine Criminal Justice Academy;
   18. Emergency room personnel;
   19. An ambulance attendant;
   20. An emergency medical technician or other licensed medical service provider;
   21. Unlicensed assistive personnel;
   22. A humane agent employed by the Department of Agriculture, Conservation and Forestry;
   23. A clergy member acquiring the information as a result of clerical professional work except for information received during confidential communications;
   24. A sexual assault counselor;
   25. A family or domestic violence victim advocate;
26. A naturopathic doctor;
27. A respiratory therapist;
28. A court-appointed guardian or conservator; or
29. A chair of a professional licensing board that has jurisdiction over mandated reporters;

B. Any person who has assumed full, intermittent or occasional responsibility for the care or custody of the adult, regardless of whether the person receives compensation;

C. Any person affiliated with a church or religious institution who serves in an administrative capacity or has otherwise assumed a position of trust or responsibility to the members of that church or religious institution, while acting in that capacity, regardless of whether the person receives compensation; or

D. Any person providing transportation services as a volunteer or employee of an agency, business or other entity, whether or not the services are provided for compensation.

The duty to report under this subsection applies to individuals who must report directly to the department. A supervisor or administrator of a person making a report under this section may not impede or inhibit the reporting, and a person making a report may not be subject to any sanction for making a report. Internal procedures to facilitate ensure confidentiality of and apprise supervisors and administrators of reports may be established as long as those procedures are not inconsistent with this chapter.

22 MRSA §3477 (1)

Permitted reporters
An animal control officer, as defined in Title 7, section 3908, subsection 4, may report to the department when that person has reasonable cause to suspect that an incapacitated or dependent adult has been or is at substantial risk of abuse, neglect or exploitation.

22 MRSA §3477 (1-A)

Reports
Reports regarding abuse, neglect or exploitation shall be made immediately by telephone to the department and shall be followed by a written report within 48 hours if requested by the department. The reports shall contain the name and address of the involved adult; information regarding the nature and extent of the abuse, neglect or exploitation; the source of the report; the person making the report; the person’s occupation; and where that person can be contacted. The report may contain any other information that the reporter believes may be helpful.

22 MRSA §3477 (2)

Confidentiality in case of treatment of individual suspected of causing abuse, neglect or exploitation
This section does not require any person acting in that person’s professional capacity to report when all of the following requirements are met:

A. The factual basis for knowing or suspecting abuse, neglect or exploitation of an adult covered under this subchapter derives from the professional's treatment of the individual suspected of causing the abuse, neglect or exploitation;

B. The treatment was sought by the individual for a problem relating to the abuse, neglect or exploitation; and

C. In the opinion of the person required to report, the abused, neglected or exploited adult's life or health is not immediately threatened.

22 MRSA §3477 (3)

Confidentiality in case of treatment of individual suspected of being abused, neglected, or exploited
This section does not require any person acting in that person’s professional capacity to report when all of the following requirements are met:

A. The factual basis for knowing or suspecting abuse, neglect or exploitation of an adult covered under this subchapter derives from the professional’s treatment of the individual suspected of being abused, neglected or exploited.

B. The treatment was sought by the individual for a problem relating to the abuse, neglect or exploitation, and
C. In the opinion of the person required to report, the individual is not incapacitated and the individual’s life or health is not immediately threatened.  

22 MRSA §3477 (4)

Permissive reporting of animal cruelty, abuse or neglect
Notwithstanding any other provision of state law imposing a duty of confidentiality, a person listed in subsection 1 may report a reasonable suspicion of animal cruelty, abuse or neglect to the local animal control officer or to the animal welfare program of the Department of Agriculture, Food and Rural Resources established pursuant to Title 7, section 3902. For purposes of this subsection, the reporter shall disclose only such limited confidential information as is necessary for the local animal control officer or animal welfare program employee to identify the animal’s location and status and the owner’s name and address. For purposes of this subsection, “cruelty, abuse or neglect” has the same meaning as provided in Title 34-B, chapter 1, subchapter 6.

22 MRSA §3477 (5)

Mandatory reporting to medical examiner for post-mortem investigation
A person required to report cases of known or suspected abuse or neglect, who knows or has reasonable cause to suspect that an adult has died as a result of abuse or neglect, shall report that fact to the appropriate authority as provided in section 3026. An adult shall not be considered to be abused or neglected solely because he was provided with treatment by spiritual means by an accredited practitioner of a recognized religious organization.

22 MRSA §3478

Optional reporting
Any person may make a report if that person knows or has reasonable cause to suspect abuse, neglect or exploitation of an incapacitated or dependent adult, or has reasonable cause to suspect that an adult is incapacitated.

22 MRSA §3479

Immunity from liability
Reporting and proceedings
A person participating in good faith in reporting under this subchapter, or in a related adult protection investigation or proceeding, is immune from any civil liability that might otherwise result from these actions.

Presumption of good faith
In a proceeding regarding immunity from liability, there shall be a rebuttable presumption of good faith.

22 MRSA §3479-A

(5) Abuse, Child
Spiritual treatment
Treatment not considered abuse or neglect
Under subchapters I to VII, a child shall not be considered to be abused or neglected, in jeopardy of health or welfare or in danger of serious harm solely because treatment is by spiritual means by an accredited practitioner of a recognized religious organization.

Treatment to be considered if requested
When medical treatment is authorized under this chapter, treatment by spiritual means by an accredited practitioner of a recognized religious organization may also be considered if requested by the child or his parent.

22 MRSA §4010
**Reporting of suspected child abuse or neglect**

**Persons required to report**
The following adult persons shall immediately report or cause a report to be made to the department when the person knows or has reasonable cause to suspect that a child has been or is likely to be abused or neglected or that a suspicious child death has occurred:

A. When acting in a professional capacity:
   1. An allopathic or osteopathic physician, resident or intern;
   2. An emergency medical services person;
   3. A medical examiner;
   4. A physician's assistant;
   5. A dentist;
   6. A dental hygienist;
   7. A dental assistant;
   8. A chiropractor;
   9. A podiatrist;
   10. A registered or licensed practical nurse;
   11. A teacher;
   12. A guidance counselor;
   13. A school official;
   14. A youth camp administrator or counselor;
   15. A social worker;
   16. A court-appointed special advocate or guardian ad litem for the child;
   17. A homemaker;
   18. A home health aide;
   19. A medical or social service worker;
   20. A psychologist;
   21. Child care personnel;
   22. A mental health professional;
   23. A law enforcement official;
   24. A state or municipal fire inspector;
   25. A municipal code enforcement official;
   26. A commercial film and photographic print processor;
   27. A clergy member acquiring the information as a result of clerical professional work except for information received during confidential communications;
   28. A chair of a professional licensing board that has jurisdiction over mandated reporters;
   29. A humane agent employed by the Department of Agriculture, Food and Rural Resources;
   30. A sexual assault counselor; and
   31. A family or domestic violence victim advocate; and
   32. A school bus driver or school bus attendant;

B. Any person who has assumed full, intermittent or occasional responsibility for the care or custody of the child, regardless of whether the person receives compensation; and

C. Any person affiliated with a church or religious institution who serves in an administrative capacity or has otherwise assumed a position of trust or responsibility to the members of that church or religious institution, while acting in that capacity, regardless of whether the person receives compensation.

Whenever a person is required to report in a capacity as a member of the staff of a medical or public or private institution, agency or facility, that person immediately shall notify either the person in charge of the institution, agency or facility or a designated agent who then shall cause a report to be made. The staff also may make a report directly to the department. If a person required to report notifies either the person
in charge of the institution, agency or facility or the designated agent, the notifying person shall acknowledge in writing that the institution, agency or facility has provided confirmation to the notifying person that another individual from the institution, agency or facility has made a report to the department. The confirmation must include, at a minimum, the name of the individual making the report to the department, the date and time of the report and a summary of the information conveyed. If the notifying person does not receive the confirmation from the institution, agency or facility within 24 hours of the notification, the notifying person immediately shall make a report directly to the department.

An employer may not take any action to prevent or discourage an employee from making a report.

22 MRSA §4011-A (1)

Optional report

Any person may make a report if that person knows or has reasonable cause to suspect that a child has been or is likely to be abused or neglected or that there has been a suspicious child death.

22 MRSA §4011-A (2)

Mental health treatment

When a licensed mental health professional is required to report under subsection 1 and the knowledge or reasonable cause to suspect that a child has been or is likely to be abused or neglected or that a suspicious child death has occurred comes from treatment of a person responsible for the abuse, neglect or death, the licensed mental health professional shall report to the department in accordance with subsection 1 and under the following conditions.

A. The department shall consult with the licensed mental health professional who has made the report and shall attempt to reach agreement with the mental health professional as to how the report is to be pursued. If agreement is not reached, the licensed mental health professional may request a meeting under paragraph B.

B. Upon the request of the licensed mental health professional who has made the report, after the department has completed its investigation of the report under section 4021 or has received a preliminary protection order under section 4034 and when the department plans to initiate or has initiated a jeopardy order under section 4035 or plans to refer or has referred the report to law enforcement officials, the department shall convene at least one meeting of the licensed mental health professional who made the report, at least one representative from the department, a
licensed mental health professional with expertise in child abuse or neglect and a representative of the district attorney's office having jurisdiction over the report, unless that office indicates that prosecution is unlikely.

C. The persons meeting under paragraph B shall make recommendations regarding treatment and prosecution of the person responsible for the abuse, neglect or death. The persons making the recommendations shall take into account the nature, extent and severity of abuse or neglect, the safety of the child and the community and needs of the child and other family members for treatment of the effects of the abuse or neglect and the willingness of the person responsible for the abuse, neglect or death to engage in treatment. The persons making the recommendations may review or revise these recommendations at their discretion.

The intent of this subsection is to encourage offenders to seek and effectively utilize treatment and, at the same time, provide any necessary protection and treatment for the child and other family members.  

22 MRSA §4011-A (4)

Photographs of visible trauma
Whenever a person is required to report as a staff member of a law enforcement agency or a hospital, that person shall make reasonable efforts to take, or cause to be taken, color photographs of any areas of trauma visible on a child.

A. The taking of photographs must be done with minimal trauma to the child and in a manner consistent with professional standards. The parent's or custodian's consent to the taking of photographs is not required.

B. Photographs must be made available to the department as soon as possible. The department shall pay the reasonable costs of the photographs from funds appropriated for child welfare services.

C. The person shall notify the department as soon as possible if that person is unable to take, or cause to be taken, these photographs.

D. Designated agents of the department may take photographs of any subject matter when necessary and relevant to an investigation of a report of suspected abuse or neglect or to subsequent child protection proceedings.

22 MRSA §4011-A (5)

Children Under 6 Months
In an infant under the age of six months or otherwise non-ambulatory, a mandated reporter must report the following conditions to Department of Health and Human Services: fractures of a bone; substantial bruising or multiple bruises; subdural hematoma; burns; poisoning, or injury resulting in substantial bleeding, soft tissue swelling, or impairment of an organ.

This subsection does not require the reporting of injuries occurring as a result of the delivery of a child attended by a licensed medical practitioner or the reporting of burns or other injuries occurring as a result of medical treatment following the delivery of the child while the child remains hospitalized following the delivery.

22 MRSA §4011-A (7)

Required report of residence with nonfamily
A person required to make a report shall report to the department if the person knows or has reasonable cause to suspect that a child is not living with the child's family. Although a report may be made at any time, a report must be made immediately if there is reason to suspect that a child has been living with someone other than the child's family for more than 6 months or if there is reason to suspect that a child has been living with someone other than the child's family for more than 12 months pursuant to a power of attorney or other nonjudicial authorization.

22 MRSA §4011-A (8)

Training requirement
A person required to make a report shall complete at least once every 4 years mandated reporter training approved by the department.
Enhanced Enforcement
DHHS is required to report to the licensing board of a professional who appears to have violated the mandatory reporting law.

Prenatal Drug Exposure
A health care provider involved in the delivery or care of an infant who the provider knows or has reasonable cause to suspect has been born affected by illegal substance abuse, is demonstrating withdrawal symptoms that require medical monitoring or care beyond standard newborn care when those symptoms have resulted from or have likely resulted from prenatal drug exposure, whether the prenatal exposure was to legal or illegal drugs, or has fetal alcohol spectrum disorders, shall notify the department of that condition in the infant.

DHHS will be required to make a report to the licensing board of professional that appears to have violated the mandatory reporting law.

Reporting procedures
Immediate report
Reports regarding abuse or neglect shall be made immediately by telephone to the department and shall be followed by a written report within 48 hours if requested by the department.

Hospitals, medical personnel and law enforcement personnel may submit emergency reports through password-protected e-mail submissions. A faxed report may also be accepted when preceded by a telephone call informing the department of the incoming fax transmission.

Information required
The reports shall include the following information if within the knowledge of the person reporting:
A. The name and address of the child and the persons responsible for his care or custody;
B. The child's age and sex;
C. The nature and extent of abuse or neglect, including a description of injuries and any explanation given for them;
D. A description of sexual abuse or exploitation;
E. Family composition and evidence of prior abuse or neglect of the child or his siblings;
F. The source of the report, the person making the report, his occupation and where he can be contacted;
G. The actions taken by the reporting source, including a description of photographs or x rays taken; and
H. Any other information that the person making the report believes may be helpful.

Immunity from liability
Reporting and proceedings
A person, including an agent of the department, participating in good faith in reporting under this subchapter or participating in a related child protection investigation or proceeding, including, but not limited to, a multidisciplinary team, out-of-home abuse investigating team or other investigating or
treatment team, is immune from any criminal or civil liability for the act of reporting or participating in the investigation or proceeding. Good faith does not include instances when a false report is made and the person knows the report is false. Nothing in this section may be construed to bar criminal or civil action regarding perjury or regarding the abuse or neglect which led to a report, investigation or proceeding.

22 MRSA §4014 (1)

Photographs and x rays
A person participating in good faith in taking photographs or x rays under this subchapter is immune from civil liability for invasion of privacy that might otherwise result from these actions.

22 MRSA §4014 (2)

Presumption of good faith
In a proceeding regarding immunity from liability, there shall be a rebuttable presumption of good faith.

22 MRSA §4014 (3)

Privileged or confidential communications
The husband-wife and physician and psychotherapist-patient privileges under the Maine Rules of Evidence and the confidential quality of communication under Title 16, section 53-B (“Privileged communications to victim advocate; family violence”); Title 20-A, sections 4008 (“Education: Privileged communications”) and 6001 (“Student records: Dissemination of information”), to the extent allowed by applicable federal law; Title 24-A, section 4224 (“Health maintenance organizations: Confidentiality; liability; access to records”); Title 32, sections 7005 (“Communication between social workers and clients”) and 18393 (“Dental professions: Confidentiality”); and Title 34-B, section 1207 (“Behavioral and developmental services: Confidentiality of information”), are abrogated in relation to required reporting, cooperating with the department or a guardian ad litem in an investigation or other child protective activity or giving evidence in a child protection proceeding. Information released to the department pursuant to this section must be kept confidential and may not be disclosed by the department except as provided in section 4008 (“DHHS reports to licensing boards”).

Statements made to a licensed mental health professional in the course of counseling, therapy or evaluation where the privilege is abrogated under this section may not be used against the client in a criminal proceeding. Nothing in this section may limit any responsibilities of the professional pursuant to this Act.

22 MRSA §4015

Discrimination
No person may be discriminated against by any employer in any way for participating in good faith in reporting under this subchapter or in a related child protection investigation or proceeding.

22 MRSA §4017

Abandoned child; safe haven provider
Definitions
A. Medical services provider: means an individual certified, registered or licensed in the healing arts, including, but not limited to, a physician, nurse, podiatrist, optometrist, chiropractor, physical therapist, dentist, psychologist, physician's assistant or emergency medical services person.
B. Safe haven provider: means:
   1. A law enforcement officer;
   2. Staff at a medical emergency room;
   3. A medical services provider; or
   4. A hospital staff member at a hospital.

22 MRSA §4018 (1)
Request for information
A person who voluntarily delivers a child less than 31 days of age to a safe haven provider and who does not express an intent to return for the child may be requested to provide information helpful to the welfare of the child. The person who accepts a child under this section may not detain the person delivering the child to obtain information.

22 MRSA §4018 (2)

Action by safe haven provider; guidelines
A safe haven provider who accepts a child under this section shall promptly notify the department of the delivery of the child, transfer the child to the department at the earliest opportunity and provide to the department all information provided by the person delivering the child to the safe haven provider. The department shall establish guidelines to assist safe haven providers concerning procedures when a child is delivered to a safe haven provider under this section.

22 MRSA §4018 (3)

Confidentiality
All personally identifiable information provided by the person delivering the child to a safe haven provider is confidential and may not be disclosed by the safe haven provider to anyone except to the extent necessary to provide temporary custody of the child until the child is transferred to the department and except as otherwise provided by court order. All health care or other information obtained by a safe haven provider in providing temporary custody of the child may also be provided to the department upon request.

22 MRSA §4018 (4)

Liability
A person or entity who accepts a child under this section or provides temporary custody of a child accepted under this section is not subject to civil, criminal or administrative liability for accepting the child or providing temporary custody of the child in the good faith belief that the action is required or authorized by this section. This subsection does not affect liability for personal injury or wrongful death, including, but not limited to, injury resulting from medical malpractice.

22 MRSA §4018 (5)

(6) Advanced Directives (Formerly Known as Living Wills; see Power of Attorney for Healthcare)

(7) Academic Medical Centers (see Research)

(8) Advertising

Advertising
A health care practitioner who advertises health care services shall disclose in an advertisement the applicable license under which the health care practitioner is authorized to provide services. The advertisement:

A. May not constitute deceptive or misleading advertising; and
B. Must include the health care practitioner's name, the type of license the practitioner holds and the common term for the practitioner's profession.

24 MRSA §2988 (2)

Identification
A health care practitioner seeing patients on a face-to-face basis shall wear a name badge or some other form of identification that clearly discloses:
A. The health care practitioner's first name or first and last name, except that if the health care practitioner is a physician, the name badge or identification must disclose the physician's first and last name; and
B. The type of license, registration or certification the health care practitioner holds, including the common term for the health care practitioner's profession.

**24 MRSA §2988 (3)**

**Complaints; disciplinary action**
A person may file a complaint with the appropriate licensing board regarding a health care practitioner who fails to provide the consumer information required in this section. A health care practitioner who violates any provision of this section engages in unprofessional conduct and is subject to disciplinary action under the applicable licensing provisions of the health care practitioner.

**24 MRSA §2988 (3)**

**AMA’s Policy and Guidance on Advertising and Publicity**

**Grounds for Discipline**
The board [of licensing] may suspend or revoke a license pursuant to Title 5, section 10004. The following are grounds for an action to refuse to issue, modify, restrict, suspend, revoke or refuse to renew the license of an individual licensed under this chapter:

A. I. Engaging in false, misleading or deceptive advertising;

**32 MRSA §3282-A (2)(I)**

(9) AIDS and HIV

**Confidentiality of test**
No person may disclose the results of an HIV test, except as follows:

A. **Subject of test.** To the subject of the test;
B. **Designated health care provider.** To a health care provider designated by the subject of the test in writing. When a patient has authorized disclosure of HIV test results to a person or organization providing health care, the patient's health care provider may make these results available only to other health care providers working directly with the patient and only for the purpose of providing direct medical or dental patient care. Any health care provider who discloses HIV test results in good faith pursuant to this subsection is immune from any criminal or civil liability for the act of disclosing HIV test results to other health care providers;
C. **Authorized person.** To a person or persons to whom the test subject has authorized disclosure in writing, except that the disclosure may not be used to violate any other provisions of this chapter;
D. **Certain health care providers.** A health care provider who procures, processes, distributes or uses a human body part donated for a purpose may, without obtaining informed consent to the testing, perform an HIV test in order to assure medical acceptability of the gift for the purpose intended. Testing pursuant to this subsection does not require pretest and post-test counseling;
E. **Research facility.** The Department of Human Services, a laboratory certified and approved by the Department of Human Services pursuant to Title 22, chapter 411, or a health care provider, blood bank, blood center or plasma center may, for the purpose of research and without first obtaining informed consent to the testing, subject any body fluids or tissues to an HIV test if the testing is performed in a manner by which the identity of the test subject is not known and may not be retrieved by the researcher;
F. **Anonymous testing sites.** To an anonymous testing site established pursuant to section 19203-B;
G. **Other agencies.** To employees of, or other persons designated by, the Department of Corrections, the Department of Human Services and the Department of Behavioral and Developmental
Services, to the extent that those employees or other persons are responsible for the treatment or care of subjects of the test. Those agencies shall adopt rules, within 90 days of August 4, 1988, pursuant to chapter 375, subchapter II, designating the persons or classes of persons to whom the test results may be disclosed. The rules of the Department of Corrections must designate those persons who may receive the results of an HIV test of a county jail inmate;

H. Bureau of Health. To the Bureau of Health, which may disclose results to other persons only if that disclosure is necessary to carry out its duties as provided in Title 22, sections 3, 7 and 42 and chapters 250 and 251;

I. Medical records. As part of a medical record when release or disclosure of that record is authorized pursuant to section 19203-D (“Medical conditions: Records”); or

J. Court ordered disclosure.
   1. A person authorized by section 19203-C (“Medical conditions: Judicial consent to HIV test”) to receive test results following an accidental exposure; or
   A victim-witness advocate authorized by section 19203-F to receive the test results of a person convicted of a sexual crime as defined in section 19203-F, subsection 1, paragraph C, who shall disclose to a victim under section 19203-F, subsection 4.

This section does not prohibit limited administrative disclosure in conjunction with a mandatory testing program of a military organization subject to Title 37-B.

Nothing in this section may be construed as prohibiting the entry of an HIV test result on the patient's medical record in accordance with this chapter.

5 MRSA §19203

Informed consent required

Individual tested

Except as provided in this section and section 19203, subsections 4 (“Certain health care providers”) and 5 (“Research facility”), an HIV test must be voluntary and undertaken only with a patient’s knowledge and understanding that an HIV test is planned. A patient must be informed orally or in writing that an HIV test will be performed unless the patient declines. Oral or written information required to be given to a patient under this subsection must include an explanation of what an HIV infection involves and the meaning of positive and negative test results. A patient must be provided the opportunity to ask questions, either orally or in writing. Informed consent is not required for repeated HIV testing by health care providers to monitor the course of established infection.

5 MRSA §19203-A (1)

Insurers

Persons required to take an HIV test by an insurer, nonprofit hospital or medical service organization or nonprofit health care plan must provide their written informed consent on forms approved by the Superintendent of Insurance. If the test is positive, post-test counseling must be provided by the person or organization requesting the test. The Superintendent of Insurance may adopt rules to define language requirements of the form.

5 MRSA §19203-A (2)

Access to medical care

A health care provider may not deny any person medical treatment or care solely for refusal to give consent for an HIV test. A health care provider may not request a person's written consent to an HIV test as a precondition to the provision of health care. All written consent to testing shall be in accordance with section 19201, subsection 5-A. This section does not prohibit a health care provider from recommending an HIV test for diagnostic or treatment purposes. A physician or other health care provider is not civilly liable for failing to have an HIV test performed for diagnostic or treatment purposes if the test was recommended and refused in writing by the patient.

5 MRSA §19203-A (3)
Occupational exposure
Consent need not be obtained when a bona fide occupational exposure creates a significant risk of infection if a court order has been obtained under section 19203-C. The fact that an HIV test was given as a result of an occupational exposure and the results of that test may not appear in any records of the person whose blood or body fluid is the source of the exposure. If the test is positive, post-test counseling must be offered. The subject of the test may choose not to be informed about the result of the test.

5 MRSA §19203-A (4)

Occupational exposure in health care setting
When a bona fide occupational exposure occurs in a health care setting, authorization to test the source patient for HIV must be obtained from that patient if the patient is present or can be contacted at the time of exposure and is capable of providing consent. At the time of exposure, if the source patient is not present and can not be contacted or is incapacitated, then any reasonably available member of the following classes of individuals, in descending order of priority, may authorize an HIV test on a blood or tissue sample from the source patient:

A. The patient's legal guardian;
B. An individual known to have power of attorney for health care for the patient;
C. An adult relative, by blood, marriage or adoption;
D. An adult with whom the patient has a meaningful social and emotional relationship; and
E. A physician who is familiar with occupational exposures to HIV.

The individual authorizing the HIV test must be informed of the nature, reliability and significance of the HIV test and the confidential nature of the test.

If the person contacted for authorization refuses to authorize the test, the test may not be conducted unless consent is obtained from the source patient or from the court pursuant to section 19203-C.

This subsection does not authorize a person described in paragraphs A to D to receive the test result. Test results must be given to the exposed person, to a personal physician if designated by the exposed person and to either the physician who authorizes the test or the health care provider who manages the occupational exposure.

The patient may choose not to be informed about the result of the HIV test. Without express patient authorization, the results of the HIV test and the fact that an HIV test was done as a result of an occupational exposure in a health care setting may not appear in the patient's health care records. The exposed individual's occupational health care record may include documentation of the occupational exposure and, if the record does not reveal the source patient's identity, the results of the source patient's HIV test.

5 MRSA §19203-A (4-A)

Judicial consent to HIV test for occupational exposure
Any person who experiences a bona fide occupational exposure may petition the District Court with jurisdiction over the facility or other place where the exposure occurred to require the person whose blood or body fluid is the source of the exposure to submit to an HIV test and to require that the results of the test be provided to the petitioner. For more information on this process, see 5 MRSA §19203-C.

Exposure from sexual crime
Consent need not be obtained when a court order has been issued under section 19203-F. The fact that an HIV test was given as a result of the exposure and the results of that test may not appear in a convicted offender's medical record. Counseling on risk reduction must be offered, but the convicted offender may choose not to be informed about the result of the test unless the court has ordered that the convicted offender be informed of the result.
Preventing HIV Transmission from Pregnant Mother to Child

Maine's HIV testing and consent law, was amended in 2011 to include a sixth subsection on the “protection of newborn infants.” The amended section of the law requires health care providers caring for pregnant women to include HIV tests in the standard set of medical tests performed, subject to the consent and procedure requirements of 5 MRSA §19203-A, sub-§ 1. That section requires that a patient must be informed orally or in writing that an HIV test will be performed, unless the patient declines.

Under the law, health care providers caring for newborn infants are also required to test the infant for HIV and ensure that the results are available within 12 hours of birth if the health care provider does not know the HIV status of the mother or the health care provider believes that HIV testing is medically necessary. There is an exception if a parent objects to the test on the grounds that it conflicts with the sincere religious or conscientious beliefs and practices of the parent. The full text of the law is available by clicking here.

Records

When a medical record entry is made concerning information of a person's HIV infection status, including the results of an HIV test, the following apply to the release of that information as a part of the medical record.

Authorized release

The person who is the subject of an HIV test, at or near the time the entry is made in the medical record, shall elect, in writing, whether to authorize the release of that portion of the medical record containing the HIV infection status information when that person's medical record has been requested. A new election may be made when a change in the person's HIV infection status occurs or whenever the person makes a new election. The release form must clearly state whether or not the person has authorized the release of that information. The person must be advised of the potential implications of authorizing the release of that information.

A. When release has been authorized, the custodian of the medical record may release, upon request, the person's medical record, including any HIV infection status information contained in the medical record. Release of HIV infection status information pursuant to this paragraph is not a violation of any of the confidentiality provisions of this chapter.
B. When release has not been authorized, the custodian of the medical record may, upon request, release that portion of the medical record that does not contain the HIV infection status information. Except as otherwise provided in this section, HIV infection status information may be released only if the person has specifically authorized a separate release of that information. A general release form is insufficient.

Authorized disclosure

A medical record containing results of an HIV test may not be disclosed, discoverable or compelled to be produced in any civil, criminal, administrative or other proceedings without the consent of the person who is the subject of an HIV test, except in the following cases:

A. Proceedings held pursuant to the communicable disease laws, Title 22, chapter 251;
B. Proceedings held pursuant to the Adult Protective Services Act, Title 22, chapter 958-A;
C. Proceedings held pursuant to the child protection laws, Title 22, chapter 1071;
D. Proceedings held pursuant to the mental health laws, Title 34-B, chapter 3, subchapter IV, article III; and
E. Pursuant to a court order upon a showing of good cause, provided that the court order limits the use and disclosure of records and provides sanctions for misuse of records or sets forth other methods for ensuring confidentiality.

5 MRSA §19203-D (2)

Utilization review; research
Nothing in this section may be interpreted to prohibit reviews of medical records for utilization review purposes by duly authorized utilization review committees or peer review organizations. Qualified personnel conducting scientific research, management audits, financial audits or program evaluation with the use of medical records may not identify, directly or indirectly, any individual patient in any report of such research, audit, evaluation or otherwise disclose the identities of persons tested in any manner.

5 MRSA §19203-D (3)

Access by health care providers
Nothing in this section may prohibit access to medical records by the designated health care provider of the person who is the subject of an HIV test in accordance with section 19203, subsection 2.

5 MRSA §19203-D (4)

Confidentiality policy (see also Health Insurance Portability and Accountability Act)
Health care providers and others with access to medical records containing HIV infection status information shall have a written policy providing for confidentiality of all patient information consistent with this chapter. That policy must require, at a minimum, action consistent with disciplinary procedures for violations of the confidentiality policy.

5 MRSA §19203-D (5)

HIV test after conviction for sexual assault

Request for testing
A person who is the victim of a sexual crime, or that person's parent, guardian or authorized representative if that person is a minor or incapacitated adult, may petition the court at any time prior to sentencing or no later than 180 days after conviction to order the convicted offender to submit to HIV testing and to order that the convicted offender be informed of the test results.

5 MRSA §19203-F (2)

Reporting and counseling
The health care facility in which a convicted offender is tested pursuant to this section shall disclose the results of the test to the victim-witness advocate, who shall disclose the result to the petitioner. The health care facility shall, upon order of the court, disclose the results of the test to the convicted offender.

5 MRSA §19203-F (4)

Counseling new HIV cases
Except as otherwise provided by this chapter, persons who test positive for HIV infection must be offered post-test counseling. Persons who are authorized by section 19203-C (“Judicial consent to HIV test”) or 19203-F (“HIV test after conviction for sexual assault”) to receive test results after exposure must be offered counseling regarding the nature, reliability and significance of the HIV test and the confidential nature of the test. Persons offered counseling under this section may decline the offer by signing a waiver stating that counseling has been offered and is being declined.

5 MRSA §19204-A

Post-test counseling
"Post-test counseling" must include:

A. Personal counseling that includes, at a minimum, a discussion of:
   1. The test results and the reliability and significance of the test results. The person providing post-test counseling shall communicate the result confidentially and through personal contact;
   Information on good preventive practices and risk reduction plans; and
Referrals for medical care and information and referrals for support services, including social, emotional support and legal services, as needed;
B. An entry in the medical record of the person being counseled summarizing the contents of the discussion; and
C. The offer of face-to-face counseling. If the subject of the test declines, the provider of the test may provide an alternative means of providing the information required by paragraph A.

Written information to person being counseled
To comply with the requirements of this section regarding post-test counseling, in addition to meeting the requirements of subsection 2, the provider of an HIV test shall give to the person being counseled a written document containing information on the subjects described in subsection 2, paragraph A.

Restrictions on requiring tests or results of tests

Employee testing
An employee or applicant for employment may not be required to submit to an HIV test or reveal whether the employee or applicant for employment has obtained an HIV test as a condition of employment or to maintain employment, except when based on a bona fide occupational qualification. The Maine Human Rights Commission shall enforce this subsection.

Employee rights
The employment status of any employee may not be affected or changed:
A. If the employee declines to be tested;
B. If the employee testifies or assists in any proceeding under this chapter;
C. If the employee asserts any other rights exercised in good faith pursuant to this chapter; or
D. Because of the result of any test taken pursuant to this chapter.

Restrictions upon revealing HIV antibody test results
An insurer, nonprofit hospital or medical services organization, nonprofit health care plan or health maintenance organization may not request any person to reveal whether the person has obtained a test for the presence of antibodies to HIV or a test to measure the virus or to reveal the results of such tests taken prior to an application for insurance coverage.

Civil liability
Any person violating this chapter is liable to the subject of the test for actual damages and costs plus a civil penalty of up to $1,000 for a negligent violation and up to $5,000 for an intentional violation, subject to Title 14, chapter 741 (“Tort claims”).

Any person may bring an action for injunctive relief for a violation of sections 19203 (“Confidentiality of test”) and 19204 (“Restrictions upon revealing HIV test results”) in addition to or instead of the penalties provided in this section. The applicant for injunctive relief under this section shall not be required to give security as a condition upon the issuance of the injunction.

Insurance (Special Restrictions in AIDS/HIV Cases)
Prohibitions
No individual or group hospital, medical or health care service contract delivered or issued for delivery in this State, other than a contract that provides benefits for specific diseases or accidental injuries only, may
provide more restrictive coverage for Acquired Immune Deficiency Syndrome, or AIDS, AIDS Related Complex, or ARC, HIV-related diseases or for related services, than for any other disease or sickness, or exclude coverage for AIDS, ARC or HIV-related diseases, except through an exclusion under which all diseases and sicknesses are treated equally.  

Test results
No nonprofit hospital or medical services organization or nonprofit health care plan may request any person to reveal whether the person has obtained a test for the presence of antibodies to HIV or a test to measure the virus or to reveal the results of such tests taken prior to an application for coverage.  

Expedited Partner Therapy (see Public Health)

(10) Ambulatory Surgical Facilities
Ambulatory surgical facility means a facility with a primary purpose of providing elective surgical care to a patient who is admitted to and discharged from the facility within the same day. In order to meet this primary purpose, a facility must at least administer anesthetic agents, maintain a sterile environment in a surgical suite and charge a facility fee separate from the professional fee. "Ambulatory surgical facility" does not include:

A. A facility that is licensed as part of a hospital;
B. A facility that provides services or accommodations for patients who stay overnight;
C. A facility existing for the primary purpose of performing terminations of pregnancies; or
D. The private office of a physician or dentist in individual or group practice, unless that facility or office is certified as a Medicare ambulatory surgical center.  

Standards
The Department of Health and Human Services rules governing Ambulatory Surgical Facilities can be found here.

Annual inspection
The department shall inspect annually ambulatory surgical facilities, except that state inspections need not be performed during a year when a Medicare inspection is performed.  

Reimbursement
2011 Public Law 657 ended MaineCare reimbursement for services provided by ambulatory surgical facilities but that funding was restored beginning July 1, 2014.

(11) American Medical Association Code of Medical Ethics and Policy Finder
https://www.ama-assn.org/about/policyfinder
https://www.ama-assn.org/delivering-care/ama-code-medical-ethics

(12) Anatomical Gifts
Select Definitions
A. Agent: means an individual:
   1. Authorized to make health care decisions on the principal’s behalf by a power of attorney for health care; or
   Expressly authorized to make an anatomical gift on the principal’s behalf by any other record signed by the principal.
B. **Anatomical gift**: means a donation of all or part of a human body to take effect after the donor’s death for the purposes of transplantation, therapy, research or education.

C. **Disinterested witness**: means a witness other than the spouse, registered domestic partner, child, parent, sibling, grandchild, grandparent or guardian of the individual who makes, amends, revokes or refuses to make an anatomical gift, or another adult who exhibited special care and concern for the individual and who is familiar with the individual’s personal values. “Disinterested witness” does not include a person to which an anatomical gift could pass under section 2951.

D. **Document of gift**: means a donor card, advance directive or other record used to make an anatomical gift. “Document of gift” also means inclusion in a donor registry.

E. **Donor registry**: means the Maine Organ Donor Registry maintained under Title 29-A, section 1402-A as well as any other electronic database that identifies donors and complies with section 2958.

F. **Eye bank**: means a person that is licensed, accredited or regulated under federal or state law to engage in the recovery, screening, testing, processing, storage or distribution of human eyes or portions of human eyes.

G. **Organ procurement organization**: means a person designated by the United States Secretary of Health and Human Services as an organ procurement organization.

H. **Procurement organization**: means an eye bank, organ procurement organization or tissue bank.

I. **Prospective donor**: means an individual who is dead or near death and has been determined by a procurement organization to have a part that could be medically suitable for transplantation, therapy, research or education. “Prospective donor” does not include an individual who has made a refusal that is known by the procurement organization.

J. **Reasonably available**: means able to be contacted by a procurement organization without undue effort and willing and able to act in a timely manner consistent with existing medical criteria necessary for the making of an anatomical gift.

K. **Recovery agency**: means an eye bank, organ procurement organization, tissue bank, educational institution or research organization that participates in or facilitates the execution of an anatomical gift.

L. **Refusal**: means a record created under section 2947 that expressly states an intent to bar other persons from making an anatomical gift of an individual’s body or part.

M. **Tissue**: means a portion of the human body other than an organ or an eye. “Tissue” does not include blood unless the blood is donated for purposes of research or education.

N. **Tissue bank**: means a person that is licensed, accredited or regulated under federal or state law to engage in the recovery, screening, testing, processing, storage or distribution of tissue.

O. **Transplant hospital**: means a hospital that furnishes organ transplants and other medical and surgical specialty services required for the care of transplant patients.

**22 MRSA §2942**

**Applicability**
This chapter applies to an anatomical gift or amendment to, revocation of or refusal to make an anatomical gift, whenever made.

**22 MRSA §2943**

**Who may make anatomic gift before donor’s death**
Subject to section 2948, an anatomical gift of a donor’s body or part may be made during the life of the donor for the purpose of transplantation, therapy, research or education in the manner provided in section 2945 by:

**22 MRSA §2944**

**Donor**
The donor, if the donor is at least 18 years of age or is under 18 years of age and is:
A. An emancipated minor; or
B. Authorized under state law to apply for a driver’s license because the donor is at least 16 years of age;

**Agent of donor**

An agent of the donor, unless the power of attorney for health care or other record prohibits the agent from making an anatomical gift;

**Parent of the donor**

A parent of the donor, if the donor is under 18 years of age and not emancipated; or

**Guardian of donor**

The donor’s guardian.

**Use of medical marijuana**

In reviewing a qualifying patient's suitability for receiving an anatomical gift, a transplant evaluator shall treat the qualifying patient's medical use of marijuana as the equivalent of the authorized use of any other medications used at the direction of a medical provider. A transplant evaluator may determine a qualifying patient to be unsuitable to receive an anatomical gift if the qualifying patient does not limit the qualifying patient's medical use of marijuana to the use of forms of prepared marijuana that are not smoked or vaporized.

**Discrimination prohibited**

Insurers may not limit or refuse coverage or otherwise discriminate against a person on the basis of that person’s status as a living organ donor.

**Manner of making anatomical gift before donor’s death**

**Donor**

A donor may make an anatomical gift:

- A. By authorizing inclusion in the donor registry;
- B. In a will; or
- C. During a terminal illness or injury of the donor, by any form of communication addressed to at least 2 other individuals who are at least 18 years of age one of whom is a disinterested witness.

**Donor or other authorized person**

A donor or other person authorized to make an anatomical gift under section 2944 may make a gift by a donor card, advance directive or other record signed by the donor or other person making the gift authorizing inclusion in the donor registry. If the donor or other person is physically unable to sign the record, the record may be signed by another individual at the direction of the donor or the other person must:

- A. Be witnessed by at least 2 other individuals who are at least 18 years of age, one of whom is a disinterested witness, who have signed at the request of the donor or the other person; and
- B. State that it has been signed and witnessed as provided in paragraph A.

**Anatomical gift not invalidated**

Revocation, suspension, expiration or cancellation of the driver’s license or identification card issued to a donor does not invalidate an anatomical gift.
Anatomical gift by will
An anatomical gift made by will takes effect upon the donor’s death whether or not the will is probated. Invalidation of the will after the donor’s death does not invalidate the gift.  

22 MRSA §2945 (4)

Amending or revoking anatomical gift before donor’s death
Donor or authorized person
Subject to section 2948, a donor or other person authorized to make an anatomical gift under section 2944 may amend or revoke an anatomical gift by:

A. A record signed by:
   1. The donor;
   2. The other person;
   Subject to subsection 2, another individual acting at the direction of the donor or the other person if the donor or other person is physically unable to sign; or

B. A later-executed document of gift that amends or revokes a previous anatomical gift or portion of an anatomical gift either expressly or by inconsistency.  

22 MRSA §2946 (1)

Individual acting at donor’s or authorized person’s direction
A record signed pursuant to subsection 1, paragraph A, subparagraph (3) must:

A. Be witnessed by at least 2 other individuals who are at least 18 years of age, one of whom is a disinterested witness, who have signed at the request of the donor or the other person; and
B. State that it has been signed and witnessed as provided in paragraph A.  

22 MRSA §2946 (2)

Revocation by destruction or cancellation of document
Subject to section 2948, a donor or other person authorized to make an anatomical gift under section 2944 may revoke the gift by the destruction or cancellation of the document of gift, or a portion of the document of gift used to make the gift, with the intent to revoke the gift.  

22 MRSA §2946 (3)

Amendment or revocation by donor during terminal illness or injury
A donor may amend or revoke an anatomical gift that was not made in a will by any form of communication during a terminal illness or injury addressed to at least 2 other individuals who are at least 18 years of age, one of whom is a disinterested witness.  

22 MRSA §2946 (4)

Amendment or revocation of gift in will
A donor who makes an anatomical gift in a will may amend or revoke the gift in the manner provided for amendment or revocation of wills or as provided in subsection 1.  

22 MRSA §2946 (5)

Refusal to make anatomical gift and effect of refusal
Refusal of individual
An individual may refuse to make an anatomical gift of the individual’s body or part by:

A. A record signed by:
   1. The individual; or
   Subject to subsection 2, another individual acting at the direction of the individual if the individual is physically unable to sign;
B. The individual’s will whether or not the will is admitted to probate or invalidated after the individual’s death; or
C. Any form of communication made by the individual during the individual’s terminal illness or injury addressed to at least 2 other individuals who are at least 18 years of age, one of whom is a disinterested witness.  

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Individual acting at direction of individual
A record signed pursuant to subsection 1, paragraph A, subparagraph (2) must:
   A. Be witnessed by at least 2 other individuals who are at least 18 years of age, one of whom is a
      disinterested witness, who have signed at the request of the individual; and
   B. State that it has been signed and witnessed as provided in paragraph A.

Amendment or revocation by individual
An individual may amend or revoke a refusal:
   A. In the manner provided in subsection 1 for making a refusal;
   B. By subsequently making an anatomical gift pursuant to section 2945 that is inconsistent with the
      refusal; or
   C. By the destruction or cancellation of the record evidencing the refusal, or the portion of the record
      used to make the refusal, with the intent to revoke the refusal.

Effect of unrevoked refusal
Except as otherwise provided in section 2948, subsection 7, in the absence of an express, contrary
indication by the individual set forth in the refusal, an individual’s unrevoked refusal to make an
anatomical gift of the individual’s body or a part bars all other persons from making an anatomical gift of
the individual’s body or the part.

Preclusive effect of anatomical gift, amendment or revocation
Person other than donor barred
Except as otherwise provided in subsection 7, in the absence of an express, contrary indication by the
donor, a person other than the donor is barred from making, amending or revoking an anatomical gift of a
donor’s body or a part if the donor made an anatomical gift of the donor’s body or part under section 2945
or an amendment to an anatomical gift of the donor’s body or the part under section 2946.

Revocation not refusal
A donor’s revocation of an anatomical gift of the donor’s body or a part under section 2946 is not a
refusal and does not bar another person specified in section 2944 or 2949 from making an anatomical gift of
the donor’s body or a part under section 2945 or 2950.

Effect of gift or amendment by person other than donor
If a person other than the donor makes an unrevoked anatomical gift of the donor’s body or a part under
section 2945 or an amendment to an anatomical gift of the donor’s body or a part under section 2946,
another person may not make, amend or revoke the gift of the donor’s body or part under section 2950.

Effect of revocation by person other than donor
A revocation of an anatomical gift of the donor’s body or a part under section 2946 by a person other than
the donor does not bar another person from making an anatomical gift of the body or a part under section
2945 or 2950.

Effect of gift of a part or for a purpose
In the absence of an express, contrary indication by the donor or other person authorized to make an
anatomical gift under section 2944:
   A. An anatomical gift of a part is neither a refusal to give another part nor a limitation on the making
      of an anatomical gift of another part at a later time by the donor or another person under section
      2945 or 2950; and
B. An anatomical gift of a part for one or more of the purposes set forth in section 2944 is not a limitation on the making of an anatomical gift of the part for any of the other purposes by the donor or any other person under section 2945 or 2950.

**Donor unemancipated minor**

If a donor who is an unemancipated minor dies under 18 years of age, a parent of the donor who is reasonably available may revoke or amend an anatomical gift of the donor’s body or part.

**Parent’s revocation of unemancipated minor’s refusal**

If an unemancipated minor who signed a refusal dies under 18 years of age, a parent of the individual who is reasonably available may revoke the individual’s refusal.

**Who may make anatomic gift of decedent’s body or part**

**Gift by members of class; priority**
Subject to subsections 2 and 3 and unless barred by subsection 4, an anatomical gift of a decedent’s body or part for purposes of transplantation, therapy, research or education may be made, in the order of priority listed, by any member of the following classes of persons who is reasonably available:

- A. An agent of the decedent at the time of death who could have made an anatomical gift under section 2944, subsection 2 immediately before the decedent’s death;
- B. The spouse of the decedent;
- C. The registered domestic partner of the decedent;
- D. Adult children on the decedent;
- E. Parents of the decedent;
- F. Adult siblings of the decedent;
- G. Adult grandchildren of the decedent;
- H. Grandparents of the decedent;
- I. An adult who exhibited special care and concern for the decedent who is familiar with the decedent’s personal values;
- J. The person or persons acting as the guardian of the person of the decedent at the time of death; and
- K. Any other person having the authority to dispose of the decedent’s body.

**Anatomical gift by member of class unless object**

If there is more than one member of a class listed in subsection 1, paragraph A, D, E, F, G, H or J entitled to make an anatomical gift, an anatomical gift may be made by a member of the class unless that member or person to which the gift can pass under section 2951 knows of an objection by another member of the class. If an objection is known, the gift may be made only by a majority of the members of the class who are reasonably available.

**Member of prior class reasonably available**

No person may make an anatomical gift if, at the time of the decedent’s death, a person in a prior class under subsection 1 is reasonably available to make or to object to the making of an anatomical gift.

**Gift barred**

An anatomical gift may not be made if doing so is barred by section 2947 or 2948.
Manner of making, amending or revoking anatomical gift of decedent’s body or part

Authorized person: document; oral communication

A person authorized to make an anatomical gift under section 2949 may make an anatomical gift by a document of gift signed by the person making the gift or that person’s oral communication that is electronically recorded or is contemporaneously reduced to a record and signed by the individual receiving the oral communication.

Amendment or revocation by prior class member

Subject to subsection 3, an anatomical gift by a person authorized under section 2949 may be amended or revoked orally or in a record by any member of a prior class who is reasonably available. If more than one member of the prior class is reasonably available, the gift may be amended or revoked only if a majority of the reasonably available members object to the amending or revoking of the gift or they are equally divided as to whether to amend or revoke an anatomical gift.

Revocation effective if known

A revocation under subsection 2 is effective only if the procurement organization or transplant hospital or the physician or technician knows of the revocation before an incision has been made to remove a part from the donor’s body or before invasive procedures have begun to prepare the recipient.

Request consent

Consent for an anatomical gift by a recovery agency under section 2949 must be documented in writing or, if secured in a telephone conversation, in a suitable recording, must disclose in plain language the specific tissue, organ or body part being donated and the purpose for which the anatomical gift will be used and must comply in all respects with rules regarding consent requirements for anatomical gifting adopted by the department pursuant to subsection 5.

Rulemaking

The department, after consultation with the Office of the Attorney General, shall adopt rules to implement this section. The rules must provide specific requirements for all recovery agencies, require federally recognized recovery agencies to demonstrate compliance with applicable federal standards governing consent to anatomical gifts and require all other recovery agencies that do not operate under federal regulation to demonstrate adherence to the consent requirements of this section. Rules adopted pursuant to this subsection are routine technical rules as defined in Title 5, chapter 375, subchapter 2-A.

Persons that may receive anatomical gift: purpose of anatomical gift

Named recipient

An anatomical gift of a body or part may be made to the following persons:

A. A named hospital, accredited medical school, dental school, college, university or organ procurement organization or other appropriate person for research or education;
B. A named individual designated by the person making the anatomical gift if the individual is the recipient of the part; or, if the part for any reason cannot be transplanted into the individual, the part passes in accordance with subsection 6 in the absence of an express, contrary indication by the person making the anatomical gift; or
C. A named eye bank or tissue bank.
Named purpose
If an anatomical gift of one or more specific parts or of all parts is made in a document of gift that does not name a person described in subsection 1 but identifies the purpose for which an anatomical gift may be used, the following rules apply.

A. If the part is an eye and the gift is for the purpose of transplantation or therapy, the gift passes to the appropriate eye bank.
B. If the part is tissue and the gift is for the purpose of transplantation or therapy, the gift passes to the appropriate tissue bank.
C. If the part is an organ and the gift is for the purpose of transplantation or therapy, the gift passes to the appropriate organ procurement organization as custodian of the organ.
D. If the part is an organ, an eye or tissue and the gift is for the purpose of research or education, the gift passes to the appropriate procurement organization.

Priority of purposes
For the purpose of subsection 2, if there is more than one purpose of an anatomical gift set forth in the document of gift but the purposes are not set forth in any priority, the gift must be used for transplantation or therapy if suitable for those purposes and, if the gift cannot be used for transplantation or therapy, the gift may be used for research or education.

No named recipient or purpose
If the anatomical gift of one or more specific parts is made in a document of gift that does not name a person described in subsection 1 and does not identify the purpose of the gift, the gift passes in accordance with subsection 6 and the decedent’s parts must be used for transplantation or therapy, if suitable, and, if not suitable, the gift may be used for research or education.

General intent
If a document of gift specifies only a general intent to make an anatomical gift by words such as “donor,” “organ donor” or “body donor” or by a symbol or statement of similar import, the gift passes in accordance with subsection 6 and the decedent’s parts must be used for transplantation or therapy, if suitable, and, if not suitable, the gift may be used for research or education.

Rules of passing anatomical gifts
For purposes of subsection 1, paragraph B and subsections 3, 4 and 5, the following rules apply.

A. If the part is an eye, the gift passes to the appropriate eye bank.
B. If the part is tissue, the gift passes to the appropriate tissue bank.
C. If the part is an organ, the gift passes to the appropriate organ procurement organization as custodian of the organ.

Passing of organ for transplantation or therapy
An anatomical gift of an organ for transplantation or therapy, other than an anatomical gift under subsection 1, paragraph B, passes to the organ procurement organization as custodian of the organ.

Custody of body or part if not passed or used
If an anatomical gift does not pass pursuant to subsections 1 to 7 or the decedent’s body or part is not used for transplantation, therapy, research or education, custody of the body or part passes to the person under obligation to dispose of the body or part.

Acceptance of gift prohibited
A person may not accept an anatomical gift if the person knows that the gift was not effectively made under section 2945 or 2950 or if the person knows that the decedent made a refusal under section 2947 that was not revoked. For purposes of this subsection if a person knows that an anatomical gift was made
on a document of gift, the person is deemed to know of any amendment or revocation of the gift or any refusal to make an anatomical gift on the same document of gift.

**Allocation of organs for transplant or therapy**
Except as otherwise provided in subsection 1, paragraph B, nothing in this chapter affects the allocation of organs for transplantation or therapy.

For more on Anatomical Gifts see 22 MRSA Chapter 710-B.

### (13) Antitrust

**Statements of Antitrust Enforcement Policy in Health Care**

See [Federal Statements of Antitrust Enforcement Policy in Health Care](#)

**Health Care Practitioner Self-referral Act**

**Definitions**

A. **Bureau**: means the Bureau of Insurance.

B. **Facility**: means any sole proprietorship, partnership, firm, corporation or other business that provides health services.

C. **Group practice**: means a group of 2 or more health care practitioners legally organized as a partnership, professional corporation, nonprofit corporation or similar association in which:

   1. Each health care practitioner who is a member, an employee or an independent contractor of the group provides substantially the full range of services that the health care practitioner routinely provides, including consultation, diagnosis or treatment, through the use of office space, facilities, equipment or personnel of the group;
   
   The services of the health care practitioners are provided through the group and payments received for health services are treated as receipts of the group; and
   
   The overhead expenses and the income from the practice are distributed by methods previously determined by the group.

D. **Health care practitioner**: means an individual regulated under the laws of this State to provide health services. "Health care practitioners" include, without limitation, acupuncturists, chiropractors, dentists, dental hygienists, nurses, occupational therapists, optometrists, pharmacists, physical therapists, physicians including allopathic and osteopathic physicians, physician assistants, podiatrists, psychologists, clinical social workers, speech therapists and audiologists or hearing aid dealers and examiners.

E. **Health services**: means diagnosis, treatment and rehabilitative services for an injured, disabled or sick person.

F. **Immediate family member**: means a health care practitioner's parent, spouse, child or child's spouse.

G. **Investment interest**: means an equity or debt security issued by a facility, including, without limitation, shares of stock in a corporation, units or other interests in a partnership, bonds, debentures, notes or other equity interests or debt instruments, except that investment interest does not include interest in a hospital licensed under state law.

H. **Investor**: means an individual who owns, whose immediate family owns or who directly or indirectly owns a controlling interest in another facility that owns an investment interest in a facility that provides health services.

I. **Office practice**: includes the facility or facilities at which a health care practitioner, on a regular basis, provides or supervises the provision of professional health services to individuals.

J. **Referral**: means a referral of a patient for health services, including, without limitation:
1. The forwarding of a patient by one health care practitioner to another health care practitioner or a facility outside the health care practitioner's office practice or group practice that provides health services; or
2. The establishment by a health care practitioner of a plan of care outside the health care practitioner's office practice or group practice that includes the provision of any health services.

22 MRSA §2084

Prohibited referrals
A health care practitioner may refer a patient to an outside facility in which that health care practitioner is an investor only when that health care practitioner directly provides health services within the facility and will be personally involved with the provision of care to the referred patient.

22 MRSA §2085 (1)

Exemption
Referrals by a health care practitioner are exempt from this chapter if the bureau determines that there is demonstrated need in the community for the facility and alternative financing is not available. A health care practitioner does not have to demonstrate a need for alternative financing if the practitioner has sufficient financial resources in the provider's practice without seeking financing from outside sources other than conventional bank loans. Demonstrated need in the community for the facility exists when:

A. There is no facility of reasonable quality that provides an appropriate service, or the bureau determines that the quality of health care services would be improved in the community, such as by providing new specialty or subspecialty services without increasing overall health care costs and utilization above levels likely to occur if such an exemption were not granted;
B. Use of existing facilities is onerous or creates too great a hardship for patients;
C. The facility is formed to own or lease medical equipment that replaces obsolete or otherwise inadequate equipment in or under the control of a hospital located in a federally designated health manpower shortage area; or
D. The facility meets other standards established by rule by the bureau, including a standard allowing the bureau to determine whether the fees charged for the health services are competitive with fees charged for those services outside the community. "Community" must be defined by rule by the bureau. The following requirements must be met to be exempt under this section.

1. Individuals who are not in a position to refer patients to a facility must be given a bona fide opportunity to invest in that facility on the same terms as those offered a referring health care practitioner.
2. A health care practitioner who invests may not be required or encouraged to make referrals to the facility or otherwise generate business as a condition of becoming or remaining an investor.
3. The facility shall market or furnish its services to investors who are referring health care practitioners and to other investors on equal terms.
4. The facility may not loan funds or guarantee loans for health care practitioners who are in a position to refer patients to that facility.
5. The income on the health care practitioner's investment must be tied to the health care practitioner's equity in the facility rather than to the volume of referrals made.
6. An investment contract between the facility and the health care practitioner may not include a covenant or noncompetition clause that prevents a health care practitioner from investing in other facilities.
7. When making a referral, a health care practitioner shall disclose to the patient being referred to the facility that health care practitioner's investment interest in that facility. If alternative facilities are reasonably available, the health care practitioner shall provide the patient with a list of alternative facilities. The health care practitioner shall inform the patient that the patient has the option to use an alternative facility and the patient will not be treated differently by the health care practitioner if the patient chooses to use another
facility. This subparagraph applies to all investors who are health care practitioners, including those who provide direct care or services for their patients in facilities outside their office practice.

8. If a 3rd-party payor requests information regarding a health care practitioner's investment interest, that information must be disclosed.

9. The facility shall establish an internal utilization review program.

10. If a health care practitioner's financial interest in a facility is incompatible with a referred patient's interest, the health care practitioner shall make alternative arrangements for that patient's care.

The bureau shall make its determination on a request for an exemption within 90 days of a completed written request.

**22 MRSA §2085 (2)**

**Exception**

It is not a violation of this chapter for a health care practitioner to refer a patient to a publicly traded facility in which that health care practitioner has an investment interest when:

A. The facility is listed for trading on the New York Stock Exchange or on the American Stock Exchange or is a national market system security traded under an automated interdealer quotation system operated by the National Association of Securities Dealers;

B. The facility, at the end of its most recent fiscal year, had total net assets of at least $50,000,000 related to the furnishing of health services;

C. Investment interest obtained after the effective date of this chapter is traded on the exchanges listed in paragraph A;

D. The facility markets or furnishes its services to investors who are referring health care practitioners and to other health care practitioners on equal terms;

E. All stock held in that facility, including stock held in the predecessor privately held facility, is of one class without preferential treatment as to status or remuneration;

F. The facility does not loan funds or guarantee loans for health care practitioners who are in a position to make referrals to a facility;

G. The income on the health care practitioner's investment is tied to the health care practitioner's equity in the facility rather than to the volume of referrals made; and

H. The investment interest does not exceed 1/2 of 1% of the facility's total equity.

**22 MRSA §2085 (3)**

**Compelling practitioner**

A health care practitioner may not compel or coerce, or attempt to compel or coerce, any other health care practitioner to violate any provision of this chapter.

**22 MRSA §2085 (4)**

**Third-party referrals**

A health care practitioner may not participate in any arrangement or plan that is designed to evade the prohibitions in this chapter by using a 3rd party to redirect referrals that are prohibited under subsection 1 if the 3rd party was not involved in the referral.

**22 MRSA §2085 (5)**

**Alternate facilities**

If compliance with the community need and alternative financing criteria is not practical, the health care practitioner shall identify to the patient reasonably available alternative facilities. The bureau, by rule, shall designate when compliance is not practical.

**22 MRSA §2085 (6)**

**Bureau opinion**

Health care practitioners may request that the bureau render an advisory opinion as to whether a referral to an existing or proposed facility under specified circumstances violates the provision of this chapter. The bureau's opinion is presumptively correct as to whether the provisions of this chapter are violated.

**22 MRSA §2085 (7)**
Health organizations
Notwithstanding any provision of this chapter, a health care practitioner may refer a patient who is a member of a health maintenance organization or a preferred provider organization licensed in this State for health services to a facility outside that health care practitioner's office or group practice in which that health care practitioner is an investor when the referral is made pursuant to a contract with the organization.

22 MRSA §2085 (8)

Penalties
A facility or a health care practitioner that makes or causes to be made a referral prohibited under section 2085 or presents or causes to be presented a bill or claim for service that the facility or health care practitioner knows or should know is prohibited by section 2085 is subject to a civil penalty of no more than $2,000 for each referral, bill or claim.

A violation of this chapter by a health care practitioner or a facility constitutes grounds for disciplinary action by the applicable licensing body.

22 MRSA §2086

(14) Audits Conducted by DHHS; Guidelines and Criteria
Auditing and adjusting of health and community service provider costs
This section governs the rules of the department and the practices of its auditors in interpreting and applying those rules with respect to payments of providers under the MaineCare program and payments by the department under grants and agreements audited pursuant to the Maine Uniform Accounting and Auditing Practices Act for Community Agencies.

Revised audit interpretations to be applied prospectively
Whenever the department's auditors revise an interpretation of a rule, agreement, circular or guideline in a manner that results in a negative adjustment of a provider's or agency's allowable costs, the revised interpretation may be applied only to provider or agency fiscal years beginning after the date of the examination report, audit report, or other written notification in which the provider or agency receives direct notice of the revised interpretation. For the fiscal year to which the report containing the revised interpretation applies, and any subsequent fiscal year ending prior to the issuance of the revised interpretation, the cost that is the subject of the revised interpretation must be considered allowable to the extent that it was allowable under the interpretation previously applied by the Office of Audit for MaineCare and Social Services, referred to in this section as "the office of audit." This subsection does not prohibit the office of audit from applying an adjustment to a fiscal year solely because that cost was not disallowed in a prior year.

22 MRSA §41-B (1)

Determination of "ordinary," "necessary" or "reasonable" costs
In making findings concerning whether costs are "ordinary," "necessary" or "reasonable," the office of audit shall consider the following criteria in conjunction with applicable state and federal rules, regulations, guidelines and agreements:

A. Whether a substantial number of providers of health care or community services in Maine incur costs of similar magnitude, frequency, quantity or price level to the costs under review.
B. Whether the expenditure is reasonably incurred to produce, accomplish, facilitate or compensate persons for providing an item or service related to the purpose of a program or activity for which the State has contracted or for which the State otherwise provides payment.
C. Whether the expenditure is comparable to one incurred by a department or agency of the State responsible for services or programs similar to those to which the finding applies.
D. Whether the expenditure is consistent with meeting special needs of the population served through innovative or specialized services offered by a particular provider.
Employee compensation and benefit costs
In evaluating the reasonableness and allowability of employee wages, salaries and benefits, the department may not disallow the costs of any employee benefits, wages or salaries if the total of those costs is reasonable under the criteria set forth in subsection 2.

Other expenses
The department shall modify its rules governing MaineCare reimbursement and other reimbursements pursuant to grants, contracts or agreements for health care providers and other agencies providing community services to allow, to the extent permitted by applicable federal law, the costs of employee information publications, health or first-aid clinics or infirmaries, recreational activities, employee counseling services and any other expenses incurred in accordance with the health care provider or other agency's established practice or custom for the improvement of working conditions, employer-employee relations, employee morale and employee performance.

Special Notes
The department has made available the decisions in all MaineCare provider appeals, including the recommendations of the hearing officer and the decision of the Commissioner, at the following public website: http://www.maine.gov/dhhs/hearings/appeals.htm.

MaineCare program integrity recovery audit contractor agreement
As of 2012, MaineCare has been authorized to enter into an agreement with a recovery audit contractor for the purpose of ensuring MaineCare program integrity, specifically to identify and reimburse to correct underpayments and to identify and recoup overpayments under the Medicaid state plan and under any waiver of the state plan. An agreement entered into under this section must provide that payment to the contractor may be made only from amounts recovered and that payments for identifying underpayments and collecting overpayments must be made on a contingent fee basis. After payments to correct underpayments and payment of any contingent fees due to the contractor, the proceeds of collections from overpayments must be deposited into the Medical Care - Payments to Providers program, Other Special Revenue Funds account in the Department of Health and Human Services for the purpose of providing state match under the federal Medicaid program.

Credible allegations of fraud; provider payment suspensions
If the department determines that there is a credible allegation of fraud by a provider under the MaineCare program, the following procedures apply.

Suspension of payments
The department shall suspend payment in whole or in part to a MaineCare provider when a suspension is necessary to comply with Section 6402(h)(2) of the federal Patient Protection and Affordable Care Act of 2010, Public Law 111-148 and 42 Code of Federal Regulations, Part 455.

Administrative appeal
A MaineCare provider may administratively appeal the department's decision to suspend payment under subsection 1.

No stay during administrative appeal
A suspension of payments under subsection 1 may not be stayed during an administrative appeal of the department's decision to suspend payment. The department may provide a fair opportunity for appropriate expedited relief from a suspension of payments consistent with federal law.
Final determination; offset
Upon a final determination that fraud has occurred and that money is owed by the MaineCare provider to the department, and 31 days after exhaustion of all administrative appeals and any judicial review available under Title 5, chapter 375, the department may retain and apply as an offset to amounts determined to be owed to the department any payments to the provider that were suspended by the department pursuant to this section. The amount retained pursuant to this subsection may not exceed the amount determined finally to be owed.

22 MRSA § 1714-E (4)

Confidentiality
Except as necessary for purposes of the investigation of fraud or the administration of the MaineCare program, the department's records regarding a determination of a credible allegation of fraud are confidential until the relevant MaineCare provider has been given notice of a suspension of payments under subsection 1.

22 MRSA § 1714-E (5)

Rules
The department shall adopt rules to implement this section, including rules to define "credible allegation of fraud" and to provide exception and appeal procedures as required by and in accordance with the requirements of federal law and regulations. If the department provides a procedure for expedited relief from suspension of payments, as authorized in subsection 3, the rules must include that procedure. Rules adopted pursuant to this subsection are routine technical rules as defined in Title 5, chapter 375, subchapter 2-A.

22 MRSA § 1714-E (6)

Repeal
This section is repealed if Section 6402(h)(2) of the federal Patient Protection and Affordable Care Act, Public Law 111-148 and 42 Code of Federal Regulations, Part 455 are invalidated by the United States Supreme Court. The department shall notify the Secretary of State, the Secretary of the Senate, the Clerk of the House of Representatives and the Revisor of Statutes if the section of law and the regulation are invalidated.

22 MRSA § 1714-E (7)

(15) Automated External Defibrillators (AED)
Definitions
A. “Automated external defibrillator” or "AED" means a medical device that combines a heart monitor and a defibrillator approved by the United States Food and Drug Administration that:
   1. Is capable of recognizing the presence or absence of ventricular fibrillation or rapid ventricular tachycardia;
   2. Is capable of determining, without intervention by an operator, whether defibrillation should be performed on an individual;
   3. Upon determination that defibrillation should be performed, automatically charges and requests delivery of an electrical impulse to an individual's heart; and

22 MRSA §2150-C (1)

Immunity
The following persons and entities are immune from civil liability for damages relating to the use, possession or purchase of an AED and arising out of acts or omissions relating to preparing for and responding to suspected sudden cardiac arrest emergencies absent gross negligence or willful or wanton misconduct:

A. Any person or entity that acquires an AED.
B. Any person or entity that owns, manages or is otherwise responsible for the premises on which an AED is located;
C. Any person who retrieves an AED in response to a perceived sudden cardiac arrest emergency;
D. Any person who uses, attempts to use or fails to use an AED in response to a perceived sudden cardiac arrest emergency;
E. Any physician or other authorized person who issues a prescription for the purchase of an AED;
F. Any person or entity that is involved with the design, management or operation of an AED program; and
G. Any person or entity that provides instruction in the use of an AED.

22 MRSA §2150-C (6)

(16) Autopsies
No embalming when autopsy authorized
The next of kin or legal representative of a person who has died may authorize an autopsy. If an autopsy is authorized, no person shall inject into or remove from any artery, vein, or cavity of the body of the person who has died any fluid, gas or other substance except by or with the permission of a pathologist, medical examiner or licensed physician in attendance.

On completion of the autopsy, the body shall be released for normal handling.

The provisions of this section do not apply to deaths within the jurisdiction of medical examiners or autopsies as authorized in Title 22, chapter 711.

A violation of this section is a Class E crime.

32 MRSA §1404-A

(17) Births
Registration of live births
A certificate of each live birth that occurs in this State must be filed with the clerk of the municipality in which the live birth occurred or with the state registrar within a reasonable period of time as specified by the department and must be registered if the certificate has been completed and filed in accordance with this section.

22 MRSA §2761

Certificate from hospital
When the live birth occurs in a hospital or an institution, or en route to the hospital or institution, the person in charge of the institution or the person's authorized designee shall obtain the personal data, prepare the certificate, certify by signature or by electronic process that the child was born alive at the place and time and on the date stated and file the certificate as directed in this section. The physician or other person in attendance shall provide the medical information required by the certificate in a timely fashion, in accordance with department rule.

22 MRSA §2761 (1)

Birth outside an institution
When a birth occurs outside an institution, the certificate must be prepared and filed by one of the following in the indicated order of priority:
A. The physician or other person in attendance at or immediately after the birth;
B. The father;
C. The mother; or
D. The person in charge of the premises where the live birth occurred.

22 MRSA §2761 (3)
Parentage
For the purposes of birth registration, the mother is deemed to be the woman who gives birth to the child, unless otherwise determined by a court of competent jurisdiction prior to the filing of the birth certificate. If the mother was married at the time of either conception or birth, or between conception and birth, the name of the husband must be entered on the certificate as the father of the child, unless paternity has been determined otherwise by a court of competent jurisdiction.

22 MRSA §2761 (3-A)

For additional information regarding the content and processing of a birth certificate see 22 MRSA §2761.

Hospital-based paternity acknowledgement

Birth center
As used in this section, "birthing center" means a hospital or other facility that provides childbirth services.

22 MRSA §2761-B (1)

Procedure
A birthing center shall provide an opportunity for all unmarried parents to complete a voluntary acknowledgement of paternity. A birthing center shall provide to each unmarried mother and alleged father, if present, written information about paternity establishment provided by the department, forms needed to voluntarily acknowledge paternity and the opportunity to speak with a person who is trained to clarify information and answer questions about paternity establishment. The birthing center shall forward all completed acknowledgement forms to the department.

22 MRSA §2761-B (2)

Birth certificates of foundlings; report

Whoever assumes the custody of a child of unknown parentage shall immediately report to the local town or city clerk in writing:

A. Date and place of finding. The date and place of finding or assumption of custody;
B. Sex, color, age. Sex; color or race; and approximate age of child;
C. Name and address of custodian. Name and address of the person or institution with whom the child has been placed for care;
D. Name. Name given to the child by the finder or custodian.

The place where the child was found or custody assumed shall be known as the place of birth and the date of birth shall be determined by approximation. The report shall constitute the certificate of birth. If the child is thereafter identified, the record of birth made in compliance herewith and any certificate issued thereon shall be null and void and so recorded.

22 MRSA §2763

Delayed birth registration
For information on delayed birth registration see 22 MRSA §2764.

Care of infants after birth
Every physician, midwife or nurse in charge shall instill or cause to be instilled into the eyes of an infant upon its birth one or 2 drops of a prophylactic solution prescribed by the department and provided without cost by the department, except an infant whose parents object to this procedure on the grounds that it conflicts with their religious tenets and practices. If one or both eyes of an infant become reddened or inflamed at any time within 4 weeks after birth, the midwife, nurse or person having charge of the infant shall report the condition of the eyes at once to a physician licensed under Title 32, chapter 36 or 48. Failure to comply with this section shall be punishable by a fine of not more than $100 or by imprisonment for not more than 6 months.
Detection of serious conditions
The department shall require hospitals, birthing centers and other birthing services to test newborn infants, or to cause them to be tested, by means of blood spot screening for the presence of treatable congenital, genetic or metabolic conditions that may be expected to result in subsequent cognitive disabilities, serious illness or death. Reports and records of those making these tests may be required to be submitted to the department in accordance with departmental rules. The department may, on request, offer consultation, training and evaluation services to those testing facilities.

22 MRSA §1531

Referrals
The department shall in a timely fashion refer newborn infants with confirmed treatable congenital, genetic or metabolic conditions or critical congenital heart disease to the Child Development Services System as defined in Title 20-A, section 7001, subsection 1-A. For additional information on referrals see 22 MRSA §1532 (2).

22 MRSA §1532 (1)

Religious objection exemption
The requirement in this section that a newborn infant be tested for the presence of treatable congenital, genetic or metabolic conditions that may be expected to result in subsequent cognitive disability does not apply to a child if the parents of that child object on the grounds that the test conflicts with their religious tenets and practices.

22 MRSA §1532 (2)

Report
A hospital, birthing center or other birthing service that tests a newborn infant pursuant to this section shall report to the department aggregate data on the testing, including but not limited to the number of infants born, the number tested for treatable congenital, genetic or metabolic conditions, the number screened for critical congenital heart disease, the results of the screening and testing and, for heart disease screening the type of screening tool used.

Screening for heart conditions
All Maine hospitals, birthing centers and other birthing services must test newborn infants by means of appropriate technology for the presence of critical congenital heart disease; rulemaking may require reporting to DHHS

22 MRSA § 1532

Newborn Hearing Program, Program requirements
Definitions
A. Birth admission: means the time after birth that the newborn remains in the hospital nursery prior to discharge.
B. Board: means the Newborn Hearing Screening Advisory Board.
C. Hearing loss: means a hearing loss of 30 decibels or more in the frequency region important for speech recognition and comprehension in one or both ears. The department may adopt rules to decrease the amount of decibels of hearing loss as technology allows for detection of hearing loss of 15 to 25 decibels in one or both ears.
D. Intervention or treatment: means the early intervention services described in the federal Individuals with Disabilities Education Act, 20 United States Code, Chapter 33, Subchapter III, Sections 1431 to 1445, as amended. "Intervention" or "treatment" includes, but is not limited to, audiological, medical or early educational services that provide a choice of methods of communication in a variety of sensory modalities.
E. Parent: means a natural parent, stepparent, adoptive parent, legal guardian or other legal custodian of a child.
F. **Person who is culturally deaf:** means a person with permanent hearing loss who identifies as a member of the deaf community and who utilizes American Sign Language as the primary mode of communication.

G. **Person who is hard-of-hearing or person who is deaf:** means a person with permanent hearing loss who communicates using aural or oral skills for accessing spoken language.

### Information to parents of children born in hospitals

A hospital shall provide information to the parents of children born in the hospital regarding the importance of screening the hearing of newborns and of receiving follow-up care. The information must explain the process of hearing screening, the likelihood of a child having a hearing loss, follow-up procedures and community resources and must include a description of the normal auditory, speech and language development process in children. The hospital must provide information about hearing screening that may be provided at the hospital or coordinated, scheduled or arranged for by the hospital. The program must provide this information prior to discharge from the birth admission to the hospital or within 3 months of discharge.

### Information to parents of children born outside of hospitals

When a newborn is delivered in a facility other than a hospital, the department shall provide information to the parents on the merits of having the hearing screening performed and on the availability of the hearing screening within 3 months of the date of birth.

### Guidelines for services for children with hearing loss and at-risk children

The department, after consultation with the board, shall establish guidelines for the provision of follow-up services for newborn children in the State who are identified as having or being at risk of developing hearing loss. These services must include, but are not limited to, diagnostic audiologic assessment, counseling and educational services for the parents and an explanation of the potential effects of the identified hearing loss on the development of the newborn's speech, language and cognitive skills as well as the potential benefits of early identification and use of spoken or sign language.

For information on reporting, see **22 MRSA §8822**. For information on tracking, see **22 MRSA §8824**.

### Burns (Arson Reporting Immunity Act)

#### Reporting by health care practitioner

**Reasonable cause to suspect; information disclosed**

A health care practitioner, as defined by Title 24, section 2502, subsection 1-A, who, as a result of the practitioner's examination or treatment of a person for a burn injury, has reasonable cause to suspect that the burn injury was sustained in connection with an act of arson, may report to the Office of the State Fire Marshal. The health care practitioner's report may include the name and address of the person examined or treated, the basis for the practitioner's suspicion and other information which, in the judgment of the practitioner, may aid in investigation by the Office of the State Fire Marshal.

**Immunity**

A health care practitioner who, acting in good faith in reporting under this section or participating in a related investigation or proceeding, makes a report pursuant to subsection 1 is immune from civil or criminal liability for the act of reporting or participating in a related investigation or proceeding. Good faith does not include instances when a false report is made and the person knows the report is false. Nothing in this section may be construed to bar criminal or civil action regarding perjury.
Presumption of good faith
In a proceeding regarding immunity from liability, there shall be a rebuttable presumption that a report made under subsection 1 was made in good faith.

Privileged or confidential communications
The physician-patient privilege under the Maine Rules of Evidence is abrogated in relation to a report authorized under subsection 1.

(19) Business Arrangements and Structures

General
There are many types of arrangements and structures available to people wishing to set up a business in Maine. Each type has its own advantages and disadvantages. The basic information below will help you understand the various types of businesses available in Maine. For more information visit the following websites:

Secretary of State, Division of Corporations Website (General information)
http://www.maine.gov/sos/cec/corp/
Office of Professional and Occupational Regulation (Licensing and Registration)
http://www.maine.gov/pfr/professionallicensing/
Maine Revenue Services
http://www.state.me.us/revenue/
US Small Business Administration (SBA)
http://www.sba.gov/
Maine Small Business Development Centers (SBA)
http://www.mainesbdc.org/
How to Organize Your Business (University of Maine Cooperative Extension Bulletin #3009)
http://extension.umaine.edu/publications/3009e/

For legal advice and/or help setting up any of these types of businesses please consult with an attorney.

Sole Proprietorship
As the name indicates, this type of business is owned and operated by one person. This person is responsible for all business aspects, including all liability and any profit or loss. This structure is easy to set up and does not require any special registrations (other than trade names, trademarks, etc.).

Partnerships
A partnership is an association of two or more people to carry on a business for profit as co-owners. This type of business allows the partners to share ownership, management and assess profits/losses to each partner. This structure is easy to set up and does not require any special registrations (other than trade names, trademarks, etc.). No written agreement is needed under law to form a partnership, though one is beneficial in avoiding misunderstandings and helps clarify the business arrangements. This type of arrangement is also known as a General Partnership.

Limited Partnerships
A limited partnership is business arrangement between one or more general partners (who manage the business and are personally liable for partnership debts) and one or more limited partners (who contribute capital, share in profits, but do not run the business and are not liable for the partnership obligations beyond contribution).
Full text of Limited Partnership Statute  
http://janus.state.me.us/legis/statutes/31/title31ch11sec0.html  
Forms and Fees  

**Limited Liability Partnership (LLP)**  
A LLP is a general partnership that has elected to limit personal liability for its general partners by registering this election with the Secretary of State.

Full text of Limited Liability Partnership Statute  
http://janus.state.me.us/legis/statutes/31/title31ch15sec0.html  
Forms and Fees  

**Corporation**  
A corporation is formed for the purpose of transacting business in the broadest sense of the word and for a profit return. This type of business requires special forms (articles of incorporations for example) to be filed with the Secretary of State.

Full text of the Maine Business Corporation Act  
http://www.mainelegislature.org/legis/statutes/13-C/title13-Cch0sec0.html  
Forms and Fees  

**Professional Service Corporation**  
A professional service corporation is a type of corporation specifically for individuals licensed to provide certain services to the public, such as accounting, law, or medicine.

Full text of the Maine Professional Service Corporation Act  
http://www.mainelegislature.org/legis/statutes/13/title13ch22-Asec0.html  
Forms and fees  

**Limited Liability Company (LLC)**  
An LLC is a cross between a traditional corporation and a partnership. This type of organization provides a flexible business structure that has the ability to limit personal liability along with the ability to assess profits and losses to individuals.

Full text of Limited Liability Company Statute  
http://janus.state.me.us/legis/statutes/31/title31ch13sec0.html  
Forms and Fees  

**“S” Corporations**  
An “S” corporation is a type of business that allows a business to limit liability while also allowing the tax burden to shift directly to the shareholders. This special type of structure is typically seen in small businesses that wish the advantages of a typical corporation with the tax benefits of a partnership (i.e. no double taxation). This structure has some special incorporation requirements in order to qualify:  
A. Must be a domestic US company  
B. Can only have one type of stock
C. Cannot have more than 35 shareholders
D. Shareholders cannot be other business
E. All shareholders must be individuals or estates
F. All shareholders must be US residents

**Non-Profit Corporation**
A nonprofit corporation is formed for the purpose of advancing a particular objective of an organization and is not established to make a profit. Generally, this includes charitable, benevolent and educational organizations. These types of organizations receive a special tax status and benefits under the IRS Tax Code Section 501 (c)(3).

As of January 2003, all nonprofit corporations in the state of Maine are either "public benefit" or "mutual benefit" corporations.

**A Public Benefit Corporation:**
- A. Is designated as a public benefit corporation by statute; or
- B. Is tax exempt under section 501(c)(3) of the Internal Revenue Code; or
- C. Is organized for a public or charitable purpose and is required to distribute assets to a similar tax exempt organization upon dissolution; or
- D. Has elected to be a public benefit corporation.

Generally, if your corporation’s mission is to serve the public or community at large, it is probably a public benefit corporation. Public benefit corporations can usually be characterized as charities.

**A Mutual Benefit Corporation:**
- A. Is any nonprofit corporation that does not fit the description of a public benefit corporation. Generally, if your corporation’s mission is to serve a limited number of members with common interests, it is probably a mutual benefit corporation.

As of January 2003, charity corporations have new filing and reporting requirements. For more information about these requirements, responsibilities of the Board of Directors and other necessary links, please visit:

**Guide for Board Members of Charitable Corporations:**

**Full text for Maine Non-Profit Corporation Act**
[http://www.mainelegislature.org/legis/statutes/13-B/title13-Bch0sec0.html](http://www.mainelegislature.org/legis/statutes/13-B/title13-Bch0sec0.html)

**Forms and Fees**

**Trade Names and Marks**
- A. **Trade Name**
  Is a word, name, symbol, device or any combination thereof used by a person to identify his business, vocation or occupation and distinguish it from the business, vocation or occupation of others.
- B. **Trademark**
  Is any word, name, symbol or device or any combination thereof adopted and used by a person to identify goods made or sold by him and to distinguish them from goods made or sold by others.
- C. **Service marks**
  Is a trademark used in the sale or advertising of services to identify the services of one person and distinguish them from the services of others.
- D. **Certification marks**
Is a mark used upon or in connection with the products or services of one or more persons other than the owner of the mark to certify regional or other origin, material, mode of manufacture, quality, accuracy or other characteristics of such goods or services, or that the work or labor on the goods or services was performed by members of a union or other organization.

E. Collective marks
Is a trademark or service mark used by the members of a cooperative, an association or other collective group or organization, and includes marks used to indicate membership in a union, an association or other organization.

These marks are entitled to be registered with the Secretary of State or with the United States Patent and Trademark Office.

Useful Links
Full Text of Trade Marks and Names Statutes: [10 MRSA Chapter 301]
Maine State Trademark page
Forms and Fees

U.S. Patent & Trademark Office

Direct Primary Care
A direct primary care service agreement is an agreement between the direct primary care provider and either an individual or the individual's representative, regardless of whether the periodic fee or other fees are paid by the individual, the individual's representative or a 3rd party.

Direct primary care service agreements are not insurance and thus not subject to regulation by the Bureau of Insurance. The written agreement must so state. Providers who offer direct primary care are not prohibited from entering into agreements with insurers to supplement the DPC agreement. [22 MRSA §1771]

(20) Cancer
Duty of physicians and hospitals
All hospitals and other health care facilities providing screening, diagnostic or therapeutic services with respect to cancer shall report to the Department of Human Services all persons diagnosed as having a malignant tumor or certain benign tumors as determined by rule no later than 6 months from the date of diagnosis. The report must include information on the person's usual occupation and industry of employment and other elements determined by rule to be appropriate. The Commissioner of Human Services shall adopt rules to implement this section. Rules adopted pursuant to this section are routine technical rules as defined in Title 5, chapter 375, subchapter 2-A.

A physician, surgeon or other health care practitioner who diagnoses or provides treatment for cancer patients, upon notification by the Department of Human Services, shall report to the department any further information requested by the department concerning any person now or formerly under the health care practitioner's care, diagnosed as having or having had a malignant tumor. A physician, surgeon or other health care practitioner who diagnoses or provides treatment for cancer patients is required to report any newly diagnosed cancer case to the department when that patient will not be referred to a reporting facility for diagnosis or treatment.

A facility or individual complying with the reporting requirements of this section is not liable for any civil damages as a result of such acts.
The requirements of this section do not apply to health care practitioners who provide treatment by spiritual means alone.  

**Cancer-incidence registry and regulations (see also Agency Rules Link)**
The Department of Human Services shall establish, maintain and operate a statewide cancer-incidence registry.

(21) **Certificate of Need**
Maine’s Certificate of Need Act provides the framework for review of proposals by or on behalf of certain health care facilities and nursing homes involving expansion of plant and equipment, the provision of new services, transfers of ownership and control and other initiatives.

For the full Maine Certificate of Need Act, including definitions, exceptions and information on the application process, see: [http://www.mainelegislature.org/legis/statutes/22/title22ch103-Asec0.html](http://www.mainelegislature.org/legis/statutes/22/title22ch103-Asec0.html)

For more information, see the Department of Health & Human Services Certificate of Need Unit: [http://www.maine.gov/dhhs/dlrs/c_o_n/](http://www.maine.gov/dhhs/dlrs/c_o_n/)

A certificate of need is generally required for:

A. Any transfer of ownership or acquisition under lease or comparable arrangement or through donation or any acquisition of control of a health care facility under lease, management agreement or comparable arrangement or through donation that would have required review if the transfer or acquisition had been by purchase, except in emergencies when that acquisition of control is at the direction of the department or except if the transfer of ownership or acquisition of control involves only entities or health care facilities that are direct or indirect subsidiaries of the same parent corporation, is between a parent corporation and its direct or indirect subsidiaries or is between entities or health care facilities all under direct or indirect ownership of or ultimate control by the same parent corporation immediately prior to the transfer or acquisition;

B. Acquisitions of major medical equipment EXCEPT for
   1. Major medical equipment being replaced by the owner; or
   2. The use of major medical equipment on a temporary basis in the case of a natural disaster, major accident or major medical equipment failure.

C. Capital expenditures. The obligation by or on behalf of a health care facility of any capital expenditure of $10,000,000 or more (EFFECTIVE 2/15/12)

D. The offering or development of any new health service.

E. The construction, development or other establishment of a new health care facility:
   1. If it requires a capital expenditure of more than $3,000,000 or i
   2. f it is a new health service. (EFFECTIVE 2/15/12).

F. An increase in the existing licensed bed complement or an increase in the licensed bed category of a health care facility, other than a nursing facility, of greater than 10%.

G. The obligation by a new or existing nursing facility, when related to nursing services provided by the nursing facility, of any capital expenditures of $5,000,000 or more EXCEPT for
   1. A nursing facility converting beds used for the provision of nursing services to beds to be used for the provision of residential care services;
   2. Capital expenditures in the case of a natural disaster, major accident or equipment failure;
   3. C. Replacement equipment, other than major medical equipment as defined in section 328, subsection 16;
   4. D. Information systems, communication systems, parking lots and garages; and
   5. E. Certain energy-efficient improvements, as described in section 334-A, subsection 4.

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(22) **Chiropractic Licensure (see also Agency Rules Links)**

**Definitions**

**Chiropractic**
Chiropractic means the art and science of identification and correction of subluxation and the accompanying physiological or mechanical abnormalities. The term subluxation, as utilized within the chiropractic health care system, means a structural or functional impairment of an intact articular unit. "Chiropractic" includes chiropractic acupuncture. Chiropractic recognizes the inherent recuperative capability of the human body as it relates to the spinal column, musculo-skeletal and nervous system.

**Chiropractic acupuncture**
Chiropractic acupuncture means the insertion of acupuncture needles through the skin at specific points. Chiropractic acupuncture is a methodology used for the correction of the soft tissue components contributing to subluxation and the accompanying physiological or mechanical abnormalities. Except as provided in section 502, chiropractic acupuncture may only be practiced by a licensee who has received a chiropractic acupuncture certification from the board.

**Chiropractic doctors**
Chiropractic doctors are health care providers functioning within their scope of practice as provided by this chapter.

**Chiropractic methodologies**
Chiropractic methodologies utilized for the identification or correction of subluxation and the accompanying physiological or mechanical abnormalities include diagnostic, therapeutic, adjustive or manipulative techniques utilized within the chiropractic profession, excluding prescriptive medication or surgery.

**Licensure and Board of Chiropractic Licensure**
The chiropractic practice is governed by [32 MRSA, Chapter 9](#) and [Board rules](#).
For examination and licensure requirements, see [32 MRSA § 551-554](#).
For information about the Board of Chiropractic Licensure and their processes, see [32 MRSA § 501-503-B](#).

(23) **Closing a Medical Practice**
Please visit the Maine Medical Association website, [www.mainemed.com](http://www.mainemed.com) or contact the Maine Medical Association for resources on closing a medical practice. The MMA has published a [Physician’s Guide to Closing a Practice](#) that includes useful checklists and resources.

(24) **Communicable (Notifiable) Diseases (see also Agency Rules Links)**

**Rules for the Control of Notification**

Who Must Report and what is required in the report

A. Health care providers/Medical laboratories/Health care facilities/Child care facilities/Educational institutions/Correctional facilities/Health Officers

1. Disease (recognition, strong suspicion, death or positive diagnostic laboratory findings);
2. Date of first onset of symptoms;
3. Patient:
   a. Name
   b. Birth date
   c. Race
d. Ethnicity
e. Sex
f. Occupation (if known)
g. Residence address
h. Phone number
i. Place of work, school or child care
j. Parent or guardian name and address
k. Parent/Guardian telephone number

4. Date of report;
5. Health care provider name, address, and phone number;
6. Name of hospital or other healthcare facility (if any); 
7. Name of person reporting (if not health care provider);
8. All diagnostic laboratory findings and dates of test relevant to the notifiable condition regardless of clinical significance;
9. Name and locating information of contacts;
10. Other information pertinent to the case, as requested by the Department.
11. If animal species- specify.
12. When the report is about a non-compliant person or a public health threat, pertinent details of how they are not complying with medical care, public health recommendations, and /or what condition or behavior is putting others at significant risk of exposure to a notifiable disease or condition.

Department of Human Services Regulation, Bureau of Health 10-144 Chapter 258 (2) (B)

Duties of health care providers and attendants
Health care providers and persons attending a case of a notifiable disease shall arrange for such precautionary measures, consistent with the rules of the Department, including examination and isolation of the case when necessary, as are required to prevent the spread of infection to other members of the household or to the community. Proper isolation or other precautionary measures may be instituted by the Department or by the health officer after consultation with the Department. Notifiable disease cases shall receive immediate treatment according to the most recently established guidelines as promulgated by the appropriate professional organization and as are generally perceived to represent the current standard of care.

Non-compliant persons shall be reported to the Department for necessary interventions.

Department of Human Services Regulation, Bureau of Health 10-144 Chapter 258 (6)

Useful Links
Maine DHHS Bureau of Infectious Diseases
List of Reportable Diseases in Maine

(25) Consent to Treatment
Informed consent to health care treatment
Disallowance of recovery on grounds of lack of informed consent
No recovery may be allowed against any physician, physician assistant, podiatrist, dentist or any health care provider upon the grounds that the health care treatment was rendered without the informed consent of the patient or the patient's spouse, parent, guardian, nearest relative or other person authorized to give consent for the patient when:

A. The action of the physician, physician assistant podiatrist or dentist in obtaining the consent of the patient or other person authorized to give consent for the patient was in accordance with the
standards of practice among members of the same health care profession with similar training and experience situated in the same or similar communities;
B. A reasonable person, from the information provided by the physician, physician assistant, podiatrist or dentist under the circumstances, would have a general understanding of the procedures or treatments and of the usual and most frequent risks and hazards inherent in the proposed procedures or treatments which are recognized and followed by other physicians, physician assistants, podiatrists or dentists engaged in the same field of practice in the same or similar communities; or
C. A reasonable person, under all surrounding circumstances, would have undergone such treatment or procedure had that person been advised by the physician, physician assistant, podiatrist or dentist in accordance with paragraphs A and B or this paragraph.

For purposes of this subsection, the physician, physician assistant, podiatrist, dentist or health care provider may rely upon a reasonable representation that the person giving consent for the patient is authorized to give consent unless the physician, podiatrist, dentist or health care provider has notice to the contrary. (Note the statute does not refer specifically to Advanced Practice Registered Nurses, but they would be included under the category of “health care provider.”)

Presumption of validity of written consent; rebuttal
A consent which is evidenced in writing and which meets the foregoing standards, and which is signed by the patient or other authorized person shall be presumed to be a valid consent. This presumption, however, may be subject to rebuttal only upon proof that such consent was obtained through fraud, deception or misrepresentation of material fact.

Mental and physical competency
A valid consent is one which is given by a person who, under all the surrounding circumstances, is mentally and physically competent to give consent.

Informed consent for breast cancer
Duty of physician
Notwithstanding section 2905, a physician who is administering the primary treatment for breast cancer shall inform the patient as provided in this section, orally and in writing, about alternative efficacious methods of treatment of breast cancer, including surgical, radiological or chemotherapeutic treatments or any other generally accepted medical treatment and the advantages, disadvantages and the usual and most frequent risks of each.

Written information
The duty to inform the patient in writing may be met by giving the patient a standardized written summary or brochure as described in subsections 3 and 4.

Standardized written summary
The standardized written summary may be developed by the Bureau of Health after consultation with the Cancer Advisory Committee.

Brochure
The brochure must be one which is approved or made available through the National Cancer Institute, the American Cancer Society, the American College of Surgeons or any other recognized professional organization approved by the Bureau of Health.
Signed form
A form, signed by the patient, indicating that the patient has been given the oral information required by
this section and a copy of the brochure or the standardized written summary shall be included in the
patient's medical record.

Extent of duty
A physician's duty to inform a patient under this section does not require disclosure of information
beyond what a reasonably well-qualified physician licensed under Title 32 would know.

Actions barred
A patient who signs a form described in subsection 5 is barred from bringing a civil action against the
physician, based on failure to obtain informed consent, but only in regard to information pertaining to
alternative forms of treatment of breast cancer and the advantages, disadvantages, and risks of each
method.

Application of this section to common law rights
Nothing in this section restricts or limits the rights of a patient under common law.

Informed consent for opioid prescribing
Health care entities that included any individual prescribers of opioid medications must have prescribing
policies in place that included procedures and practices related to risk assessment, informed consent and
counseling on the risk of opioid use.

(26) Continuing Medical Education (see Agency Rules Links)
Continuing Medical Education
Each physician licensed by this Board who is actively practicing medicine and surgery shall complete
during each biennial licensing period, a minimum of one hundred (100) credit hours of continuing
medical education subject to the following:
A. At least forty (40) hours must be in Category 1, as defined by the Board rules;
B. The total one hundred (100) hours may be in Category 1;
C. No more than sixty (60) credit hours may be in Category 2, as defined by the Board rules;
D. At least three (3) hours every two years must be related to opioid prescribing.

If appropriate CME is not completed and submitted, then an Inactive status license renewal will be issued
unless the Board has granted an extension of time or deferment as described in the Board Rules.

Department of Professional and Financial Regulation, Board of Licensure in Medicine, Rule
Chapter 1, §8

(27) Controlled Substances Prescription Monitoring (see Opioid Medication Prescribing)

(28) Covenants Not to Compete
In your business negotiations, you may have encountered a “non-competition agreement” or a “covenant
not to compete” – a term of an employment agreement limiting a professional’s right to practice in the
geographic region of the employer for a period of time. Such employment contract provisions are not
uncommon in the Maine marketplace. Judicial decisions of non-competition agreements depend very
much on the facts of the case. However, the Code of Medical Ethics frowns on these terms because they are likely to reduce the availability of medical care to the public. Opinion 9.02, Restrictive Covenants and the Practice of Medicine states the following about non-competition agreements:

Covenants-not-to-compete restrict competition, disrupt continuity of care, and potentially deprive the public of medical services. The Council on Ethical and Judicial Affairs discourages any agreement which restricts the right of a physician to practice medicine for a specified period of time or in a specified area upon termination of an employment, partnership or corporate agreement. Restrictive covenants are unethical if they are excessive in geographic scope or duration in the circumstances presented, or if they fail to make reasonable accommodation of patients’ choice of physician. (VI, VII)

Issued prior to April 1977; Updated June 1994 and June 1998.

The leading Maine case on the interpretation of non-competition agreements in the health care context is Brignull v. Albert, a 1995 case involving two optometrists where the Maine Supreme Judicial Court enforced the non-competition agreement.

In Brignull, the plaintiff optometrist had offices in Ellsworth and Bar Harbor. The employment agreement between the parties provided that the defendant could not practice on Mount Desert Island or within 20 miles of Ellsworth for 4 years after leaving the practice and it provided for $30,000 in liquidated damages for violation of the provision. Within 2 years of departing the practice, the defendant opened an office within 2 miles of the plaintiff’s Ellsworth office and in the first 6 months saw 210 of the plaintiff’s patients. The Law Court found that the non-competition agreement was reasonable and that the liquidated damages provision was enforceable. It stated that the reasonableness of such an agreement is a question of law in each case depending on the duration, geographic area, and interests to be protected by the agreement. Simply protecting the employer from business competition would not, in the Court’s view, be a legitimate interest to be protected by a non-competition agreement. The legitimate interests identified in Brignull were to prevent the employee from taking existing patients and to protect the good will of the business.

You may review the AMA’s Code of Medical Ethics at the AMA website, https://www.ama-assn.org/delivering-care/ama-code-medical-ethics

**General**

Contractual agreements that restrict practice or competition, i.e. “covenants not to compete” are not unusual arrangements between professionals and other businesspersons. These restrictive covenants are commonly found in contracts regarding the sale or dissolution of a business or a professional practice. Although there are no state laws or regulations concerning the enforceability of such covenants, Maine courts have generally upheld the agreements where the restrictions regarding period of time and geographic area are found to be reasonable.

Examples:

A. A covenant never again to practice in Maine would most likely deemed unreasonable by the court.

B. A covenant not to practice for 3 years within a 10-mile radius would probably be deemed reasonable.
(29) Crimes

Cooperation with law enforcement

Reporting Crime Against Practitioner or on Premises

A healthcare practitioner or facility may disclose, or when required by law must disclose, health care information without authorization to disclose under the circumstances stated in this subsection or in subsection 11. Disclosures may be made without authorization to federal, state or local governmental entities in order to protect the public health and welfare when reporting is required or authorized by law, to report a suspected crime against the health care practitioner or facility or to report information that the health care facility's officials or health care practitioner in good faith believes constitutes evidence of criminal conduct that occurred on the premises of the health care facility or health care practitioner or to federal, state or local governmental entities if the health care practitioner or facility that is providing diagnosis, treatment or care to an individual has determined in the exercise of sound professional judgment that the disclosure is required by section 1727.

22 MRSA § 1711-C(6)(E)

Criminal Defendants Undergoing Treatment in a Health Care Facility

Note: None of these provisions of Maine state law relieve a health care practitioner from any privacy obligations under HIPAA. Whenever there is a conflict between the two, HIPAA must govern behavior. A hospital licensed under chapter 404 or 405 shall make a good faith effort to cooperate with law enforcement agencies, with limitations set forth in the statute.

Service of protection from abuse order

A law enforcement agency may request that a hospital provide access to a defendant who is receiving care in the hospital for the purpose of serving a protection from abuse order pursuant to Title 19–A, section 4006, subsection 6.

A. The hospital shall provide the law enforcement agency with an opportunity to serve the defendant personally with the order at a time the hospital determines is clinically appropriate with due consideration to the medical condition of the defendant.

B. A hospital may disclose that the defendant is a patient to facilitate service under this section regardless of patient consent.

22 MRSA § 1727 (1)

Notice of upcoming release

A law enforcement agency may request that a hospital provide notice to the law enforcement agency that a person is to be released from the hospital so that the law enforcement agency may arrest the person.

A. The hospital shall provide notice that the person is to be released from the hospital if the person was transported or was caused to be transported to the hospital by the law enforcement agency.

B. The information contained in the notice provided by the hospital must be no more than the minimum amount necessary to satisfy the requirements of this subsection.

22 MRSA § 1727 (2)

Required consistency with federal requirements

A hospital may provide access under subsection 1 and information under subsection 2 only if the request is consistent with the provisions of 45 Code of Federal Regulations, Section 164.512 (2015) and 42 Code of Federal Regulations, Part 2 (2015).

22 MRSA § 1727 (3)

Immunity; no cause of action

A hospital, hospital agent, employee or other person who in good faith and without gross negligence provides access or information to a law enforcement agency as required by this section or cooperates in an investigation or a criminal or judicial proceeding related to the requirements of this section is immune
from civil and criminal liability and professional licensure action arising out of or related to compliance with this section. This section does not create a cause of action against the hospital, hospital agent, employee or other person for failure to comply with this section.

22 MRSA § 17276 (4)

**Failure to report treatment of a gunshot wound**
A person is guilty of failure to report treatment of a gunshot wound if, being a health care practitioner or emergency medical services person, that person treats a human being for a wound apparently caused by the discharge of a firearm and knowingly fails to report the same to a law enforcement agency immediately by the quickest means of communication.

17-A MRSA §512

**Reports of death**
**Persons suspecting medical examiner case**
Any person who becomes aware of a suspected medical examiner case shall immediately notify a law enforcement officer or the Office of Chief Medical Examiner. As used in this subsection, "person" means a natural person, including a public servant, and a corporation, partnership, unincorporated association or any other nonhuman legal entity, including any governmental unit.

22 MRSA §3026 (1)

**Medical examiners suspecting medical examiner case**
Any medical examiner who becomes aware of a death caused by physical injury, or in which physical injury is the suspected cause, shall immediately notify the Office of Chief Medical Examiner and the appropriate law enforcement agency. The agency shall notify the district attorney for the district in which the body is located.

22 MRSA §3026 (3)

**Cases involving or suspected of involving physical injury attributable to criminal conduct**
Any law enforcement officer or medical examiner who becomes aware of a death involving physical injury attributable to criminal conduct, or in which physical injury attributable to criminal conduct is suspected, other than vehicular manslaughter, in addition to complying with the notification requirements in subsection 3, shall immediately notify the Attorney General.

22 MRSA §3026 (4)

**Employment leave for victims of violence**
**Definition**
For purposes of this subchapter, the terms "daughter," "son," "parent" and "spouse" have the same meanings as those terms have under federal regulations adopted pursuant to 29 United States Code, Section 2654, as in effect on January 1, 2002. An employer may require an employee to provide reasonable documentation of the family relationship, which may include a statement from the employee, a birth certificate, a court document or similar documents.

26 MRSA §850 (1-A)

**Required leave**
An employer must grant reasonable and necessary leave from work, with or without pay, for an employee to:

A. Prepare for and attend court proceedings;
B. Receive medical treatment or attend to medical treatment for a victim who is the employee's daughter, son, parent or spouse; or
C. Obtain necessary services to remedy a crisis caused by domestic violence, sexual assault or stalking.

The leave must be needed because the employee or the employee's daughter, son, parent or spouse is a victim of violence, assault, sexual assaults under Title 17-A, chapter 11, stalking or any act that would support an order for protection under Title 19-A, chapter 101.2.

26 MRSA §850 (1)
Exceptions
Subsection 1 is not violated if:

A. The employer would sustain undue hardship from the employee's absence;
B. The request for leave is not communicated to the employer within a reasonable time under the circumstances; or
C. The requested leave is impractical, unreasonable or unnecessary based on the facts then made known to the employer.

Civil penalties
If notice of a violation of this section is given to the employer and the Department of Labor within 6 months of the occurrence, the Department of Labor may assess penalties as follows:

A. For denial of leave in violation of this section, a fine of up to $1,000 for each violation of this section may be assessed. A fine assessed under this paragraph must be paid to the Treasurer of State. Additionally, the employer shall pay liquidated damages to the affected individual in an amount equal to 3 times the amount of total assessed fines; and
B. For termination in connection with an individual exercising a right granted by this section, the affected individual may elect to receive:
   1. Liquidated damages pursuant to paragraph A; or
   2. Reemployment with the employer with back wages.

Endangering welfare of dependent person
A. A person is guilty of endangering the welfare of a dependent person if:
   1. The person recklessly endangers the health, safety or mental welfare of a dependent person who is unable to perform self-care because of advanced age or physical or mental disease, disorder or defect. Violation of this paragraph is a Class D crime; or
   2. The person intentionally or knowingly endangers the health, safety or mental welfare of a dependent person who is unable to perform self-care because of advanced age or physical or mental disease, disorder or defect. Violation of this paragraph is a Class C crime.
B. As used in this section, "endanger" includes a failure to act only when the defendant has a legal duty to protect the health, safety or mental welfare of the dependent person. For purposes of this section, a legal duty may be inferred if the defendant has assumed responsibility for the care of the dependent person.

Deaths (Uniform Determination of Death Act)
Determination of death
An individual who has sustained either (1) irreversible cessation of circulatory and respiratory functions, or (2) irreversible cessation of all functions of the entire brain, including the brain stem, is dead. A determination of death must be made in accordance with accepted medical standards.

Registration of fetal deaths
Except as authorized by the department or as required under section 1596, a certificate of each death of a fetus of 20 or more weeks of gestation must be filed with the State Registrar of Vital Statistics or the clerk of the municipality where the delivery occurred. Such registration must occur within 14 days after delivery and prior to removal of the fetus from the State.

Certificate filed by funeral director
The funeral director or other authorized person in charge of the disposition of the dead fetus or its removal from the State is responsible for filing the certificate. In the absence of such a person, the physician, the certified nurse midwife, the nurse practitioner or other person in attendance at or after the...
delivery shall be responsible for filing the certificate. The person responsible for filing the certificate shall obtain the personal data from the best qualified person or source available and shall present the certificate to the person responsible for completing the medical certification of the cause of death.

Medical certificate by physician
The medical certification must be completed and signed within 5 days after delivery by the physician in attendance at or after the delivery, except when an inquiry as to the cause of fetal death is required by law.

Medical certificate by medical examiner
When the fetal death occurs without medical attendance upon the mother at or after delivery, or when inquiry as to the cause of fetal death is required by law, the medical examiner shall complete and sign the medical certification within 5 days after delivery. A certification need not be completed before the remains are ready for release.

Certificate from hospital or institution
When the fetal death occurs in a hospital or an institution, the person in charge of the hospital or institution or the person authorized to obtain the medical data shall prepare the certificate, certify by signature or by electronic process that the fetal death occurred at the place and time and on the date stated and file the certificate as directed in this section. The physician or other person in attendance shall provide the medical information required on the certificate in a timely fashion, as specified by department rule.

Registration of deaths
Except as authorized by the department, a certificate of each death which occurs in this State shall be filed with the State Registrar of Vital Statistics or clerk of the municipality where death occurred within a reasonable period of time, as specified by department regulation, after the day on which death occurred and prior to the removal of the body from the State.

Certificate filed by funeral director
The funeral director or other authorized person in charge of the disposition of the dead human body or its removal from the State is responsible for filing the certificate. He shall obtain the personal data from the best qualified person or source available and he shall present the certificate to the physician or medical examiner responsible for completing the medical certification of the cause of death.

Definitions
A. Life-sustaining procedure: means any medical procedure or intervention that, when administered to a qualified patient, will serve only to prolong the dying process and does not include nutrition and hydration.
B. Terminally ill patient: means a patient who has been diagnosed as having an incurable or irreversible condition that, without the administration of life-sustaining procedures, will, in the opinion of the attending health care provider, result in death within a short time.
C. Health care provider: means a physician authorized to practice in this State, nurse practitioner, or physician assistant.

Medical certificate by physician, nurse practitioner, or physician assistant
The medical certification of the cause of death must be completed and signed in a timely manner, as specified by Department rule, by a physician, nurse practitioner, or physician assistant authorized to practice in the State who has knowledge of the patient's recent medical condition, in accordance with department regulations and other laws detailing who can certify and in what time frame, except when the death falls under the jurisdiction of the medical examiner as provided in section 3025. If the patient was a resident of a nursing home licensed under section 1817 at the time of death and if the health care provider
in charge of the patient's care or another health care provider designated by the health care provider in charge had not examined the patient within 48 hours prior to death, or within 2 weeks prior to death in the case of a terminally ill patient, the health care provider in charge or another health care provider designated by the health care provider in charge shall examine the body prior to completing the certification of death process. Any health care provider who fails to complete the medical certification of the cause of death fully and in a timely manner, or who fails to examine the body of a nursing home resident prior to certifying cause of death as required by this section must be reported to the Board of Licensure in Medicine, the Board of Osteopathic Licensure or the State Board of Nursing, whichever is appropriate, by the State Registrar of Vital Statistics of the Department of Health and Human Services.

Medical certification
Notwithstanding subsection 2, with respect to a person who dies within the State naturally and for whom the physician, nurse practitioner, or physician assistant was the attending health care provider, the medical certification of the cause of death may be completed and signed by a physician, nurse practitioner, or physician assistant authorized to practice at the U.S. Department of Veterans’ Affairs hospital at Togus or at another federal medical facility within the State or by a physician, physician assistant, or advanced practice registered nurse licensed to practice in New Hampshire, Vermont or Massachusetts who, at the request of the Chief Medical Examiner, is willing to do so.

Medical certificate by medical examiner
When a death occurs under circumstances that make it a medical examiner case as defined in section 3025, or when inquiry as to the cause of death is required by law, the medical examiner or the Office of the Chief Medical Examiner shall complete the medical certification of the cause of death and sign the death certificate. A certification need not be completed before the remains are ready for release.

The medical examiner or the Office of the Chief Medical Examiner is responsible for the identity of the deceased and the time, date, place, cause, manner and circumstances of death on the death certificate. Entries may be left "pending" if further study is needed; or, at the specific direction of the Attorney General relative to cases under investigation by the Attorney General's office, entries must be left "withheld" until such time as the Attorney General, in the Attorney General's sole discretion, determines that any criminal investigation and prosecution will not be harmed by public disclosure of such information. Notwithstanding section 2706, subsection 4, unless directed otherwise by the Attorney General as specified in this subsection, this information for which the medical examiner is responsible may be made available to the general public by the Office of Chief Medical Examiner.

Correction of errors on death statistic records filed under chapter 711
Certificates of death in medical examiner cases, as defined in section 3025, may be completed or amended at any time by means described in rule by the department to the Office of Chief Medical Examiner. Either the Chief Medical Examiner or the medical examiner assigned to the case may sign the forms or submit an electronic amendment or file a certificate using the electronic death registration system in accordance with section 2847. A person authorized by the Chief Medical Examiner may amend a certificate of death with respect to the time, date, place and circumstances of death. The medical examiner assigned shall submit the form or electronic amendment to the Office of the Chief Medical Examiner for filing with the State Registrar of Vital Statistics. These forms or electronic amendments may be filed at any time after death and need not include a summary description of the evidence in support of the completion or amendment.

Electronic death registration system
A certificate of death required to be filed by any person authorized under section 2842 pursuant to this chapter must (as of 7/1/18) be filed using the electronic death registration system maintained by the State
Registrar of Vital Statistics…The State Registrar of Vital Statistics shall adopt rules to carry out the purposes of this section.

22 MRSA § 2847

**DHHS Rules for the Medical Certification of the Cause of Death (see Agency Rules Links)**

**Maine Elder Death Analysis Review Team**
The Maine Elder Death Analysis Review Team in the Attorney General’s Office is expanded to 16 members from 13, and now includes a sexual assault nurse examiner, a physician and an emergency medical services professional.

For a list of the composition of the team see 5 MRSA §200-H (1).

(31) **Delegation; see Staffing & Supervision**

(32) **Dentists & Dental Hygenists**
For the statutes governing the practice of dentists, dental hygenists, dental assistants, independent practice dental hygenists, dentuists and dental radiographers, see Title 32, Chapter 143 of the Maine Revised Statutes. See also the Agency Rules Links.

Protocols for professional review committeeThe Board of Dental Examiners may establish protocols for the operation of a professional review committee as defined in Title 24, section 2502, subsection 4-A. The protocols must include the committee reporting information the board considers appropriate regarding reports received, contracts or investigations made and the disposition of each report, provided that the committee is not required to disclose any personally identifiable information. The protocols may not prohibit an impaired dentist from seeking alternative forms of treatment.

32 MRSA §18323 (8)

(33) **Dirigo Health Program**

The Dirigo Health Program was Governor John Baldacci’s signature health care reform program enacted by the legislature in 2003 to address the “three legs” of the health care reform “stool:” access (the DirigoChoice product), quality (the Maine Quality Forum), and cost (Advisory Council on Health Systems Development, State Health Plan, and Certificate-of-Need [CON] program). You can learn more about the Dirigo Health Agency on its website: [http://www.dirigohealth.maine.gov](http://www.dirigohealth.maine.gov). The Dirigo health insurance programs ended as of December 31, 2013 as the Health Insurance Marketplace created by the Affordable Care Act was phased in.

You can find the Dirigo statute, 24-A M.R.S.A., Chapter 87 here: [http://www.mainelegislature.org/legis/statutes/24-A/title24-Ach87sec0.html](http://www.mainelegislature.org/legis/statutes/24-A/title24-Ach87sec0.html)

**Maine Quality Forum**
The Maine Quality Forum (MQF) was established by the Governor and the legislature in September 2003. The MQF has been charged with: collecting research, promoting best practices, collecting and publishing comparative quality data, promoting electronic technology, promoting healthy lifestyles and reporting to consumers and the Legislature. The Maine Quality Forum's mission is to advocate for high quality healthcare and help each Maine citizen make informed healthcare choices. You can find more information about the quality improvement role of the Maine Quality Forum on its website: [http://www.mainequalityforum.gov/](http://www.mainequalityforum.gov/)
Disabilities

Americans with Disabilities Act

Accessibility of doctors’ offices, clinics, and other health care providers is essential in providing medical care to people with disabilities. Due to barriers, individuals with disabilities are less likely to get routine preventative medical care than people without disabilities. Accessibility is not only legally required, it is important medically so that minor problems can be detected and treated before turning into major and possibly life-threatening problems.

The Americans with Disabilities Act of 1990 (ADA) is a federal civil rights law that prohibits discrimination against individuals with disabilities in every day activities, including medical services. Section 504 of the Rehabilitation Act of 1973 (Section 504) is a civil rights law that prohibits discrimination against individuals with disabilities on the basis of their disability in programs or activities that receive federal financial assistance, including health programs and services. These statutes require medical care providers to make their services available in an accessible manner.

The ADA requires access to medical care services and the facilities where the services are provided. Private hospitals or medical offices are covered by Title III of the ADA as places of public accommodation. Public hospitals and clinics and medical offices operated by state and local governments are covered by Title II of the ADA as programs of the public entities. Section 504 covers any of these that receive federal financial assistance, which can include Medicare and Medicaid reimbursements. The standards adopted under the ADA to ensure equal access to individuals with disabilities are generally the same as those required under Section 504.

Both Title II and Title III of the ADA and Section 504 require that medical care providers provide individuals with disabilities:

A. full and equal access to their health care services and facilities; and
B. reasonable modifications to policies, practices, and procedures when necessary to make health care services fully available to individuals with disabilities, unless the modifications would fundamentally alter the nature of the services (i.e. alter the essential nature of the services).

In addition, all buildings, including those built before the ADA went into effect, are subject to accessibility requirements for existing facilities. Under Title III, existing facilities are required to remove architectural barriers where such removal is readily achievable. Barrier removal is readily achievable when it is easily accomplishable and able to be carried out without much difficulty or expense. If barrier removal is not readily achievable, the entity must make its services available through alternative methods, if those methods are readily achievable. Under Title II, a public entity must ensure that its program as a whole is accessible; this may entail removing architectural barriers or adopting alternative measures, such as relocating activities to accessible locations. This same program accessibility standard applies under Section 504.

For further guidance on complying with the ADA, see:
- The U.S. Department of Justice Primer for Small Businesses
- US Department of Justice Access to Medical Care for Individuals with Motility Disabilities
- US Department of Justice ADA Homepage

Maine Human Rights Act (Select sections)

Many of the anti-discrimination sections of the Maine Human Rights Act are similar to those of the provisions found in the American’s with Disabilities Act. The Maine Human Rights Act is found at 5 MRSA, Chapter 337 and prevents discrimination in employment, housing, education, access to public accommodations and credit transactions on the basis of an individual’s characteristics including physical or
mental disability. For more information about the Maine Human Rights Act and Maine Human Rights Commission, see: http://www.maine.gov/mhrc/about/index.htm

(35) Driving, Reporting of Impaired Drivers

Optional reporting of drivers operating under the influence of intoxicating liquor or drugs

Persons who may report
If, while acting in a professional capacity, a medical or osteopathic physician, resident, intern, emergency medical services person, medical examiner, physician's assistant, dentist, dental hygienist, dental assistant or registered or licensed practical nurse knows or has reasonable cause to believe that a person has been operating a motor vehicle, hunting or operating a snowmobile, all-terrain vehicle or watercraft while under the influence of intoxicants and that motor vehicle, snowmobile, all-terrain vehicle or watercraft or a hunter has been involved in an accident, that person may report those facts to a law enforcement official.

29-A MRSA §2405 (1)

Immunity from liability
A person participating in good faith in reporting under this section, or in participating in a related proceeding, is immune from criminal or civil liability for the act of reporting or participating in the proceeding.

Nothing in this section may be construed to bar criminal or civil action regarding perjury.

In a proceeding regarding immunity from liability, there is a rebuttable presumption of good faith.

29-A MRSA §2405 (2)

Privileged or confidential communications
The physician-patient privileges under the Maine Rules of Evidence and the confidential quality of communication under Title 24-A, section 4224 and Title 32, section 1092-A are abrogated in relation to required reporting or other proceeding.

29-A MRSA §2405 (3)

Reporting of Those Unsafe to Drive Due to Age, Medical Condition
A member of the [Department of Motor Vehicles Medical Advisory] board or other person making an examination and report of opinion, recommendation or advice to the Secretary of State in good faith is immune from criminal or civil liability for so doing. A physician or other person who becomes aware of a physical, mental or emotional impairment that appears to present an imminent threat to driving safety and reports this information to the Secretary of State in good faith is immune from criminal or civil liability for so doing. The immunity for damages under this subsection applies only to the extent that this immunity is not in conflict with federal law or regulation.

29-A M.R.S.A. §1258 (6)

Contact the Maine Medical Association for further resources and training materials on working with senior drivers.

(36) Drugs, Prescription

Prescriber Laws & Rules (select sections)

Prescription Monitoring Program
The state legislature passed a law in 2003 that requires information about all transactions for Schedule II, III, and IV controlled substances dispensed in Maine to be reported to the state government. Pharmacies – both in and out of the state – submit data weekly. The data is then cleaned and added to a relational database. Using patients’ names and birth dates, registered users of the database can log on to the web site at www.maine.gov/pmp to look up their patients online. Clinicians have immediate access to a patient’s
history with controlled substances freely available at their fingertips. (To register to become a requester or submitter of data, go to www.maine.gov/pmp and follow the instructions there.)

In 2016, the significant changes to the opioid prescribing laws included a requirement to check the prescription monitoring program. (See Opioid Medication Prescribing for more detail.)

For more information about the prescription monitoring program, see http://www.maine.gov/dhhs/samhs/osa/data/pmp/index.htm

For rules applying to the program, see: http://www.maine.gov/dhhs/samhs/osa/data/pmp/prescriber.htm

**Automatic enrollment**
A law passed in 2014 directs the Prescription Monitoring Program to streamline the process for enrollment in the PMP. Based on the recommendations of the Substance Abuse Services Commission the law states that the PMP must implement a mechanism for prescribers of controlled substances to be enrolled automatically as a part of the process of applying for or renewing a professional license. The change impacts prescribers who are registering with the program for the first time or are renewing registration. The new law also repeals the section of statute that made registration mandatory as of March 1, 2014, if 90% or more of a class of prescribers had not registered.

22 MRSA § 7249(5)

**Identification of persons prescribing medicines on hospital prescription blanks**
Any practitioner who writes a prescription upon a prescription blank of a hospital or clinic shall sign that practitioner's name and cause that name to be printed, stamped or typed on the blank.

This section applies to any physician's assistant or registered nurse who writes a prescription while working under the control or supervision of a physician. In case of the physician's assistant or registered nurse, the name of the physician under whom the assistant or nurse works shall be printed, stamped or typed on the blank.

32 MRSA §13786

**Prohibition on excessive copayments or charges**
Copayments or other charges may not exceed claim cost of any prescription drug.

24-A MRSA §4317 (13)

**Prescription drug price transparency**
By December 1, 2018 and annually thereafter, the Maine Health Data Organization shall provide a report containing the following information about prescription drugs, both brand name and generic:

A. The 25 most frequently prescribed drugs in the State;
B. The 25 costliest drugs as determined by the total amount spent on those drugs in the State; and
C. The 25 drugs with the highest year-over-year cost increases as determined by the total amount spent on those drugs in the State.

22 MRSA §8712 (5)

**Possession of drug samples**
No person may purchase manufacturers' drug samples from any person for purposes of resale. If those samples are given gratuitously to a registered pharmacist, qualified assistant pharmacist or medical practitioner, any such sample may be given to any person, provided that any such sample is kept in containers suitably labeled to conform to the Federal Food and Drug Act and the state food and drug laws and provided that this gift shall be subject to the laws relating to the sale of drugs.

32 MRSA §13789
**Acquiring drugs by deception**

A person is guilty of acquiring drugs by deception if, as a result of deception, the person obtains or exercises control over a prescription for a scheduled drug or what the person knows or believes to be a scheduled drug, which is in fact a scheduled drug.

17-A MRSA §1108 (1)

**Deception defined**

As used in this section, “deception” has the same meaning as in section 354, subsection 2 and includes:

A. Failure by a person, after having been asked by a prescribing health care provider or a person acting under the direction or supervision of a prescribing health care provider, to disclose the particulars of every narcotic drug or prescription for a narcotic drug issued to that person by a different health care provider within the preceding 30 days; or

B. Furnishing a false name or address to a prescribing health care provider or a person acting under the direction or supervision of a prescribing health care provider.

17-A MRSA §1108 (2)

**Reporting the Crime of Acquiring Drugs by Deception**

For the purposes of this section, information communicated to a prescribing health care provider, or a person acting under the direction or supervision of a prescribing health care provider, in an effort to violate this section, including a violation by procuring the administration of a scheduled drug by deception, may not be deemed a privileged communication.

17-A MRSA §1108 (3)

**Immunity from criminal or civil liability**

A prescribing health care provider, or a person acting under the direction or supervision of a prescribing health care provider, who knows or has reasonable cause to believe that a person is committing or has committed deception may report that fact to a law enforcement officer. A person participating in good faith in reporting under this subsection, or in participating in a related proceeding, is immune from criminal or civil liability for the act of reporting or participating in the proceeding.

17-A MRSA §1108 (6)

**Reporting Crime Against Practitioner or on Premises**

A health care practitioner or facility may disclose, or when required by law must disclose, health care information without authorization to disclose under the circumstances stated in this subsection or as provided in subsection 11. Disclosure may be made without authorization as follows: (E) To federal, state or local governmental entities in order to protect the public health and welfare when reporting is required or authorized by law, to report a suspected crime against the health care practitioner or facility or to report information that the health care facility's officials or health care practitioner in good faith believes constitutes evidence of criminal conduct that occurred on the premises of the health care facility or health care practitioner, or to federal, state or local governmental entities if the health care practitioner or facility that is providing diagnosis, treatment or care to an individual has determined, in the exercise of sound professional judgment, that the disclosure is required by section 1726.

22 MRSA § 1711-C (6)

**Use of Controlled Substances for the Treatment of Pain (Licensing Boards Chapter 21)**

For the full guidance on Use of Controlled Substances For Treatment of Pain, see the Board of Licensure in Medicine Rule Chapter 21.

**Principles of Proper Patient Management**

The Board of Licensure in Medicine has adopted the following criteria when evaluating the clinician’s treatment of pain including the use of controlled substances. Each of these principles is essential in the treatment of patients with pain.
The Board of Licensure in Medicine, the Board of Osteopathic Licensure, and the Board of Nursing have established very specific joint rules under “Chapter 21” that apply to all prescribers licensed by those boards. Those rules require the following:

**Universal Precautions**
1. Patient evaluation
   a. Medical history & physical (specific items to be included are listed in rule)
   b. Risk assessment (specific items to be included are listed in rule)

**Treatment Plan**
Requirements for initiating or continuing prior opioid therapy
- Start with lowest possible dose and titrate to effect based on documented functional assessment
- Immediate release rather than extended release when initiating
- Begin as trial for no more than 30 days, evaluate harms and benefits within 28 days
- Dosage limits
- Dosage limit exemptions (as in statute and DHHS rules)
- Electronic prescriptions, time limits, and rules for palliative care exemption
- Naloxone for high risk patients
- Avoid concurrent opioid and benzodiazepine prescription whenever possible

Periodic review of treatment efficacy, with frequency based on risk level (specified requirements for review)

Consult or refer for additional evaluation or treatment as necessary

Coordination of care with other clinicians who have narcotic contract with patient

Tapering and/or managed withdrawal or treatment if opioid therapy is discontinued

**Informed Consent**
- Written and signed
- Minimum contents
- Benefits: pain reduction, improved physical and psychological function
- Risks
- Side effects
- Effect on vehicle operation
- Allergic reactions
- Medication interaction
- Likelihood of tolerance or dependence
- Risks of misuse, addiction and potentially fatal overdose
- Potential for withdrawal symptoms
- Risk of fatal overdose due to accidental exposure, especially to children
- Risk of use during pregnancy

**Use of PMP**

**Treatment Agreement**
List of required provisions, including:
- Disclose all medical conditions and medications
• Responsibility to be discreet about possessing narcotics
• Take only as prescribed
• Prescribing policies and expectations, including use of single pharmacy and policy on early and after-hour refills
• Disclose any other opioids received from other clinicians
• Keep scheduled appointments, comply with random pill counts and urine/blood testing
• Statement that clinician may notify authorities if concerned re: illegal activity
• Statement that violation of agreement may result in decrease or termination of opioid prescriptions
• Clinician must document all violations and response to violation, along with rationale for any changes in treatment plan

**Toxicological Drug Screens**
If prescription for 90+ days, documented screen prior to treatment initiation and randomly thereafter, at least annually

**Pill Counts**
“An additional tool”, not clear if required or merely encouraged. Must be documented

**Medical Records**
Specific list of what records must contain

**CDC Guidelines**
Prescribers “must be aware of and follow” US CDC guidelines on opioids and chronic pain prescribing

**Continuing Education**
All prescribers licensed by Board of Licensure in Medicine must take 3 hours of Category 1 CME on opioid prescribing, regardless of whether they prescribe opioids.

All prescribers licensed by Board of Osteopathic Licensure or Board of Nursing must take 3 hours of Category 1 CME on opioid prescribing as a condition of prescribing opioids.

*Board of Licensure in Medicine and Board of Osteopathic Licensure Chapter 21 (§3)*

**Limits on coverage of opioids by MaineCare**
• If a patient has new acute pain, and opioids are allowed, prescribers will only be able to prescribe them for up to 28 days (cumulative maximum). If the patient needs more than 15 days of opioid treatment, it is considered treatment for chronic pain.
• Opioid treatment for chronic pain requires the following:
  o Thorough history and physical exam, including an opioid therapy risk assessment, at the initial visit for pain management;
  o PMP review by the prescriber to verify no concomitant narcotic use, documented in the medical record;
  o Treatment plan that addresses realistic goals for pain and function, includes nonpharmacologic and non-opioid pharmacologic therapy, includes strategies to mitigate risk, and includes a plan for how therapy will be discontinued if
  o benefits do not outweigh risks
  o Review of the treatment plan at least every 90 days;
  o Counseling the patient about side effects, risks and benefits of opioid use at the initial visit, annually, and any time there is a dosage change;
• Urine drug test of other toxicology test before the start of treatment and considered at least quarterly, on a random basis;
• Immediate-release opioids must be tried before extended-release, long acting opioids whenever possible;
• Benefits and harms of continued opioid therapy must be reviewed with patient at least every 6 months, face-to-face;
• Prescribe accompanying naloxone if there are risk factors such as mental health disorder, substance use disorder, medical condition that increases sensitivity, or concurrent benzodiazepine use;
• If clinically indicated, a written treatment agreement outlining the patient’s responsibilities, consequence of loss or shortage of medications, consequence of obtaining similar prescriptions from other prescribers, and an agreement to use only one pharmacy.

• Prescribers are required to follow Maine law and prescribing guidelines and prescription monitoring program rules.

• Prior authorizations, as follows:
  • For acute pain, a face-to-face visit at initial prescription. Each authorization will allow up to 7 days’ coverage.
  • Prior authorization required after a total of 7 days of opioids prescribed for treatment of acute pain within a calendar year.
  • For chronic pain, prior authorization requires referral to 2 or more alternative treatment options; actively waiting for alternative treatment to begin or active participation in alternative treatment; alternative treatment progress notes in the record; and no clinically meaningful improvement in function or pain within the last prescription dose period.
  • Alternative treatment options include nonpharmacologic treatments, such as physical therapy, occupational therapy, osteopathic manipulation, chiropractic treatment, outpatient counseling, psychological therapies, Eye Movement Desensitization and Reprocessing (EMDR), and nonopioid pharmacologic treatments, such as acetaminophen, NSAIDS, gabapentin, and selected antidepressants. Benzodiazepines will not be considered an alternative treatment option.
  • Prescriptions shall not exceed 30 days, and prior authorizations shall not exceed 6 months.

• Medical records shall include the following:
  • History and physical exam;
  • Diagnostic, therapeutic and laboratory results;
  • Documentation of urine drug tests and review of results with patient;
  • Evaluations and consultations;
  • PMP reviews;
  • Treatment objectives’
  • Discussion of risks and benefits;
  • Informed consent;
  • Treatments;
  • All medications, including date, type, dosage, and quantity prescribed;
  • Instructions and agreements;
  • Treatment planning update every 90 days including nonpharmacological and non-opioid pharmacological treatment, plans to mitigate risk, and plan to discontinue treatment if benefits do not outweigh risks.
Buprenorphine and Buprenorphine combination products for substance use disorder treatment covered by MaineCare

MaineCare’s coverage of buprenorphine, a Schedule III narcotic, is subject to strict limitations on members qualified to receive the drug, rules regarding prior authorization, and clearly defined maximum daily dosages. MaineCare covers multiple formulations of the drug buprenorphine only for a member who has a diagnosis of Substance Use Disorder (SUD). Prescribers must have XDEA number or military service identification number, noted on every prescription for a controlled substance. Confidentiality regulations relating to treatment for substance use disorder must be followed, along with federal and state prescribing guidelines.

Treatment requires comprehensive screening and assessment, including educational needs, vocational rehabilitation needs, employment needs, medical support services, psychosocial support services, economic and legal support services, and other special needs. Patients must be counseled on side effects, risks and benefits, and all available options.

PMP review is required.

Urine drug tests or other medically appropriate toxicology test are required initially and randomly thereafter, with a minimum of 8 such tests per year.

Detailed treatment plan must include dosage for induction and maintenance phases, projected frequency of office visits, projected counseling and referral, treatment goals, and conditions under which treatment is to be discontinued. Plan must be updated every 90 days.

Patient must be referred to counseling.

Written treatment agreements are required if clinically indicated.

Patient must also agree to the following:

- Return to prescriber’s office as instructed during induction period;
- Random screenings;
- Active participation in substance use disorder counseling;
- Appearance within 24 hours of receiving a random screening call;
- Bring medications to all prescriber appointments to be counted;
- Avoid all “illegal or inappropriate” substances of abuse;
- Provide random urine samples, and meet with monitoring physician if test is positive;
- Buprenorphine kept at home must be locked in a safe place to prevent accidental use by others, especially children.

In case of involuntary termination of treatment, patient must be referred to other treatment providers, or withdrawal must be managed appropriately to minimize withdrawal discomfort if treatment is not being transferred.

Requirements for medical records, prior authorizations, and other opioid prescribing requirements must be followed.

For more information, visit the Provider Section of the MaineCare website.
**Authorized possession by individuals**
A person to whom or for whose use any scheduled drug, prescription drug or controlled substance has been prescribed, sold or dispensed for a legitimate medical purpose by a physician, dentist, podiatrist, pharmacist or other person acting in the usual course of professional practice and authorized by law or rule to do so and the owner or the person having the custody or control of any animal for which any scheduled drug, prescription drug or controlled substance has been prescribed, sold or dispensed for a legitimate veterinary medical purpose by a licensed veterinarian acting in the usual course of professional veterinary practice may lawfully possess the drug or substance, except when in use, only in the container in which it was delivered by the person selling or dispensing the drug or substance. “When in use” includes reasonable repackaging for more convenient legitimate medical use.

**22 MRSA §2383-B (1)**

**Others lawfully in possession**
Except as otherwise authorized or restricted, the following persons are authorized to possess, furnish and have control of scheduled or prescription drugs, controlled substances or hypodermic apparatuses:

A. Common carriers or warehouse operators while engaged in lawfully transporting or storing prescription drugs or hypodermic apparatuses or any of their employees acting within the scope of their employment;

B. Employees or agents of persons lawfully entitled to possession who have temporary, incidental possession while acting within the scope of their employment or agency;

C. Persons whose possession is for the purpose of aiding public officers in performing their official duties while acting within the scope of their employment or duties;

D. Law enforcement officers while acting within the scope of their employment and official duties

E. Physicians, dentists, podiatrists, pharmacists or other persons authorized by law or rule to administer, dispense, prescribe or sell scheduled or prescription drugs, controlled substances or hypodermic apparatuses while acting within the course of their professional practice;

F. With regard to the possession or furnishing of hypodermic apparatuses, persons authorized by the Bureau of Health pursuant to a hypodermic apparatus exchange program, certified under chapter 252-A while acting within the scope of their employment under such programs; and

G. Persons conducting research at a school of pharmacology that is accredited or is a candidate for accreditation in good standing.

**22 MRSA §2383-B (2)**

**Naloxone**

**Prescription; Possession; Administration**

A. A health care professional may directly or by standing order prescribe naloxone hydrochloride to an individual at risk of experiencing an opioid-related drug overdose.

A-1 pharmacist may dispense naloxone hydrochloride in accordance with protocols established under Title 32, section 13815 to an individual at risk of experiencing an opioid-related drug overdose.

A-2. A pharmacist may prescribe and dispense naloxone under protocols established by the Board of Pharmacy under 32 MRSA §13815.

B. An individual to whom naloxone hydrochloride is prescribed or dispensed in accordance with paragraph A or A-1 may provide the naloxone hydrochloride so prescribed or dispensed to a member of that individual's immediate family to possess and administer to the individual if the family member believes in good faith that the individual is experiencing an opioid-related drug overdose.

C. A health care professional may directly or by standing order prescribe naloxone hydrochloride to a member of an individual's immediate family or a friend of the individual or to another person in a position to assist the individual if the individual is at risk of experiencing an opioid-related drug overdose.

C-1 A pharmacist may prescribe and dispense naloxone hydrochloride in accordance with protocols established under Title 32, section 13815 to a member of an individual's immediate family or a
friend of the individual or to another person in a position to assist the individual if the individual is at risk of experiencing an opioid-related drug overdose.

D. If a member of an individual's immediate family, friend of the individual or other person is prescribed or provided naloxone hydrochloride in accordance with paragraph C or C-1, that family member, friend or other person may administer the naloxone hydrochloride to the individual if the family member, friend or other person believes in good faith that the individual is experiencing an opioid-related drug overdose.

Nothing in this subsection affects the provisions of law relating to maintaining the confidentiality of medical records.

Community-based drug overdose prevention programs; standing orders for naloxone

Acting under standing orders from a licensed health care professional authorized by law to prescribe naloxone hydrochloride, a public health agency that provides services to populations at high risk for a drug overdose may establish an overdose prevention program in accordance with rules adopted by the department and the provisions of this subsection.

A. Notwithstanding any other provision of law, an overdose prevention program established under this subsection may store and dispense naloxone hydrochloride without being subject to the provisions of Title 32, chapter 117 (Maine Pharmacy Act) as long as these activities are undertaken without charge or compensation.

B. An overdose prevention program established under this subsection may distribute unit-of-use packages of naloxone hydrochloride and the medical supplies necessary to administer the naloxone hydrochloride to a person who has successfully completed training provided by the overdose prevention program that meets the protocols and criteria established by the department, so that the person may possess and administer naloxone hydrochloride to an individual who appears to be experiencing an opioid-related drug overdose.

Immunity

The following provisions provide immunity for actions taken in accordance with this section.

A. A health care professional or a pharmacist, acting in good faith and with reasonable care, is immune from criminal and civil liability and is not subject to professional disciplinary action for storing, dispensing or prescribing naloxone hydrochloride in accordance with this section or for any outcome resulting from such actions.

B. A person, acting in good faith and with reasonable care, is immune from criminal and civil liability and is not subject to professional disciplinary action for possessing or providing to another person naloxone hydrochloride in accordance with this section or for administering naloxone hydrochloride in accordance with this section to an individual whom the person believes in good faith is experiencing an opioid-related drug overdose or for any outcome resulting from such actions.

Pharmacy/Dispenser Laws & Rules (select sections)

Generic and therapeutically equivalent substitution

A written prescription issued by a practitioner in this State may contain a box in the lower right-hand corner of the prescription form. The following words must appear to the left of this box: "Any drug which is the generic and therapeutic equivalent of the drug specified above in this prescription must be dispensed, provided that no check mark (✓) has been handwritten in the box in the lower right-hand corner."

Except with regard to a patient who is paying for a drug with the patient’s own resources, any pharmacist receiving a prescription in which no handwritten check mark (✓) is found in the box provided shall
substitute a generic and therapeutically equivalent drug for the drug specified on the prescription if the substituted drug is distributed by a business entity doing business in the United States that is subject to suit and the service of legal process in the United States and the price of the substituted drug does not exceed the price of the drug specified by the practitioner; except that, when the cost of a prescription is to be reimbursed under the MaineCare program pursuant to Title 22, chapter 855, the pharmacist shall substitute a generic and therapeutically equivalent drug only when the Department of Health and Human Services has determined that the substitute drug would be a more cost-effective alternative than the drug prescribed by the practitioner. Except for prescribed drugs listed under the Comprehensive Drug Abuse Prevention and Control Act of 1970, 21 United States Code, Section 812, as amended, as Schedule II drugs, with regard to a patient who is paying for a drug with the patient’s own resources, a pharmacist shall inquire about the patient’s preference for either the brand-name drug or generic and therapeutically equivalent drug and dispense the drug that the patient prefers.

Except with regard to a patient who is paying for a drug with the patient’s own resources, if a written prescription issued by a practitioner in this State does not contain the box described in this section, a pharmacist shall substitute a generic and therapeutically equivalent drug for the drug specified on the prescription if the substituted drug is distributed by a business entity doing business in the United States that is subject to suit and the service of legal process in the United States and the price of the substituted drug does not exceed the price of the drug specified by the practitioner, unless a practitioner has handwritten on the prescription form, along with the practitioner’s signature, “dispense as written,” “DAW,” “brand” “brand necessary” or “brand medically necessary”; except that, when the cost of a prescription is to be reimbursed under the MaineCare program pursuant to Title 22, chapter 855, the pharmacist shall substitute a generic and therapeutically equivalent drug only when the Department of Health and Human Services has determined that the substitute drug would be a more cost-efficient alternative than the drug prescribed by the practitioner. Except for prescribed drugs listed under the Comprehensive Drug Abuse Prevention and Control Act of 1970, 21 United States Code, Section 812, as amended, as Schedule II drugs, with regard to a patient who is paying for a drug with the patient’s own resources, a pharmacist shall inquire about the patient’s preference for either the brand-name drug or generic and therapeutically equivalent drug and dispense the drug that the patient prefers.

Any pharmacist who substitutes a generic and therapeutically equivalent drug under this section shall inform the person to whom the drug is dispensed of the substitution. When any substitution is made under this section, the pharmacist shall cause the name of the generic and therapeutically equivalent drug, the name or abbreviation of the drug manufacturer or distributor of that substitute drug and all other information as required by section 13794 to appear on the container label of the drug dispensed.

This section does not apply to prescriptions ordered by practitioners for patients in hospitals when those prescriptions are filled by a hospital pharmacy or in any institution where a formulary system is established.  

32 MRSA §13781

Patient profile record system regulation
A patient profile record system shall be maintained in all pharmacies for persons for whom prescriptions are dispensed. The patient profile record system shall be devised to enable the immediate retrieval of information necessary for the dispensing pharmacist to identify previously dispensed medication at the time a prescription is presented for dispensing. One profile record or document may be maintained for all members of a family living at the same address and possessing the same family name. The following information shall be recorded:

A. Name. The family name and the first name of the person for whom the medication is intended;
B. Address. The address to correspond to the name in subsection 1;
C. Age group. An indication of the patient's age group, that is, infant, child or adult;
D. **Original date of dispensing.** The original date the medication is dispensed pursuant to the receipt of a practitioner's prescription;

E. **Prescription identification.** The number or designation identifying the prescription;

F. **Prescriber's name.** The name of the person prescribing the drug or device;

G. **Drug information.** The name, strength and quantity of the drug; and

H. **Initials of pharmacist; date of refill.** The initials of the dispensing pharmacist and the date of dispensing the medication as a renewal or refill, if those initials and that date are not recorded on the back of the original prescription.

The pharmacist shall attempt to ascertain and shall record any allergies and idiosyncrasies of the patient and any chronic conditions which may relate to drug utilization as communicated to the pharmacy by the patient.

Upon receipt of a prescription, a pharmacist shall examine the patient's profile record before dispensing the medication to determine the possibility of a harmful drug interaction or reaction. Upon recognizing a potentially harmful reaction or interaction, the pharmacist shall take appropriate action to avoid or minimize the problem which may include consultation with the practitioner.

A patient profile record must be maintained for a period of not less than the amount of time required under federal Medicare laws, beginning from the date of the last entry in the profile record. As used in this section, "Medicare" means the Health Insurance for the Aged Act, Title XVIII of the Social Security Amendments of 1965, as amended.

*Under HIPAA, records are required to be maintained for a minimum period of 6 years unless another state or federal law, regulation or other legal requirement extends this period.*

32 MRSA §13785

**Photographic proof of identification**

As a precondition to filling any prescription or dispensing any drug, a pharmacist or person acting at the direction of a pharmacist may demand, inspect and record proof of identification, including valid photographic identification, from any patient presenting a prescription or any person acting on behalf of the patient or person purchasing a targeted methamphetamine precursor. Valid photographic identification includes but is not limited to the following:

A. A valid Maine motor vehicle operator's license;

B. A valid Maine identification card issued under Title 29-A, section 1410;

C. A valid United States passport;

D. A valid passport or motor vehicle operator's license of another state, territory, possession or foreign country only if it:
   1. Contains a photograph of the traveler or licensee;
   2. Is encased in tamper-resistant plastic or otherwise possesses indicia of tamper-resistance; and
   3. Identifies the traveler's or licensee's date of birth; or

E. Other valid, tamper-resistant, photographic identification as provided in rules adopted by the board pursuant to section 13722, subsection 1, paragraph A and in accordance with Title 5, chapter 375.

32 MRSA §13795 (1)

**Refusal to fill prescription or dispense drug**

A pharmacist or person acting at the direction of a pharmacist may exercise discretion and refuse to fill any prescription or dispense any drug if unsatisfied as to the legitimacy or appropriateness of any prescription presented, the validity of any photographic identification or the identity of any patient presenting a prescription or any person acting on behalf of the patient, or the intention of the customer to use the drug targeted methamphetamine precursor according to the instructions for use. A pharmacist or person acting at the direction of a pharmacist may make a report to a law enforcement agency when that
person has reasonable cause to suspect that a prescription is not legitimate or appropriate, that a person has presented photographic identification that is not valid or that a customer has the intention to use a drug or targeted methamphetamine precursor in a manner inconsistent with the instructions for use.

**Record keeping - Methamphetamine**

With regard to purchases of targeted methamphetamine precursors, a pharmacy may keep a log of information about the purchaser, which may include name, date of birth, address and amount of targeted methamphetamine precursors purchased.

**Security requirements**

The Department of Public Safety, after consultation with the Board of Osteopathic Licensure, the Board of Licensure in Medicine and the Board of Pharmacy, shall adopt rules that establish security requirements for all written prescriptions for schedule II drugs issued by health care providers.

**Out-of-state prescription security requirements**

Notwithstanding any law or rule to the contrary, a prescription for a schedule II drug written by an out-of-state practitioner on a prescription blank that does not comply with the requirements for a security prescription blank, as defined in the Department of Public Safety rule pursuant to subsection 1, may be filled by a pharmacist only if:

A. The pharmacist receives and makes a record of oral confirmation of the validity of the prescription from the out-of-state practitioner or the practitioner's agent and the pharmacist makes a reasonable effort to determine that the oral confirmation came from the practitioner or the practitioner's agent, which may include a telephone call to the practitioner's telephone number listed in a telephone directory or other directory or other good faith efforts to confirm the identity of the person giving the oral confirmation; and

B. The pharmacist demands, inspects and records a valid photographic identification from any person presenting a prescription or receiving a filled prescription unless:

1. The person is the patient for whom the prescription is written;
2. The person's identity is personally known to the pharmacist; and
3. The pharmacist confirms by reviewing the pharmacy records that the pharmacist has previously demanded, inspected and recorded a valid photographic identification from the person.

**Partial filling of out-of-state prescriptions**

The partial filling of a prescription for a schedule II drug written by an out-of-state practitioner on a prescription blank that does not comply with the requirements for a security prescription blank, as defined in the Department of Public Safety rule pursuant to subsection 1, is permissible if the pharmacist is unable after reasonable effort to obtain the oral confirmation described in subsection 2 in the case of the practitioner's office being closed during nights, weekends or holidays. The partial filling is limited to a 72-hour supply of the controlled substance. The remaining portion of the prescription may be filled within the 72-hour period upon obtaining the oral confirmation. No further quantity may be filled beyond the 72 hours without a new prescription.

**Sale of poisonous drugs**

Each licensed pharmacist who sells a poison shall affix to the package sold a label plainly marked with the name and address of the store and the word "POISON" and the name of the poison sold, and shall enter at the time of sale in a permanently bound book to be kept for that purpose the name and address of the purchaser, the date of sale, the name of the poison and the quantity sold and the person making the sale shall sign the entry. This section shall not apply to sales on prescription of practitioners, sales at wholesale to pharmacists or sales to hospitals, colleges or public institutions.
Using drugs not in prescription
If a pharmacist knowingly uses any drugs or ingredients in preparing or compounding a written or oral prescription of any practitioner different from those named in the prescription, that use shall constitute a civil violation for which a forfeiture of not more than $1,000 nor less than $50 may be adjudged.

Return of drugs
A drug or pharmaceutical preparation that has been dispensed on prescription may be returned to pharmacy stock after being in possession and under the control of another person and may be dispensed again if the drug is packaged in an unbroken, sealed container or if, in the case of a hospital, a licensed pharmacist determines that the drug has not been impaired.

Partial fill of opioid prescriptions
A pharmacist may, upon request by the patient, fill a prescription for opioid medication by dispensing less than the total amount prescribed. The pharmacist must report that fact to the prescriber. When a partial fill is dispensed, the remainder of the prescription is void and may not be filled at a later time without a new prescription.

Adulterating and selling drugs
Whoever fraudulently adulterates, for the purpose of sale, any drug or medicine or sells any fraudulently adulterated drug or medicine, knowing the same to be adulterated, shall be punished by a fine of not more than $1,000 or by imprisonment for not more than 11 months. These adulterated drugs and medicines shall be forfeited and destroyed under the direction of the court.

Labeling of prescriptions
Every drug dispensed pursuant to prescription, whether for a legend drug or not, must carry on the label the following information:
A. The prescription number;
B. The date of filling;
C. The patient's name;
D. Directions for use;
E. The name and strength of the drug and the amount dispensed, including either the brand name of the drug or, if a generic and therapeutically equivalent drug is dispensed it must be in accordance with section 13781;
F. The beyond use date of the drug;
G. The name of the practitioner prescribing the drug; and
H. The name, address and telephone number of the pharmacy where the prescription was compounded and dispensed.
For purposes of this section, "beyond use date" means a date beyond which the contents of the prescription are not recommended to be used.

Unused Pharmaceutical Disposal Program
The Maine Drug Enforcement Agency shall create a system for the return of unused pharmaceuticals. The system must use prepaid mailing envelopes into which the unused pharmaceuticals are placed and returned to a single collection location. The prepaid mailing envelopes must be made available to the public at various locations, including, but not limited to, pharmacies, physicians' offices and post offices. The agency may randomly assess the toxicity of materials received under the program as long as the assessment results do not identify the patient, person who mailed the material, prescriber or pharmacy.
Advertising, Marketing & Costs

Advertising
It is lawful for any pharmacy, pharmacist or other licensee of the board to advertise to the public the current retail price charged for any drugs, medicines or appliances as defined in the United States Code, Title 21, Section 3211 (g) (1) which bears the legend “Caution: Federal law prohibits dispensing without prescription.” The advertising may be according to either the brand name or the generic name of the drug. No media advertising of any drugs included in the United States Comprehensive Drug Abuse Prevention and Control Act of 1970, 84 Stat. 1236, is permitted.

Price disclosure required
A pharmacist or pharmacy technician employed by a pharmacy shall disclose upon the request of any person making an inquiry in person or by telephone the price of any brand or generic drug sold by that pharmacy.

Sale by certain methods prohibited
It shall be unlawful for any person to sell, distribute, vend or otherwise dispose of any drug, medicine or pharmaceutical or medical preparation by means of any public exhibition, entertainment, performance, carnival or by vending machines.

Maine RX Plus Program
The Maine Rx Plus Program, referred to in this subchapter as the "program," is established to reduce prescription drug prices and to improve the quality of health care for residents of the State. The program is administered by the department and must utilize manufacturer rebates and pharmacy discounts to reduce prescription drug prices.

Prescription Drug Price Reduction Act
The Legislature finds that affordability is critical in providing access to prescription drugs for Maine residents. This subchapter is enacted by the Legislature as a positive measure to make prescription drugs more affordable for qualified Maine residents, thereby increasing the overall health of Maine residents, promoting healthy communities and protecting the public health and welfare of Maine residents.

Emergency drug pricing
In order to achieve the public health purposes listed in section 2691, maximum retail prices for prescription drugs sold in Maine may be established pursuant to this section.

Profiteering in Prescription Drugs
Prescription drugs are a necessity of life. Profiteering in prescription drugs is unlawful and is subject to the provisions of this section. The provisions of this section apply to manufacturers, distributors and labelers of prescription drugs.

Marketing costs
Maine’s law requiring manufacturers to report marketing costs for prescription drugs was repealed in 2011 in an attempt to make Maine law consistent with federal law. See 2011 Public Law, Chapter 461.

Confidentiality
Prescription Privacy Law
A carrier or prescription drug information intermediary may not license, use, sell, transfer or exchange for value, for any marketing purpose, prescription drug information that identifies directly or indirectly the individual who is prescribed the prescription drug.

32 MRSA §13782
32 MRSA §13782-A (1)
32 MRSA §13792
22 MRSA §2681 (1)
22 MRSA §2691
22 MRSA §2693
22 MRSA §2697
22 MRSA §1711-E (2)
Maine’s law requiring confidentiality of prescription drug information that identifies the prescriber was repealed in 2012 in an attempt to make Maine’s law consistent with federal law. See 2011 Public Law, Chapter 494.

Prescription Monitoring Program Data Confidentiality – Office of Chief Medical Examiner
The office of Chief Medical Examiner may access prescription monitoring information for the purpose of conducting an investigation or inquiry into the cause, manner and circumstances of death in a medical examiner case as described in section 3025. Prescription monitoring information in the possession or under the control of the Office of Chief Medical Examiner is confidential and, notwithstanding section 3022, may not be disseminated. Information that is not prescription monitoring information and is separately acquired following access to prescription monitoring information pursuant to this paragraph remains subject to protection or dissemination in accordance with section 3022.

22 MRSA §7250 (4)(f)

The chief medical officer, medical director or other administrative prescriber employed by a licensed hospital, insofar as the information relates to prescriptions written by prescribers employed by that licensed hospital.

22 MRSA §7250 (4)(k)

Methadone—Consent to Report
Patients being treated with methadone for substance use disorder may consent to that treatment being reported to the Prescription Monitoring Program. The patient may refuse or withdraw consent at any time.

22 MRSA §7250 (4)

Prescription Drug Academic Detailing Program
Creation
By January 1, 2008, the department shall establish a prescription drug academic detailing program, referred to in this section as "the program," to enhance the health of residents of the State, to improve the quality of decisions regarding drug prescribing, to encourage better communication between the department and health care practitioners participating in publicly funded health programs and to reduce the health complications and unnecessary costs associated with inappropriate drug prescribing. (The program, known as the Maine Independent Clinical Information Program [MICIS], is operated by the Maine Medical Association.)

22 MRSA §2685

Program Components
Program components must include outreach and education regarding the therapeutic and cost-effective use of prescription drugs as issued in peer-reviewed scientific, medical and academic research publications and made available to prescribers and dispensers of drugs in the State, including through written information and through personal visits from program staff. To the extent possible, program components must also include information regarding clinical trials, pharmaceutical efficacy, adverse effects of drugs, evidence-based treatment options and drug marketing approaches that are intended to circumvent competition from generic and therapeutically equivalent drugs. Academic detailers shall observe standards of conduct in their educational materials and written and oral presentations as established by rules adopted by the department that are consistent with the following federal regulations regarding labeling and false and misleading advertising: the Food and Drug Administration labeling requirements of 21 Code of Federal Regulations, Part 201 (2007) and prescription drug advertising provisions of 21 Code of Federal Regulations, Part 202 (2007) and the Office of the Inspector General's Compliance Program Guidance for Pharmaceutical Manufacturers issued in April 2003, as amended. The rules must require academic detailers to disclose evidence-based information about the range and cost of appropriate drug treatment options and the health benefits and risks of all appropriate drugs.

22 MRSA §2685 (3)
Program Coverage
The program must provide outreach and education to prescribers and dispensers who participate in, contract with or are reimbursed by state-funded health care programs, including but not limited to the MaineCare program, the Maine Rx Plus Program, Dirigo Health insurance, the elderly low-cost drug program and the state employee health insurance program. The program may provide outreach and education to carriers, health plans, hospitals, employers and other persons interested in the program on a subscription or fee-paying basis under rules adopted by the department.  

22 MRSA §2685 (4)

For more information, see the Maine Medical Association website on Academic Detailing or the website for the Maine Independent Clinical Information Service (MICIS).

Controlled Substances Act, Select Sections

General
Any person planning on using any controlled substance, as listed in the USC Schedules of Controlled Substances, must register with the United States Department of Justice, Drug Enforcement Administration prior to the uses of any controlled substance.

A list of DEA Scheduled Controlled Substances along with common names and DEA numbers can be found on the DEA Website

An Exempt List of controlled substances can be found as part of the DEA’s Diversion Control Program

Persons required to register

A. Period of registration
   1. Every person who manufactures or distributes any controlled substance or list I chemical, or who proposes to engage in the manufacture or distribution of any controlled substance or list I chemical, shall obtain annually a registration issued by the Attorney General in accordance with the rules and regulations promulgated by him.
   2. Every person who dispenses, or who proposes to dispense, any controlled substance, shall obtain from the Attorney General a registration issued in accordance with the rules and regulations promulgated by him. The Attorney General shall, by regulation, determine the period of such registrations. In no event, however, shall such registrations be issued for less than one year nor for more than three years.

B. Authorized activities
   Persons registered by the Attorney General under this subchapter to manufacture, distribute, or dispense controlled substances or list I chemicals are authorized to possess, manufacture, distribute, or dispense such substances or chemicals (including any such activity in the conduct of research) to the extent authorized by their registration and in conformity with the other provisions of this subchapter.

C. Exceptions
   The following persons shall not be required to register and may lawfully possess any controlled substance or list I chemical under this subchapter:
   1. An agent or employee of any registered manufacturer, distributor, or dispenser of any controlled substance or list I chemical if such agent or employee is acting in the usual course of his business or employment.
   2. A common or contract carrier or warehouseman, or an employee thereof, whose possession of the controlled substance or list I chemical is in the usual course of his business or employment.
   3. An ultimate user who possesses such substance for a purpose specified in section 802(25) of this title.

D. Waiver
The Attorney General may, by regulation, waive the requirement for registration of certain manufacturers, distributors, or dispensers if he finds it consistent with the public health and safety.

E. Separate registration
A separate registration shall be required at each principal place of business or professional practice where the applicant manufactures, distributes, or dispenses controlled substances or list I chemicals.

F. Inspection
The Attorney General is authorized to inspect the establishment of a registrant or applicant for registration in accordance with the rules and regulations promulgated by him. 21 USCA §822

Controlled Substances Registration Materials
DEA Registration Procedures
DEA Registration Categories and Fees
DEA Registration Forms
DEA Change Request
Registration FAQ's

Schedules of Controlled Substances, Defined

A. Schedule I
   1. The drug or other substance has a high potential for abuse.
   2. The drug or other substance has no currently accepted medical use in treatment in the United States.
   3. There is a lack of accepted safety for use of the drug or other substance under medical supervision.

B. Schedule II
   1. The drug or other substance has a high potential for abuse.
   2. The drug or other substance has a currently accepted medical use in treatment in the United States or a currently accepted medical use with severe restrictions.
   3. Abuse of the drug or other substances may lead to severe psychological or physical dependence.

C. Schedule III
   1. The drug or other substance has a potential for abuse less than the drugs or other substances in schedules I and II.
   2. The drug or other substance has a currently accepted medical use in treatment in the United States.
   3. Abuse of the drug or other substance may lead to moderate or low physical dependence or high psychological dependence.

D. Schedule IV
   1. The drug or other substance has a low potential for abuse relative to the drugs or other substances in schedule III.
   2. The drug or other substance has a currently accepted medical use in treatment in the United States.
   3. Abuse of the drug or other substance may lead to limited physical dependence or psychological dependence relative to the drugs or other substances in schedule III.

E. Schedule V
   1. The drug or other substance has a low potential for abuse relative to the drugs or other substances in schedule IV.
   2. The drug or other substance has a currently accepted medical use in treatment in the United States.
3. Abuse of the drug or other substance may lead to limited physical dependence or psychological dependence relative to the drugs or other substances in schedule IV.

Distributors of controlled substances in schedule I or II
The Attorney General shall register an applicant to distribute a controlled substance in schedule I or II unless he determines that the issuance of such registration is inconsistent with the public interest. In determining the public interest, the following factors shall be considered:

A. Maintenance of effective control against diversion of particular controlled substances into other than legitimate medical, scientific, and industrial channels;
B. Compliance with applicable State and local law;
C. Prior conviction record of applicant under Federal or State laws relating to the manufacture, distribution, or dispensing of such substances;
D. Past experience in the distribution of controlled substances; and
E. Such other factors as may be relevant to and consistent with the public health and safety.

NOTE: While this section normally does not apply to medical practitioners, it is important to note that Medical Marijuana (as described under the Maine Marijuana Act) is listed as a schedule I drug.

Limits of authorized activities
Registration granted under subsections …(b) of this section shall not entitle a registrant to (1) manufacture or distribute controlled substances in schedule I or II other than those specified in the registration, or (2) manufacture any quantity of those controlled substances in excess of the quota assigned pursuant to section 826 of this title.

Distributors of controlled substances in schedule III, IV or V
The Attorney General shall register an applicant to distribute a controlled substance in schedule III, IV or V unless he determines that the issuance of such registration is inconsistent with the public interest. In determining the public interest, the following factors shall be considered:

A. Maintenance of effective control against diversion of particular controlled substances into other than legitimate medical, scientific, and industrial channels;
B. Compliance with applicable State and local law;
C. Prior conviction record of applicant under Federal or State laws relating to the manufacture, distribution, or dispensing of such substances;
D. Past experience in the distribution of controlled substances,
E. Such other factors as may be relevant to and consistent with the public health and safety.

Denial, revocation, or suspension of registration
A. Grounds
A registration pursuant to section 823 of this title to manufacture, distribute, or dispense a controlled substance or a list I chemical may be suspended or revoked by the Attorney General upon a finding that the registrant:

1. Has materially falsified any application filed pursuant to or required by this subchapter or subchapter II of this chapter;
2. Has been convicted of a felony under this subchapter or subchapter II of this chapter or any other law of the United States, or of any State, relating to any substance defined in this subchapter as a controlled substance or a list I chemical;
3. Has had his State license or registration suspended, revoked, or denied by competent State authority and is no longer authorized by State law to engage in the manufacturing, distribution, or dispensing of controlled substances or list I chemicals or has had the
suspension, revocation, or denial of his registration recommended by competent State authority;
4. Has committed such acts as would render his registration under section 823 of this title inconsistent with the public interest as determined under such section; or
5. Has been excluded (or directed to be excluded) from participation in a program pursuant to section 1320a-7(a) of title 42. A registration pursuant to section 823(g)(1) of this title to dispense a narcotic drug for maintenance treatment or detoxification treatment may be suspended or revoked by the Attorney General upon a finding that the registrant has failed to comply with any standard referred to in section 823(g)(1) of this title.

B. Limits of revocation or suspension
The Attorney General may limit revocation or suspension of a registration to the particular controlled substance or list I chemical with respect to which grounds for revocation or suspension exist.

C. Service of show cause order; proceedings
Before taking action pursuant to this section, or pursuant to a denial of registration under section 823 of this title, the Attorney General shall serve upon the applicant or registrant an order to show cause why registration should not be denied, revoked, or suspended. The order to show cause shall contain a statement of the basis thereof and shall call upon the applicant or registrant to appear before the Attorney General at a time and place stated in the order, but in no event less than thirty days after the date of receipt of the order. Proceedings to deny, revoke, or suspend shall be conducted pursuant to this section in accordance with subchapter II of chapter 5 of title 5. Such proceedings shall be independent of, and not in lieu of, criminal prosecutions or other proceedings under this subchapter or any other law of the United States.

D. Suspension of registration in cases of imminent danger
The Attorney General may, in his discretion, suspend any registration simultaneously with the institution of proceedings under this section, in cases where he finds that there is an imminent danger to the public health or safety. A failure to comply with a standard referred to in section 823(g)(1) of this title may be treated under this subsection as grounds for immediate suspension of a registration granted under such section. A suspension under this subsection shall continue in effect until the conclusion of such proceedings, including judicial review thereof, unless sooner withdrawn by the Attorney General or dissolved by a court of competent jurisdiction.

E. Suspension and revocation of quotas
The suspension or revocation of a registration under this section shall operate to suspend or revoke any quota applicable under section 826 of this title.

F. Disposition of controlled substances or list I chemicals
In the event the Attorney General suspends or revokes a registration granted under section 823 of this title, all controlled substances or list I chemicals owned or possessed by the registrant pursuant to such registration at the time of suspension or the effective date of the revocation order, as the case may be, may, in the discretion of the Attorney General, be placed under seal. No disposition may be made of any controlled substances or list I chemicals under seal until the time for taking an appeal has elapsed or until all appeals have been concluded except that a court, upon application therefor, may at any time order the sale of perishable controlled substances or list I chemicals. Any such order shall require the deposit of the proceeds of the sale with the court. Upon a revocation order becoming final, all such controlled substances or list I chemicals (or proceeds of sale deposited in court) shall be forfeited to the United States; and the Attorney General shall dispose of such controlled substances or list I chemicals in accordance with section 881(e) of this title. All right, title, and interest in such controlled substances or list I chemicals shall vest in the United States upon a revocation order becoming final.

G. Seizure or placement under seal of controlled substances or list I chemicals
The Attorney General may, in his discretion, seize or place under seal any controlled substances or list I chemicals owned or possessed by a registrant whose registration has expired or who has
ceased to practice or do business in the manner contemplated by his registration. Such controlled substances or list I chemicals shall be held for the benefit of the registrant, or his successor in interest. The Attorney General shall notify a registrant, or his successor in interest, who has any controlled substance or list I chemical seized or placed under seal of the procedures to be followed to secure the return of the controlled substance or list I chemical and the conditions under which it will be returned. The Attorney General may not dispose of any controlled substance or list I chemical seized or placed under seal under this subsection until the expiration of one hundred and eighty days from the date such substance or chemical was seized or placed under seal.

21 USCA §824

Labeling and packaging

A. Symbol
   It shall be unlawful to distribute a controlled substance in a commercial container unless such container, when and as required by regulations of the Attorney General, bears a label (as defined in section 321(k) of this title) containing an identifying symbol for such substance in accordance with such regulations. A different symbol shall be required for each schedule of controlled substances.

B. Unlawful distribution without identifying symbol
   It shall be unlawful for the manufacturer of any controlled substance to distribute such substance unless the labeling (as defined in section 321(m) of this title) of such substance contains, when and as required by regulations of the Attorney General, the identifying symbol required under subsection (a) of this section.

C. Warning on label
   The Secretary shall prescribe regulations under section 353(b) of this title which shall provide that the label of a drug listed in schedule II, III, or IV shall, when dispensed to or for a patient, contain a clear, concise warning that it is a crime to transfer the drug to any person other than the patient.

D. Containers to be securely sealed
   It shall be unlawful to distribute controlled substances in schedule I or II, and narcotic drugs in schedule III or IV, unless the bottle or other container, stopper, covering, or wrapper thereof is securely sealed as required by regulations of the Attorney General.

21 USCA §825

Records and reports of registrants

A. Inventory
   Except as provided in subsection (c) of this section -
   1. Every registrant under this subchapter shall, on May 1, 1971, or as soon thereafter as such registrant first engages in the manufacture, distribution, or dispensing of controlled substances, and every second year thereafter, make a complete and accurate record of all stocks thereof on hand, except that the regulations prescribed under this section shall permit each such biennial inventory (following the initial inventory required by this paragraph) to be prepared on such registrant's regular general physical inventory date (if any) which is nearest to and does not vary by more than six months from the biennial date that would otherwise apply;
   2. On the effective date of each regulation of the Attorney General controlling a substance that immediately prior to such date was not a controlled substance, each registrant under this subchapter manufacturing, distributing, or dispensing such substance shall make a complete and accurate record of all stocks thereof on hand; and
   3. On and after May 1, 1971, every registrant under this subchapter manufacturing, distributing, or dispensing a controlled substance or substances shall maintain, on a current basis, a complete and accurate record of each such substance manufactured,
received, sold, delivered, or otherwise disposed of by him, except that this paragraph shall not require the maintenance of a perpetual inventory.

B. Availability of records
Every inventory or other record required under this section shall be in accordance with, and contain such relevant information as may be required by, regulations of the Attorney General, be maintained separately from all other records of the registrant, or alternatively, in the case of nonnarcotic controlled substances, be in such form that information required by the Attorney General is readily retrievable from the ordinary business records of the registrant, and be kept and be available, for at least two years, for inspection and copying by officers or employees of the United States authorized by the Attorney General.

C. Nonapplicability
The foregoing provisions of this section shall not apply –
1. to the prescribing of controlled substances in schedule II, III, IV, or V by practitioners acting in the lawful course of their professional practice unless such substance is prescribed in the course of maintenance or detoxification treatment of an individual; or
2. to the administering of a controlled substance in schedule II, III, IV, or V unless the practitioner regularly engages in the dispensing or administering of controlled substances and charges his patients, either separately or together with charges for other professional services, for substances so dispensed or administered or unless such substance is administered in the course of maintenance treatment or detoxification treatment of an individual;
3. to the use of controlled substances, at establishments registered under this subchapter which keep records with respect to such substances, in research conducted in conformity with an exemption granted under section 355(i) or 360b(j) of this title;
4. to the use of controlled substances, at establishments registered under this subchapter which keep records with respect to such substances, in pre-clinical research or in teaching; or
5. to the extent of any exemption granted to any person, with respect to all or part of such provisions, by the Attorney General by or pursuant to regulation on the basis of a finding that the application of such provisions (or part thereof) to such person is not necessary for carrying out the purposes of this subchapter. Nothing in the Convention on Psychotropic Substances shall be construed as superseding or otherwise affecting the provisions of paragraph (1)(B), (2), or (3) of this subsection.

D. Periodic reports to Attorney General
Every manufacturer registered under section 823 of this title shall, at such time or times and in such form as the Attorney General may require, make periodic reports to the Attorney General of every sale, delivery or other disposal by him of any controlled substance, and each distributor shall make such reports with respect to narcotic controlled substances, identifying by the registration number assigned under this subchapter the person or establishment (unless exempt from registration under section 822(d) of this title) to whom such sale, delivery, or other disposal was made.

E. Reporting and record keeping requirements of drug conventions
In addition to the reporting and record keeping requirements under any other provision of this subchapter, each manufacturer registered under section 823 of this title shall, with respect to narcotic and non-narcotic controlled substances manufactured by it, make such reports to the Attorney General, and maintain such records, as the Attorney General may require to enable the United States to meet its obligations under articles 19 and 20 of the Single Convention on Narcotic Drugs and article 16 of the Convention on Psychotropic Substances. The Attorney General shall administer the requirements of this subsection in such a manner as to avoid the unnecessary imposition of duplicative requirements under this subchapter on manufacturers subject to the requirements of this subsection.
F. Investigational uses of drugs; procedures
   Regulations under sections 355(i) and 360(j) of this title, relating to investigational use of drugs, shall include such procedures as the Secretary, after consultation with the Attorney General, determines are necessary to insure the security and accountability of controlled substances used in research to which such regulations apply.

G. Change of address
   Every registrant under this subchapter shall be required to report any change of professional or business address in such manner as the Attorney General shall by regulation require.

H. Reporting requirements for GHB
   In the case of a drug product containing gamma hydroxybutyric acid for which an application has been approved under section 355 of this title, the Attorney General may, in addition to any other requirements that apply under this section with respect to such a drug product, establish any of the following as reporting requirements:
   1. That every person who is registered as a manufacturer of bulk or dosage form, as a packager, repackager, labeler, relabeler, or distributor shall report acquisition and distribution transactions quarterly, not later than the 15th day of the month succeeding the quarter for which the report is submitted, and annually report end-of-year inventories.
   2. That all annual inventory reports shall be filed no later than January 15 of the year following that for which the report is submitted and include data on the stocks of the drug product, drug substance, bulk drug, and dosage forms on hand as of the close of business December 31, indicating whether materials reported are in storage or in process of manufacturing.
   3. That every person who is registered as a manufacturer of bulk or dosage form shall report all manufacturing transactions both inventory increases, including purchases, transfers, and returns, and reductions from inventory, including sales, transfers, theft, destruction, and seizure, and shall provide data on material manufactured, manufactured from other material, use in manufacturing other material, and use in manufacturing dosage forms.
   4. That all reports under this section must include the registered person's registration number as well as the registration numbers, names, and other identifying information of vendors, suppliers, and customers, sufficient to allow the Attorney General to track the receipt and distribution of the drug.
   5. That each dispensing practitioner shall maintain for each prescription the name of the prescribing practitioner, the prescribing practitioner's Federal and State registration numbers, with the expiration dates of these registrations, verification that the prescribing practitioner possesses the appropriate registration to prescribe this controlled substance, the patient's name and address, the name of the patient's insurance provider and documentation by a medical practitioner licensed and registered to prescribe the drug of the patient's medical need for the drug. Such information shall be available for inspection and copying by the Attorney General.
   6. That section 830(b)(3) of this title (relating to mail order reporting) applies with respect to gamma hydroxybutyric acid to the same extent and in the same manner as such section applies with respect to the chemicals and drug products specified in subparagraph (A)(i) of such section.

Prescriptions
   A. Schedule II substances
      Except when dispensed directly by a practitioner, other than a pharmacist, to an ultimate user, no controlled substance in schedule II, which is a prescription drug as determined under the Federal Food, Drug, and Cosmetic Act (21 U.S.C. §§ 301 et seq.), may be dispensed without the written prescription of a practitioner, except that in emergency situations, as prescribed by the Secretary by regulation after consultation with the Attorney General, such drug may be dispensed upon

21 USCA §827
oral prescription in accordance with section 503(b) of that Act (21 U.S.C. § 353(b)). Prescriptions shall be retained in conformity with the requirements of section 827 of this title. No prescription for a controlled substance in schedule II may be refilled.

B. Schedule III and IV substances
Except when dispensed directly by a practitioner, other than a pharmacist, to an ultimate user, no controlled substance in schedule III or IV, which is a prescription drug as determined under the Federal Food, Drug, and Cosmetic Act (21 U.S.C. §§ 301 et seq.), may be dispensed without a written or oral prescription in conformity with section 503(b) of that Act (21 U.S.C. § 353(b)). Such prescriptions may not be filled or refilled more than six months after the date thereof or be refilled more than five times after the date of the prescription unless renewed by the practitioner.

C. Schedule V substances
No controlled substance in schedule V which is a drug may be distributed or dispensed other than for a medical purpose.

D. Non-prescription drugs with abuse potential
Whenever it appears to the Attorney General that a drug not considered to be a prescription drug under the Federal Food, Drug, and Cosmetic Act (21 U.S.C. §§ 301 et seq.) should be so considered because of its abuse potential, he shall so advise the Secretary and furnish to him all available data relevant thereto.

21 USCA §829

(37) Emergency Care and Assistance
For definitions, see 32 MRSA §83

Immunity from civil liability (Good Samaritan Law)
Notwithstanding any inconsistent provisions of any public or private and special law, any person who voluntarily, without the expectation of monetary or other compensation from the person aided or treated, renders first aid, emergency treatment or rescue assistance to a person who is unconscious, ill, injured or in need of rescue assistance, shall not be liable for damages for injuries alleged to have been sustained by such person nor for damages for the death of such person alleged to have occurred by reason of an act or omission in the rendering of such first aid, emergency treatment or rescue assistance, unless it is established that such injuries or such death were caused willfully, wantonly or recklessly or by gross negligence on the part of such person. This section shall apply to members or employees of nonprofit volunteer or governmental ambulance, rescue or emergency units, whether or not a user or service fee may be charged by the nonprofit unit or the governmental entity and whether or not the members or employees receive salaries or other compensation from the nonprofit unit or the governmental entity. This section shall not be construed to require a person who is ill or injured to be administered first aid or emergency treatment if such person objects thereto on religious grounds. This section shall not apply if such first aid or emergency treatment or assistance is rendered on the premises of a hospital or clinic.

14 MRSA §164

Asthma inhalers and epinephrine pens
Municipal employees and volunteers that operate or assist in any municipal recreational program or camp may receive training on how to administer asthma inhalers and epinephrine pens. Municipal employees and volunteers may possess and administer prescribed asthma inhalers and epinephrine pens in order to provide emergency aid.

30-A MRSA §3108

Immunity from civil liability for volunteer activities
Health care practitioners
Notwithstanding any inconsistent provision of any public or private and special law, an individual is not liable for an injury or death arising from medical services provided as described in this subsection unless
the injury or death was caused willfully, wantonly or recklessly or by gross negligence of the individual if that individual is:

A. A licensed health care practitioner who voluntarily, without the expectation or receipt of monetary or other compensation either directly or indirectly, provides professional services within the scope of that health care practitioner's licensure:
   1. To a nonprofit organization;
   2. To an agency of the State or any political subdivision of the State;
   3. To members or recipients of services of a nonprofit organization or state or local agency;
   4. To support the State's response to a public health threat as defined in Title 22, section 801, subsection 10;
   5. To support the State's response to an extreme public health emergency as defined in Title 22, section 801, subsection 4-A; or
   6. To support the State's response to a disaster as defined in Title 37-B, section 703, subsection 2; or

B. An emergency medical services' person who voluntarily, without the expectation or receipt of monetary or other compensation either directly or indirectly, provides emergency medical services within the scope of that person's licensure:
   1. To support the State's response to a public health threat as defined in Title 22, section 801, subsection 10;
   2. To support the State's response to an extreme public health emergency as defined in Title 22, section 801, subsection 4-A; or
   3. To support the State's response to a disaster as defined in Title 37-B, section 703, subsection 2.

24 MRSA §2904 (1)

Retired Physicians, Podiatrists and Dentists
Notwithstanding any inconsistent provision of any public or private and special law, a licensed physician, podiatrist or dentist who has retired from practice and who voluntarily, without the expectation or receipt of monetary or other compensation either directly or indirectly, provides professional services within the scope of that physician's, podiatrist's or dentist's licensure is not liable for an injury or death arising from those services unless the injury or death was caused willfully, wantonly or recklessly by the physician, podiatrist or dentist for professional services provided:

A. To a nonprofit organization;
B. To an agency of the State or any political subdivision of the State;
C. To members or recipients of services of a nonprofit organization or state or local agency;
D. To support the State's response to a public health threat as defined in Title 22, section 801, subsection 10;
E. To support the State's response to an extreme public health emergency as defined in Title 22, section 801, subsection 4-A; or
F. To support the State's response to a disaster as defined in Title 37-B, section 703, subsection 2.

The extended immunity under this subsection applies only if the licensed physician, podiatrist or dentist is retired from practice, possessed an unrestricted license in the relevant profession and had not been disciplined by the licensing board in the previous 5 years at the time of the act or omission causing the injury.

24 MRSA §2904 (2)

Hiring of Healthcare Personnel during Emergency Circumstances
Healthcare Workforce
A private institution is immune from civil penalties and liability for any actions arising from allegations of inadequate investigation prior to that institution's hiring or engagement of a licensed health care worker, including but not limited to allegations of negligent hiring, credentialing or privileging, for services provided within the scope of that health care worker's licensure in response to an extreme public health emergency as defined in section 801, subsection 4-A; a health emergency declared pursuant to
section 802, subsection 2; or a disaster as defined in Title 37-B, section 703, subsection 2, as long as the private institution hires or engages the services of the licensed health care worker in accordance with this subsection. When hiring or engaging the services of a health care worker:

A. The private institution shall first make a reasonable attempt to contact the appropriate occupational or professional licensing board within or affiliated with the Department of Professional and Financial Regulation for any available information about the health care worker; and

B. A private institution may rely on:
   1. Information available from the occupational and professional licensing boards within or affiliated with the Department of Professional and Financial Regulation regarding appropriate screening of the worker, such as background investigation, primary source verification or credentialing;
   2. The representation of a volunteer health care worker registry that is operated or certified in accordance with federal or state requirements regarding appropriate screening of the worker that is registered on that registry, such as background investigation, primary source verification or credentialing;
   3. The representation of the employing or privileging entity regarding appropriate screening of the worker that, at the time of hiring or engagement, is employed or privileged by any entity in any state, such as background investigation, primary source verification, credentialing or privileging; or
   4. The representation of a retired or unemployed worker’s most recent employer or privileging entity if that employment or privileging occurred within the previous 24 months.

A private institution that complies with this subsection may hire or engage the services of a licensed health care worker and is deemed in compliance with all state licensing standards. The private institution shall initiate the standard preemployment screening process within 48 hours of the official termination of the extreme public health emergency as defined in section 801, subsection 4-A or disaster as defined in Title 37-B, section 703, subsection 2.

See also, Employment During Extreme Public Health Emergency
Found in Public Health chapter of Physician’s Guide to Maine Law

Treatment at the Scene
Treatment to be in accord with regional medical orders
When an ambulance service or nontransporting emergency medical service is present at an accident or other situation in which a person or persons require emergency medical treatment, the medical treatment of the patients must be carried out in accordance with any rules adopted under this chapter, any protocols as defined in section 83, subsection 19 and any verbal orders given under the system of delegation established by the regional medical director; except that:

A. When a patient is already under the supervision of a personal physician or a physician's assistant or nurse practitioner supervised by that physician and the physician, physician's assistant or nurse practitioner assumes the care of the patient, then for as long as the physician, physician's assistant or nurse practitioner remains with the patient, the patient must be cared for as the physician, physician's assistant or nurse practitioner directs. The emergency medical services persons shall assist to the extent that their licenses and protocol allow; and

B. A patient is not required to accept treatment to which the patient does not consent.

Emergency medical persons
Basic and advanced skills
With advice from and in consultation with the Medical Direction and Practices Board, the board may provide, by rule, which skills, techniques and judgments constitute a basic emergency medical treatment.
Advanced emergency medical treatment
With the advice and consultation noted in subsection 1, the board may provide, by rule, which advanced skills, techniques and judgments may be supervised by a physician by means of standing orders, by voice radio and by other means. In every case, advanced emergency medical treatment must be given in accordance with protocols adopted by the Medical Direction and Practices Board. The board may establish by rule appropriate licensure levels for advanced emergency medical technicians and fix the qualifications for persons to hold those licenses.

Ambulance services and nontransporting medical services
Ambulance services and nontransporting medical services to be licensed
Every ambulance service and nontransporting emergency medical service must be licensed, operate in accordance with the rules adopted and protocols developed for services under this chapter and carry the equipment called for in those rules.

Care of patient
Whenever an ambulance transports a patient from the scene of an emergency, the patient must be cared for by a physician, by a flight nurse or by a person licensed under this chapter to provide emergency medical care. Whenever an ambulance transports a patient from a hospital or other health care facility to another place, the patient must be cared for by:

A. The physician in charge of the patient's case, by a person licensed under this chapter or by a professional nurse; or
B. A licensed practical nurse, or other person appropriately trained to care for the patient, acting under orders from the patient's physician.

The person specified in this subsection as caring for the patient shall accompany the patient in the portion of the ambulance where the patient rides.

Air transportation
Any patient transported by air must be flown on a service licensed under Federal Aviation Regulations, Part 135 or Part 121. In such an instance, the flight is deemed to be an air ambulance and the patient must be cared for as provided in subsection 2.

Limitation of liability for ambulance service
Except as otherwise expressly provided by statute, all governmental entities shall be immune from suit on any and all tort claims seeking recovery of damages. When immunity is removed by this chapter, any claim for damages shall be brought in accordance with the terms of this chapter.

Community Paramedicine Programs
The board may establish community paramedicine services. As used in this subsection, “community paramedicine” means the practice by an emergency medical services provider primarily in an out-of-hospital setting of providing episodic patient evaluation, advice and treatment directed at preventing or improving a particular medical condition, within the scope of practice of the emergency medical services provider as specifically requested or directed by a physician.

The board shall establish by rule the requirements and application and approval process of community paramedicine services established pursuant to this subsection. At a minimum, an emergency medical services provider, including, but not limited to, an ambulance service or nontransporting emergency medical service, that conducts community paramedicine services shall work with an identified primary care medical director, have an emergency medical services medical director and collect and submit data and written reports to the board, in accordance with requirements established by the board.
### Trauma-incidence registry

**Registry**
The board shall maintain a statewide trauma-incidence registry that meets the requirements of the federal Trauma Care Systems Planning and Development Act of 1990, Public Law 101-590, Section 1, 104 Stat. 2915. The board shall adopt rules to define trauma.

### Reporting by physicians and hospitals

Physicians and hospitals may report trauma information to the board as follows.

- **A.** A hospital may report to the board information regarding persons diagnosed as suffering from trauma. Trauma reports should be made no later than 30 days from the date of diagnosis or the date of discharge from the hospital, whichever is later.
- **B.** A physician, upon request of the board, may report to the board any further information requested by the board concerning any person now or formerly under that physician's care who was diagnosed as having suffered from trauma.
- **C.** A physician or hospital that reports in good faith in accordance with this section is not liable for any civil damages for making the report.

### Immunity for supervision and training

**Emergency medical treatment supervision**

No physician functioning within the medical control system established by the regional medical director and practicing in a hospital to or from which patients are transported under section 86 or health care practitioner under such a physician's supervision who gives oral or written instructions to a basic emergency medical services person or an advanced emergency medical technician for the provision of emergency medical treatment outside the hospital may be civilly liable for negligence as a result of issuing the instructions, if the instructions were in accordance with the protocol for the patient's reported condition. For the purpose of aiding in establishing the use of a protocol that permits the immunity provided in this subsection, the following provisions apply:

- **A.** The basic emergency medical services person or advanced emergency medical technician to whom the instructions are given shall document those instructions on the state ambulance run record; and
- **B.** The physician or health care practitioner giving the instructions shall maintain a medical control log documenting those instructions at the time they were given and shall sign the log.

The immunity provided in this subsection extends to the hospital in which the physician described in this subsection is practicing or the health care practitioner described in this subsection is being supervised.

### Emergency medical services persons' training

Except as otherwise provided in this subsection, no hospital, physician or health care practitioner providing an emergency medical services course, refresher course or continuing education course approved by Maine Emergency Medical Services may be vicariously liable for the civil liability of a person enrolled in the course to a person receiving emergency medical treatment during the course.

The immunity provided by this subsection does not apply if the person enrolled in the course is an employee of the hospital, physician or health care practitioner seeking immunity under this subsection.

### Emergency Medical Treatment and Active Labor Act (EMTALA)

**Definitions**

- **A.** Emergency medical condition: means
1. A medical condition manifesting itself by acute symptoms of sufficient severity (including severe pain) such that the absence of immediate medical attention could reasonably be expected to result in –
   a. placing the health of the individual (or, with respect to a pregnant woman, the health of the woman or her unborn child) in serious jeopardy,
   b. serious impairment to bodily functions, or
   c. serious dysfunction of any bodily organ or part; or
   d. With respect to a pregnant woman who is having contractions –
      i. that there is inadequate time to effect a safe transfer to another hospital before delivery, or
      ii. that transfer may pose a threat to the health or safety of the woman or the unborn child.

B. Participating hospital: means hospital that has entered into a provider agreement under section 1395cc of this title.

C. To stabilize: means, with respect to an emergency medical condition described in paragraph (1)(A), to provide such medical treatment of the condition as may be necessary to assure, within reasonable medical probability, that no material deterioration of the condition is likely to result from or occur during the transfer of the individual from a facility, or, with respect to an emergency medical condition described in paragraph (1)(B), to deliver (including the placenta).

D. Stabilized: means, with respect to an emergency medical condition described in paragraph (1)(A), that no material deterioration of the condition is likely, within reasonable medical probability, to result from or occur during the transfer of the individual from a facility, or, with respect to an emergency medical condition described in paragraph (1)(B), that the woman has delivered (including the placenta).

E. Transfer: means the movement (including the discharge) of an individual outside a hospital's facilities at the direction of any person employed by (or affiliated or associated, directly or indirectly, with) the hospital, but does not include such a movement of an individual who (A) has been declared dead, or (B) leaves the facility without the permission of any such person.

F. Hospital: includes a rural primary care hospital (as defined in section 1395x(mm)(1) of this title).

G. Comes to the emergency department: means, with respect to an individual who is not a patient, the individual—
   1. Has presented at a hospital's dedicated emergency department, as defined in this section, and requests examination or treatment for a medical condition, or has such a request made on his or her behalf. In the absence of such a request by or on behalf of the individual, a request on behalf of the individual will be considered to exist if a prudent layperson observer would believe, based on the individual's appearance or behavior, that the individual needs examination or treatment for a medical condition;
   2. Has presented on hospital property, as defined in this section, other than the dedicated emergency department, and requests examination or treatment for what may be an emergency medical condition, or has such a request made on his or her behalf (except for certain outpatients as specified in paragraph (d)(3) of this section). In the absence of such a request by or on behalf of the individual, a request on behalf of the individual will be considered to exist if a prudent layperson observer would believe, based on the individual's appearance or behavior, that the individual needs emergency examination or treatment;
   3. Is in a ground or air ambulance owned and operated by the hospital for presentation for examination and treatment for a medical condition at a hospital's dedicated emergency department, even if the ambulance is not on hospital grounds. This provision does not apply: (1) If the ambulance is operating under communitywide EMS protocols that direct
it to transport the individual to a hospital other than the hospital that owns the ambulance; for example, to the nearest hospital. In this latter case, the individual is considered to have come to the emergency department of the hospital to which the individual is transported, at the time the individual is brought onto hospital property; or (2) the ambulance is operated at the direction of a physician who is not employed or otherwise affiliated with the hospital that owns the ambulance.

4. Is in a nonhospital-owned ambulance on hospital property for presentation for examination and treatment for a medical condition at a hospital's dedicated emergency department. An individual in a nonhospital-owned ambulance off hospital property is not considered to have come to the hospital's emergency department, even if a member of the ambulance staff contacts the hospital by telephone or telemetry communications and informs the hospital that they want to transport the individual to the hospital for examination and treatment. In the latter circumstance, the hospital may deny access if it is in "diversionary status," that is, it does not have the staff or facilities to accept any additional emergency patients. If, however, the ambulance staff disregards the hospital's instructions and transports the individual onto hospital property, the individual is considered to have come to the emergency department.

H. **Dedicated emergency department:** means any department or facility of the hospital, regardless of whether it is located on or off the main hospital campus, that meets at least one of the following requirements:
   1. It is licensed by the State in which it is located under applicable State law as an emergency room or emergency department;
   2. It is held out to the public (by name, posted signs, advertising, or other means) as a place that provides care for emergency medical conditions on an urgent basis without requiring a previously scheduled appointment; or
   3. During the calendar year immediately preceding the calendar year in which a determination under this section is being made, based on a representative sample of patient visits that occurred during that calendar year, it provides at least one-third of all of its outpatient visits for the treatment of emergency medical conditions on an urgent basis without requiring a previously scheduled appointment.

I. **Hospital property:** means the entire main hospital campus as defined in Sec. 413.65(b) of this chapter, including the parking lot, sidewalk, and driveway, but excluding other areas or structures of the hospital's main building that are not part of the hospital, such as physician offices, rural health centers, skilled nursing facilities, or other entities that participate separately under Medicare, or restaurants, shops, or other nonmedical facilities.

J. **Hospital with an emergency department:** means a hospital with a dedicated emergency department (as defined in this paragraph (b)).

K. **Patient:** means –
   1. An individual who has begun to receive outpatient services as part of an encounter, as defined in Section 410.2 of this chapter, other than an encounter that the hospital is obliged by this section to provide;
   2. An individual who has been admitted as an inpatient, as defined by this section.

**Medical screening requirement**

In the case of a hospital that has a hospital emergency department, if any individual (whether or not eligible for benefits under this subchapter) comes to the emergency department and a request is made on the individual's behalf for examination or treatment for a medical condition, the hospital must provide for an appropriate medical screening examination within the capability of the hospital's emergency department, including ancillary services routinely available to the emergency department, to determine whether or not an emergency medical condition (within the meaning of subsection (e)(1) of this section) exists.
Necessary stabilizing treatment for emergency medical conditions and labor

A. In general
If any individual (whether or not eligible for benefits under this subchapter) comes to a hospital and the hospital determines that the individual has an emergency medical condition, the hospital must provide either -
1. within the staff and facilities available at the hospital, for such further medical examination and such treatment as may be required to stabilize the medical condition, or
2. for transfer of the individual to another medical facility in accordance with subsection (c) of this section.

B. Refusal to consent to treatment
A hospital is deemed to meet the requirement of paragraph (1)(A) with respect to an individual if the hospital offers the individual the further medical examination and treatment described in that paragraph and informs the individual (or a person acting on the individual's behalf) of the risks and benefits to the individual of such examination and treatment, but the individual (or a person acting on the individual's behalf) refuses to consent to the examination and treatment. The hospital shall take all reasonable steps to secure the individual's (or person's) written informed consent to refuse such examination and treatment.

C. Refusal to consent to transfer
A hospital is deemed to meet the requirement of paragraph (1) with respect to an individual if the hospital offers to transfer the individual to another medical facility in accordance with subsection (c) of this section and informs the individual (or a person acting on the individual's behalf) of the risks and benefits to the individual of such transfer, but the individual (or a person acting on the individual's behalf) refuses to consent to the transfer. The hospital shall take all reasonable steps to secure the individual's (or person's) written informed consent to refuse such transfer.

Restricting transfers until individual stabilized

A. Rule
If an individual at a hospital has an emergency medical condition which has not been stabilized (within the meaning of subsection (c)(3)(B) of this section), the hospital may not transfer the individual unless -
1. the individual (or a legally responsible person acting on the individual's behalf) after being informed of the hospital's obligations under this section and of the risk of transfer, in writing requests transfer to another medical facility,
2. a physician (within the meaning of section 1395x(r)(1) of this title) has signed a certification that based upon the information available at the time of transfer, the medical benefits reasonably expected from the provision of appropriate medical treatment at another medical facility outweigh the increased risks to the individual and, in the case of labor, to the unborn child from effecting the transfer, or
3. if a physician is not physically present in the emergency department at the time an individual is transferred, a qualified medical person (as defined by the Secretary in regulations) has signed a certification described in clause (ii) after a physician (as defined in section 1395x(r)(1) of this title), in consultation with the person, has made the determination described in such clause, and subsequently countersigns the certification; and
4. the transfer is an appropriate transfer (within the meaning of paragraph (2)) to that facility.

A certification described in clause (ii) or (iii) of subparagraph (A) shall include a summary of the risks and benefits upon which the certification is based.

B. Appropriate transfer
An appropriate transfer to a medical facility is a transfer -
1. in which the transferring hospital provides the medical treatment within its capacity which minimizes the risks to the individual's health and, in the case of a woman in labor, the health of the unborn child;
2. in which the receiving facility -
   b. has available space and qualified personnel for the treatment of the individual, and
   c. has agreed to accept transfer of the individual and to provide appropriate medical treatment;
3. in which the transferring hospital sends to the receiving facility all medical records (or copies thereof), related to the emergency condition for which the individual has presented, available at the time of the transfer, including records related to the individual's emergency medical condition, observations of signs or symptoms, preliminary diagnosis, treatment provided, results of any tests and the informed written consent or certification (or copy thereof) provided under paragraph (1)(A), and the name and address of any on-call physician (described in subsection (d)(1)(C) of this section) who has refused or failed to appear within a reasonable time to provide necessary stabilizing treatment;
4. in which the transfer is effected through qualified personnel and transportation equipment, as required including the use of necessary and medically appropriate life support measures during the transfer; and
5. which meets such other requirements as the Secretary may find necessary in the interest of the health and safety of individuals transferred.

Preemption
The provisions of this section do not preempt any State or local law requirement, except to the extent that the requirement directly conflicts with a requirement of this section.

Nondiscrimination
A participating hospital that has specialized capabilities or facilities (such as burn units, shock-trauma units, neonatal intensive care units, or (with respect to rural areas) regional referral centers as identified by the Secretary in regulation) shall not refuse to accept an appropriate transfer of an individual who requires such specialized capabilities or facilities if the hospital has the capacity to treat the individual.

No delay in examination or treatment
A participating hospital may not delay provision of an appropriate medical screening examination required under subsection (a) of this section or further medical examination and treatment required under subsection (b) of this section in order to inquire about the individual's method of payment or insurance status.

Whistleblower protections
A participating hospital may not penalize or take adverse action against a qualified medical person described in subsection (c)(1)(A)(iii) of this section or a physician because the person or physician refuses to authorize the transfer of an individual with an emergency medical condition that has not been stabilized or against any hospital employee because the employee reports a violation of a requirement of this section.
Special responsibilities of Medicare hospitals in emergency cases

Applicability

In the case of a hospital that has an emergency department, if an individual (whether or not eligible for Medicare benefits and regardless of ability to pay) "comes to the emergency department", as defined in paragraph (b) of this section, the hospital must--

A. Provide an appropriate medical screening examination within the capability of the hospital's emergency department, including ancillary services routinely available to the emergency department, to determine whether or not an emergency medical condition exists. The examination must be conducted by an individual(s) determined qualified by hospital bylaws or rules and regulations and who meet the requirements of Sec. 482.55 of this chapter concerning emergency services personnel and direction; and

B. If an emergency medical condition is determined to exist, provide any necessary stabilizing treatment, as defined in paragraph (d) of this section, or an appropriate transfer as defined in paragraph (e) of this section. If the hospital admits the individual as an inpatient for further treatment, the hospital's obligation under this section ends as specified by paragraph (d)(2) of this section.

42 CFR §489.24 (a)

Use of dedicated emergency department for nonemergency services

If an individual comes to a hospital's dedicated emergency department and a request is made on his or her behalf for examination or treatment for a medical condition, but the nature of the request makes it clear that the medical condition is not of an emergency nature, the hospital is required only to perform such screening as would be appropriate for any individual presenting in that manner, to determine that the individual does not have an emergency medical condition.

42 CFR §489.24 (c)

Necessary stabilizing treatment for emergency medical conditions

A. General.

If any individual (whether or not eligible for Medicare benefits) comes to a hospital and the hospital determines that the individual has an emergency medical condition, the hospital must provide either—

1. Within the capabilities of the staff and facilities available at the hospital, for further medical examination and treatment as required to stabilize the medical condition; or
2. For transfer of the individual to another medical facility in accordance with paragraph (e) of this section.

B. Application to inpatients--admitted emergency patients.

1. When an individual has been screened under paragraph (a) of this section and found to have an emergency medical condition, and the hospital admits the patient as an inpatient in good faith in order to stabilize the emergency medical condition, the hospital is relieved of further responsibility to the individual under this section.

2. A hospital has no responsibility under this section with respect to an inpatient who was admitted for elective (nonemergency) diagnosis or treatment.

3. A hospital is required by the conditions of participation for hospitals under part 482 of this chapter to provide care to its inpatients in accordance with those conditions of participation.

C. Refusal to consent to treatment

A hospital meets the requirements of paragraph (d)(1)(i) of this section with respect to an individual if the hospital offers the individual the further medical examination and treatment described in that paragraph and informs the individual (or a person acting on the individual's behalf) of the risks and benefits to the individual of the examination and treatment, but the individual (or a person acting on the individual's behalf) refuses to consent to the examination and treatment. The medical record must contain a description of the examination, treatment, or both if applicable, that was refused by or on behalf of the individual. The hospital must take all
reasonable steps to secure the individual's written informed refusal (or that of the person acting on his or her behalf). The written document should indicate that the person has been informed of the risks and benefits of the examination or treatment, or both.

D. **Delay in examination or treatment**

1. A participating hospital may not delay providing an appropriate medical screening examination required under paragraph (a) of this section or further medical examination and treatment required under paragraph (d)(1) of this section in order to inquire about the individual's method of payment or insurance status.

2. A participating hospital may not seek, or direct a patient to seek, authorization from the individual's insurance company for screening or stabilization services to an individual until after the hospital has provided the appropriate medical screening examination required under paragraph (a) of this section, and initiated any further medical examination and treatment that may be required to stabilize the emergency medical condition under paragraph (d)(1) of this section.

3. An emergency physician is not precluded from contacting the patient's physician at any time to seek advice regarding the patient's medical history and needs that may be relevant to the medical treatment and screening of the patient, as long as this consultation does not inappropriately delay services required under paragraph (a) or paragraph (d)(1) of this section.

4. Hospitals may follow reasonable registration processes for individuals for whom examination or treatment is required by this section, including asking whether an individual is insured and, if so, what that insurance is, as long as that inquiry does not delay screening or treatment. Reasonable registration processes may not unduly discourage individuals from remaining for further evaluation.

E. **Refusal to consent to transfer**

A hospital meets the requirements of paragraph (d)(1)(ii) of this section with respect to an individual if the hospital offers to transfer the individual to another medical facility in accordance with paragraph (e) of this section and informs the individual (or a person acting on his or her behalf) of the risks and benefits to the individual of the transfer, but the individual (or a person acting on the individual's behalf) refuses to consent to the transfer. The hospital must take all reasonable steps to secure the individual's written informed refusal (or that of a person acting on his or her behalf). The written document must indicate the person has been informed of the risks and benefits of the transfer and state the reasons for the individual's refusal. The medical record must contain a description of the proposed transfer that was refused by or on behalf of the individual.

42 CFR §489.24 (d)

F. **Availability of on-call physicians**

Each hospital must maintain an on-call list of physicians on its medical staff in a manner that best meets the needs of the hospital's patients. Physicians, including specialists and subspecialists, are not required to be on call at all times. The hospital must have written policies and procedures in place to respond to situations in which a particular specialty is not available or the on-call physician cannot respond because of circumstances beyond the physician's control.

42 CFR §489.24 (j)

(38) **Ethics, Code of Medical Ethics of the American Medical Association**

See: https://www.ama-assn.org/delivering-care/ama-code-medical-ethics

(39) **Experimental Treatment**

(see Treatment of Terminally Ill Patients, Experimental/ Investigational)
(40) Fees and Charges
See AMA Opinions
AMA Code of Medical Ethics Chapter 11.00, Opinions on Financing and Delivery of Health Care
AMA Policy Finder

(41) Genetic Information and Use Thereof
See AMA Opinions
AMA Code of Medical Ethics, Chapter 4.0, Opinions on Genetics and Reproductive Medicine; Chapter 7.0, Opinions on Research and Innovation
AMA Policy Finder

Maine Laws

Definitions
A. Genetic characteristic: means any inherited gene or chromosome, or alteration of a gene or chromosome that is scientifically or medically believed to predispose an individual to a disease, disorder or syndrome or to be associated with a statistically significant increased risk of development of a disease, disorder or syndrome.

B. Genetic information: means the information concerning genes, gene products or inherited characteristics that may be obtained from an individual or family member.

C. Genetic test: means a test for determining the presence or absence of an inherited genetic characteristic in an individual, including tests of nucleic acids such as deoxyribonucleic acid, or DNA, ribonucleic acid, or RNA, or mitochondrial DNA, and tests of chromosomes or proteins in order to identify a predisposing genetic characteristic.

5 MRSA §19301
Employment discrimination prohibited
An employer may not fail or refuse to hire, discharge or otherwise discriminate against an employee or applicant for employment with respect to the compensation, terms or conditions of employment on the basis of genetic information concerning that individual or because of the individual's refusal to submit to a genetic test or make available the results of a genetic test or on the basis that the individual received a genetic test or genetic counseling, except when based on a bona fide occupational qualification.

5 MRSA §19302
Discrimination in health, hospital and dental insurance
An insurer, nonprofit hospital and medical service organization or health maintenance organization that issues individual or group hospital, health or dental insurance may not discriminate against an individual or eligible dependent on the basis of genetic information or the refusal to submit to a genetic test or make available the results of a genetic test or on the basis that the individual or eligible dependent received a genetic test or genetic counseling in the issuance, withholding, extension or renewal of any hospital confinement or other health insurance, as defined by the superintendent, by rule, or in the fixing of the rates, terms or conditions for insurance, or in the issuance or acceptance of any application for insurance. This subsection does not apply to accidental injury, specified disease, hospital indemnity, disability, long-term care and other limited benefit health insurance policies and contracts.

24-A MRSA §2159-C
Confidentiality
All of the state and federal, statutes, rules and regulations governing confidentiality, use of medical information, disclosures, authorizations and mandatory reports apply to the use of genetic information. Please see the sections on Confidentiality, Consent to Treatment, HIPAA, Medical Records, and Mandatory Reports for more information.
(42) Gifts

Prohibition
Pharmaceutical and device manufacturers and wholesalers are prohibited from offering or giving to health care practitioners cash gifts in any amount or gifts for which reciprocity is expected or implied. There are exceptions for:

1. Noncash items of minimal value that will directly benefit patients, such as drug samples, educational materials, and “modest” meals and refreshments in connection with presentations about risks, benefits and appropriate uses of prescription drugs or medical devices;
2. Funding to academic institutions and residency and fellowship programs, so long as the funding sources are identified; or
3. Reasonable honoraria and reimbursement of reasonable expenses for professional or educational conferences or meetings.

32 MRSA §13759

(43) Good Samaritan Law (see Emergency Care and Assistance)

(44) Health Care Pricing

Price Transparency Requirements
A health care entity shall have available to patients the prices of the health care entity's most frequently provided health care services and procedures. The prices stated must be the prices that the health care entity charges patients directly when there is no insurance coverage for the services or procedures or when reimbursement by an insurance company is denied. The prices stated must be accompanied by descriptions of the services and procedures and the applicable standard medical codes or current procedural technology codes used by the American Medical Association.

A health care entity shall inform patients about the availability of prices for the most frequently provided health care services and procedures.

A health care entity shall prominently display, in a location that is readily accessible to patients, information on the price transparency tools available from the publicly accessible website of the Maine Health Data Organization established pursuant to chapter 1683 to assist consumers with obtaining estimates of costs associated with health care services and procedures.

A health care entity that does not routinely render services directly to patients in an office setting may satisfy this subsection by providing the information on its publicly accessible website.

A health care entity making a referral or recommendation for a comparable health care service (24-A MRSA §4318-A) must give the patient written notice of the right to receive services from a different provider. The entity shall comply with this paragraph by providing a written notice at the time the health care entity recommends or refers a patient for a health care service or procedure that may qualify as a comparable health care service. A written notice provided under this paragraph must include a notification that, prior to obtaining the recommended service, the patient may review the health care price transparency tool provided by the patient's carrier or contact the patient's carrier directly via a toll-free telephone number so that the patient may consider whether the recommended provider of the comparable health care service represents the best value for the patient. A written notice provided under this paragraph must also include a description of the service or the applicable standard medical codes or current procedural terminology codes used by the American Medical Association sufficient to allow the carrier to assist the patient in comparing prices for the comparable health care service.
**Prohibition on balance billing for “surprise” bills**

"Surprise bill" means a bill for health care services, other than emergency services, received by an enrollee for covered services rendered by an out-of-network provider, when such services were rendered by that out-of-network provider at a network provider, during a service or procedure performed by a network provider or during a service or procedure previously approved or authorized by the carrier and the enrollee did not knowingly elect to obtain such services from that out-of-network provider. "Surprise bill" does not include a bill for health care services received by an enrollee when a network provider was available to render the services and the enrollee knowingly elected to obtain the services from another provider who was an out-of-network provider.

With respect to a surprise bill:

A. A carrier shall require an enrollee to pay only the applicable coinsurance, copayment, deductible or other out-of-pocket expense that would be imposed for health care services if the services were rendered by a network provider;

B. A carrier shall reimburse the out-of-network provider or enrollee, as applicable, for health care services rendered at the average network rate under the enrollee's health care plan as payment in full, unless the carrier and out-of-network provider agree otherwise; and

C. Notwithstanding paragraph B, if a carrier has an inadequate network, as determined by the superintendent, the carrier shall ensure that the enrollee obtains the covered service at no greater cost to the enrollee than if the service were obtained from a network provider or shall make other arrangements acceptable to the superintendent.

An out-of-network provider reimbursed for a surprise bill under Title 24-A, section 4303-C, subsection 2, paragraph B may not bill an enrollee for health care services beyond the applicable coinsurance, copayment, deductible or other out-of-pocket cost expense that would be imposed for the health care services if the services were rendered by a network provider under the enrollee's health plan.

**Uninsured Patients**

On request of an uninsured patient, a health care entity as defined in section 1718-B shall provide, within a reasonable time, an estimate of the total price of medical services to be rendered directly by that health care entity during a single medical encounter. If the health care entity is unable to provide an accurate estimate because the amount of the medical services to be consumed is unknown in advance, the health care entity shall provide a brief description of the basis for determining the total price of that particular medical service. If a single medical encounter will involve medical services to be rendered by one or more 3rd party health care entities, the health care entity shall identify each 3rd party health care entity to enable the uninsured patient to seek an estimate of the total price of medical services to be rendered directly by each health care entity to that patient. When providing an estimate as required by this section, a health care entity must notify the uninsured patient of any charity care policies adopted by the health care entity.

**Itemized bills – Hospitals**

Each hospital licensed by the State under chapter 405 shall inform all patients, or their legal guardians, in writing, at the time of the patient's discharge, that it will provide an itemized bill upon their request.
The request may be made by the patient or his legal guardian at discharge or at any time within 7 years after discharge.

The hospital shall provide an itemized bill to the person making the request within 30 days of the request. Notwithstanding this section, effective July 1, 1985, each hospital shall itemize on the hospital bill of each patient the cost of nursing services provided to that patient.

**22 MRSA §1712**

**Prohibition on payment for health care facility mistakes or preventable adverse events**

**Definitions**

A. **Health Care Facility**: means a hospital or ambulatory surgical center licensed under chapter 405.

B. **Mistake or preventable adverse event**: see **22 MRSA §1721** for full list of events within the health care facility’s control to avoid.

**22 MRSA §1721 (1)**

**Prohibition**

A health care facility is prohibited from knowingly charging a patient or the patient’s insurer or the patient’s employer as defined in Title 39-A, section 102, subsection 12 for health care services it provided as a result of or to correct a mistake or preventable adverse event caused by that health care facility.

**22 MRSA §1721 (2)**

**Patient education**

A health care facility is required to inform patients of the prohibition on payment for health care facility mistakes or preventable adverse events.

**22 MRSA §1721 (3)**

**Financial disclosure – Hospitals**

Each hospital licensed under this chapter must annually publicly disclose:

A. **IRS Form 990**: The federal Internal Revenue Service Form 990, including all related disclosable schedules, for the hospital and for each tax-exempt entity related to the hospital that is required by federal law to file that form with the Internal Revenue Service; and

B. **IRS Form 1120**: The federal Internal Revenue Service Form 1120 for each for-profit entity in which the hospital has a controlling interest.

Information required to be disclosed under this section must be submitted by the hospital to the department within 5 months after the end of the hospital's fiscal year or within 5 months after the date on which the entity files the applicable form with the Internal Revenue Service. The department shall make available for public inspection and photocopying copies of all documents required by this section and shall post those documents on the department's publicly accessible website.

**22 MRSA §1819-A**

(45) **Health Insurance Portability and Accountability Act of 1996 (HIPAA)**

In plain English, HIPAA allows people to maintain their health insurance coverage (the continuity part) while allowing them the ability to change jobs or health plans (the portability part). In addition, HIPAA provides certain protections around a person’s protected health information and limits who has access to that information and how it can be transmitted, mailed, e-mailed etc.
Administrative Simplification
Main sections
Of all the sections of HIPAA, the Administrative Simplification section affects the largest portion of the healthcare industry. This section is composed of four main parts plus an enforcement component. These provisions are:

A. Electronic Health Transactions Standards (Code Sets)
B. Unique identifiers for providers, employers, and health plans
C. Security of health information and electronic signature standards (the Security Rule)
D. Privacy and confidentiality standards (the Privacy Rule)
E. Enforcement Provisions

View the Combined Regulation Text of HIPAA, Unofficial Version

HITECH Amendments
Congress passed the Health Information Technology for Economic and Clinical Health (HITECH) Act in 2009 as part of the American Recovery and Reinvestment Act. HITECH amended and added to HIPAA, including new civil money penalties for HIPAA Privacy and Security Rule violations and breach notification obligations.

Final Omnibus HIPAA Rule
A final updated Omnibus HIPAA rule was released by the federal government on January 17, 2013. You can view a copy of the Final Omnibus Rule here. As promised, the omnibus rule embodied four final rules:

• modifications to the HIPAA Privacy and Security rules mandated in the Health Information for Economic and Clinical Health (HITECH) Act;
• changes to the HIPAA enforcement rule;
• final regulations concerning reporting of data breaches; and
• modifications to the Privacy Rule as required in the Genetic Information Nondiscrimination Act (GINA).

The rule became effective March 26, 2013.

Enforcement
The Office of Civil Rights will handle enforcement of HIPAA violations. Penalties for HIPAA violations range from monetary fines to imprisonment, based on the nature and severity of the violation.

For more information on the HIPAA enforcement provisions, please go to the Office of Civil Rights Enforcement Page.

Useful Resources
U.S. DHHS Office of Civil Rights
HIPAA Frequently Asked Questions
How to File a Complaint
AMA HIPAA Compliance Resources

For more information about complying with HIPAA, including sample forms and trainings, contact the Maine Medical Association at 622-3374 or visit http://www.mainemed.com/education-info-cme/hipaa
(46) Hospital and Health Care Provider Cooperation Act

Legislative findings and intent
The Legislature finds that it is necessary and appropriate to encourage hospitals and other health care
providers to cooperate and enter into agreements that will facilitate cost containment, improve quality of
care and increase access to health care services. This Act provides processes for state review of overall
public benefit, for approval through certificates of public advantage and for continuing supervision. It is
the intent of the Legislature that a certificate of public advantage approved under this chapter provide
state action immunity under applicable federal antitrust laws.

22 MRSA §1842

For the full Hospital and Health Care Provider Cooperation Act, see 22 MRSA Chapter 405-A.
For more information on filing and maintaining certificates of public advantage, see 22 MRSA §1844.
For more information on the continuing supervision of holders of certificates of public advantage and
revocation of a certificates, see 22 MRSA §1845.

(47) Hypodermic Apparatus, Sale of

Authorized seller
A hypodermic apparatus, as defined in Title 17-A, section 1101, subsection 2, may be sold only by a
manufacturer or dealer of embalming supplies, manufacturer or dealer of medical or dental supplies,
wholesale druggist, manufacturing pharmacist, pharmacist, veterinarian, agricultural supply store or
manufacturer of surgical instruments.

32 MRSA §13787-A (1)

Purchaser
Any person who is 18 years of age or older may purchase a hypodermic apparatus from a seller described
in subsection 1.

32 MRSA §13787-A (2)

Criminal immunity
Immunity from criminal prosecution is governed by the following:
A. A seller described in subsection 1 is "expressly authorized" within the meaning of Title 17-A,
section 1110, subsection 1, paragraph A.
B. A seller described in subsection 1 or a purchaser described in subsection 2 is "expressly
authorized" within the meaning of Title 17-A, section 1111, subsection 1, paragraph A.

32 MRSA §13787-A (3)

Immunity limited
This section does not limit prosecution for violation of any law prohibiting or regulating the use,
possession, dispensing, distribution or promotion of controlled substances, scheduled drugs or drug
paraphernalia.

32 MRSA §13787-A (4)

Medicaid not affected
This section does not diminish, expand or otherwise affect Medicaid reimbursement for hypodermic
apparatuses.

32 MRSA §13787-A (5)

(48) Hysterectomy

According to federal regulations:

Additional condition for Federal Financial Participation (FFP)
A. FFP is not available in expenditures for any sterilization or hysterectomy unless the Medicaid
agency, before making payment, obtained documentation showing that the requirements of this
subpart were met. This documentation must include a consent from, an acknowledgement of
receipt of hysterectomy information or a physician's certification under Sec. 441.255(d)(2), as applicable.

(49) Immunization

Immunization Requirements
For immunization requirements for health care employees, providers and facilities, as well as day care, school, college and campground entry, see the Maine Immunization Program website.

Universal Childhood Immunization Program
The Universal Childhood Immunization Program was established by 2009 Public Law 595 to provide all children from birth until 19 years of age in the State with access to a uniform set of vaccines as determined and periodically updated by the Maine Vaccine Board. Vaccine purchase is funded by an assessment on health insurance carriers and 3rd-party administrators registered under Title 24-A of the Maine Revised Statutes. For information about the Maine Universal Immunization program see the Maine Vaccine Board Website. See also 22 MRSA §1066, establishing the program.

Pharmacist Administration (see Pharmacy Practice)

(50) Impaired Physician, Reporting of

Drug and Alcohol Misuse Reporting
Every state professional society shall establish a professional competence committee of its members pursuant to written bylaws approved by the society's governing board. The committee shall receive, investigate and determine the accuracy of any report made to the society of any member physician's acts amounting to gross or repeated medical malpractice, misuse of alcohol, drugs or other substances that may result in the member physician's performing services in a manner that endangers the health or safety of patients or professional.

Any professional competence committee within this State and any physician licensed to practice or otherwise lawfully practicing within this State shall, and any other person may, report the relevant facts to the appropriate board relating to the acts of any physician in this State if, in the opinion of the committee, physician or other person, the committee or individual has reasonable knowledge of acts of the physician amounting to gross or repeated medical malpractice, misuse of alcohol, drugs or other substances that may result in the physician's performing services in a manner that endangers the health or safety of patients, professional incompetence, unprofessional conduct or sexual misconduct identified by board rule. The failure of any such professional competence committee or any such physician to report as required is a civil violation for which a fine of not more than $1,000 may be adjudged.

Except for specific protocols developed by a board pursuant to Title 32, section 1073, 2596-A or 3298, a physician, dentist or committee is not responsible for reporting misuse of alcohol or, drugs or other substances or professional incompetence or malpractice as a result of physical or mental infirmity or by the misuse of alcohol or, drugs or other substances discovered by the physician, dentist or committee as a result of participation or membership in a professional review committee or with respect to any information acquired concerning misuse of alcohol or, drugs or other substances or professional incompetence or malpractice as a result of physical or mental infirmity or by the misuse of alcohol or, drugs or other substances, as long as that information is reported to the professional review committee.

Nothing in this section may prohibit an impaired physician or dentist from seeking alternative forms of treatment.

24 MRSA §2504
24 MRSA §2505
Provider, entity and carrier reports

A health care provider or health care entity shall, within 60 days, report in writing to the disciplined practitioner's board or authority the name of any licensed, certified or registered employee or person privileged by the provider or entity whose employment or privileges have been revoked, suspended, limited or terminated or who resigned while under investigation or to avoid investigation for reasons related to clinical competence or unprofessional conduct, together with pertinent information relating to that action. Pertinent information includes a description of the adverse action, the name of the practitioner involved, the date, the location and a description of the event or events giving rise to the adverse action, identification of the complainant and the medical records involved. Upon request, the following information must be released to the board or authority within 20 days of receipt of the request: medical records relating to the event or events; written statements signed or prepared by any witness or complainant to the event; and related correspondence between the practitioner and the provider or entity. The report must include situations in which employment or privileges have been revoked, suspended, limited or otherwise adversely affected by action of the health care practitioner while the health care practitioner was the subject of disciplinary proceedings, and it also must include situations where employment or privileges have been revoked, suspended, limited or otherwise adversely affected by act of the health care practitioner in return for the health care provider or health care entity terminating such proceeding. Any reversal, modification or change of action reported pursuant to this section must be reported immediately to the practitioner's board or authority, together with a brief statement of the reasons for that reversal, modification or change. The failure of any health care provider or health care entity to report as required is a civil violation for which a fine of not more than $5,000 may be adjudged. Carriers providing managed care plans are subject to the reporting requirements of this section when they take adverse actions against a practitioner's credentials or employment for reasons related to clinical competence or unprofessional conduct that may adversely affect the health or welfare of the patient.

24 MRSA §2506

Society reports

Any professional society within this State which takes formal disciplinary action against a member relating to professional ethics, professional incompetence, moral turpitude, or drug or alcohol abuse shall, within 60 days of the action, report in writing to the appropriate board the name of the member, together with pertinent information relating to the action. The report shall include situations in which membership or privileges have been revoked, suspended, limited or otherwise adversely affected by action of the health care practitioner while the health care practitioner was under investigation or the subject of proceedings and it shall also include situations where membership or privileges have been revoked, suspended, limited or otherwise adversely affected by an act of the health care practitioner in return for the professional society's not conducting or for its ceasing such investigation proceeding. The report shall include situations under which an individual under societal investigation resigns during that pending investigation. The failure of any such society to report as required is a civil violation for which a fine of not more than $1,000 may be adjudged.

24 MRSA §2507

Effect of filing

The filing of a report with the board pursuant to this chapter, investigation by the board or any disposition by the board may not, in and of itself, preclude any action by a hospital or other health care facility or health care entity or professional society comprised primarily of physicians to suspend, restrict or revoke the privileges or membership of the physician.

24 MRSA §2508

Board records

Record of physicians

Each board shall create and maintain a permanent record of the names of all physicians licensed by it or otherwise lawfully practicing in this State and subject to the board's jurisdiction along with an individual historical record for each physician relating to reports or other information furnished the board under this
chapter or otherwise pursuant to law. The record may include, in accordance with rules established by the board, additional items relating to a physician's record of medical practice as will facilitate proper periodic review of the physician's professional competency.

**Reports dismissed without disciplinary action; removal and destruction**

If the board dismisses any report submitted to it without imposing disciplinary action, the report must be removed from the physician's individual historical record and destroyed, unless the report has been placed on file for a specified amount of time pursuant to Title 10, section 8003, subsection 5, paragraph E. Reports placed on file pursuant to Title 10, section 8003, subsection 5, paragraph E may only be removed and destroyed upon the expiration of the specified amount of filing time.

**Forms; acceptance of other forms**

The board shall provide forms for filing reports pursuant to this chapter. Reports submitted in other forms shall be accepted by the board.

**Disclosure to physician**

A physician shall be provided with a written notice of the substance of any information received pursuant to this chapter and placed in his individual historical record.

**Examination of records by physician; response to information**

A physician or his authorized representative shall have the right, upon request, to examine the physician's individual historical record which the board maintains pursuant to this chapter, and to place into the record a statement of reasonable length of the physician's view of the correctness or relevance of any information existing in the record. The statement shall at all times accompany that part of the record in contention. This subsection shall not apply to material submitted to the board in confidence prior to licensure by the board.

**Court action for amendment or destruction**

With the exception of orders of the board relating to disciplinary action, and reports placed on file for a specified amount of time pursuant to Title 10, section 8003, subsection 5, paragraph E, a physician has the right to seek through court action pursuant to the Maine Rules of Civil Procedure the amendment or destruction of any part of that physician's historical record in the possession of the board. When a physician initiates court action under this subsection, the board shall notify the persons who have filed complaints of the physician's request to amend these complaints or expunge them from the record. Notice to complainants must be sent to the last known address of the complainants. The notice must contain the name and address of the court to which a complainant may respond, the specific change in the complaint that the physician is seeking or the complaint that the physician seeks to expunge, and the length of time that the complainant has to respond to the court. The board shall provide complainants with at least 60 days' notice from the date the notice is sent in which to respond.

**Confidentiality of information**

The board shall provide forms for filing reports pursuant to this chapter. Reports submitted in other forms shall be accepted by the board.

**Disclosure to physician**

A physician shall be provided with a written notice of the substance of any information received pursuant to this chapter and placed in his individual historical record.

**Examination of records by physician; response to information**

A physician or his authorized representative shall have the right, upon request, to examine the physician's individual historical record which the board maintains pursuant to this chapter, and to place into the record a statement of reasonable length of the physician's view of the correctness or relevance of any information existing in the record. The statement shall at all times accompany that part of the record in contention. This subsection shall not apply to material submitted to the board in confidence prior to licensure by the board.

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**Confidentiality of information**

Any reports, information or records received and maintained by the board pursuant to this chapter, including any material received or developed by the board during an investigation shall be confidential, except for information and data that is developed or maintained by the board from reports or records received and maintained pursuant to this chapter or by the board during an investigation and that does not identify or permit identification of any patient or physician; provided that the board may disclose any confidential information only:

A. In a disciplinary hearing before the board or in any subsequent trial or appeal of a board action or order relating to such disciplinary hearing;
B. To governmental licensing or disciplinary authorities of any jurisdiction or to any health care providers or health care entities located within or outside this State that are concerned with granting, limiting or denying a physician's privileges, but only if the board includes along with the transfer an indication as to whether or not the information has been substantiated by the board;

C. As required by section 2509, subsection 5 (“Examination of records by physician”);

D. Pursuant to an order of a court of competent jurisdiction; or

E. To qualified personnel for bona fide research or educational purposes, if personally identifiable information relating to any patient or physician is first deleted.

Confidentiality of orders in disciplinary proceedings
Orders of the board relating to disciplinary action against a physician, including orders or other actions of the board referring or scheduling matters for hearing, shall not be confidential.

Confidentiality of letters of guidance or concern
Letters of guidance or concern issued by the board pursuant to Title 10, section 8003, subsection 5, paragraph E, are not confidential.

Availability of confidential information
In no event may confidential information received, maintained or developed by the board, or disclosed by the board to others, pursuant to this chapter, or information, data, incident reports or recommendations gathered or made by or on behalf of a health care provider pursuant to this chapter, be available for discovery, court subpoena or introduced into evidence in any medical malpractice suit or other action for damages arising out of the provision or failure to provide health care services. This confidential information includes reports to and information gathered by a professional review committee.

Penalty
Any person who unlawfully discloses such confidential information possessed by the board shall be guilty of a Class E crime.

Physician-patient privilege; proceedings by board
The physician-patient privilege shall, as a matter of law, be deemed to have been waived by the patient and shall not prevail in any investigation or proceeding by the board acting within the scope of its authority, provided that the disclosure of any information pursuant to this subsection shall not be deemed a waiver of such privilege in any other proceeding.

Confidentiality of professional competence review records
Except as otherwise provided by this chapter, all professional competence review records are privileged and confidential and are not subject to discovery, subpoena or other means of legal compulsion for their release to any person or entity and are not admissible as evidence in any civil, judicial or administrative proceeding. Information contained in professional competence review records is not admissible at trial or deposition in the form of testimony by an individual who participated in the written professional competence review process. Nothing in this section may be read to abrogate the obligations to report and provide information under section 2506, nor the application of Title 32, sections 2599 and 3296.

Protection; waiver
This chapter's protection may be invoked by a professional competence committee or by the subject of professional competence review activity in any civil, judicial or administrative proceeding. This section's protection may be waived only by a written waiver executed by an authorized representative of the professional competence committee.
**Adverse professional competence review action**
Subsection 1 does not apply in a proceeding in which a physician contests an adverse professional competence review action against that physician, but the discovery, use and introduction of professional competence review records in such a proceeding does not constitute a waiver of subsection 1 in any other or subsequent proceedings seeking damages for alleged professional negligence against the physician who is the subject of such professional competence review records.  

24 MRSA §2510-A (2)

**Defense of professional competence committee**
Subsection 1 does not apply in a proceeding in which a professional competence committee uses professional competence review records in its own defense, but the discovery, use and introduction of professional competence review records in such a proceeding does not constitute a waiver of subsection 1 in the same or other proceeding seeking damages for alleged professional negligence against the physician who is the subject of such professional competence review records.  

24 MRSA §2510-A (3)

**Waiver regarding individual**
Waiver of subsection 1 in a proceeding regarding one physician does not constitute a waiver of subsection 1 as to other physicians.  

24 MRSA §2510-A (4)

**Release to other review bodies, agencies, accrediting bodies**
A professional competence committee may furnish professional competence review records or information to other professional review bodies, state or federal government agencies and national accrediting bodies without waiving any privilege against disclosure under section 2510-A.  

24 MRSA §2510-B (1)

**Release to physician**
A professional competence committee may furnish professional competence review records to the physician who is the subject of the professional competence review activity and the physician's attorneys, agents and representatives without waiving any privilege against disclosure under section 2510-A.  

24 MRSA §2510-B (2)

**Release of directory information**
A professional competence committee may furnish directory information showing membership, clinical privileges, provider panel or other practice status of a physician with the health care entity to anyone without waiving the privilege against disclosure under section 2510-A.  

24 MRSA §2510-B (3)

**Immunity**
Any person acting without malice, any physician, podiatrist, health care provider, health care entity or professional society, any member of a professional competence committee or professional review committee, any board or appropriate authority and any entity required to report under this chapter are immune from civil liability:

- **Reporting:** For making any report or other information available to any board, appropriate authority, professional competence committee or professional review committee pursuant to law;
- **Assisting in preparation:** For assisting in the origination, investigation or preparation of the report or information described in subsection 1; or
- **Assisting in duties:** For assisting the board, authority or committee in carrying out any of its duties or functions provided by law.  

24 MRSA §2511

**MMA’s Medical Professional Health Program**
A Medical Professional Health Program is conducted by the Maine Medical Association under protocols developed with the Board of Licensure in Medicine, the Board of Osteopathic Licensure, the Board of Dental Examiners, the Board of Pharmacy, the Board of Nursing and the Board of Veterinary Medicine. If you, or a colleague, have a potential impairment, please contact the Medical Professional Health Program at (207) 623-9266
The program has a website at: https://www.mainemphp.org/

(51) Insurance Regulations (Select Sections)

Health Care Bill of Rights
Part A requires carriers to provide a toll-free telephone number that certificate holders can call to determine if a policy has been cancelled or reinstated after payment of the premium. It requires carriers to provide notice to plan enrollees regarding any exclusions or limits of coverage for childhood immunizations. Part A also requires carriers to post at least 5 individual and 5 small group health plans on its publicly accessible website for comparison purposes and sets minimum standards for explanation of benefits documents used by carriers.
Part B establishes standards for provider profiling programs used by carriers.
Part C requires carriers and health maintenance organizations to include certain information about product offerings in the annual report supplement to the Department of Professional and Financial Regulation, Bureau of Insurance.
Part D extends the notice period for all carriers to notify policyholders of proposed rate increases. It also permits the Attorney general to request a rate hearing regarding proposed rate increases for individual health plans.
Part E requires health maintenance organizations to disclose loss information upon request from contract holders in the same manner as insurance companies. Part E also authorizes the Superintendent of Insurance to adopt rules requiring small group health carriers to offer standardized small group health plans. Part E also requires the Superintendent of Insurance to undertake market conduct exams of health insurance companies no less frequently than once every 3 years, beginning in 2010.
Part F requires a carrier replacing a previous carrier to honor any prior authorizations for prescription drugs for an enrollee undergoing a course of treatment for a period of 6 months.

For the full text see Public Laws 2009, Chapter 439. The law is found in various sections of Title 24-A, the Maine Insurance Code.

Provider Profiling Programs Used by Insurance Companies
An insurer delivering or issuing for delivery any individual health insurance policy, group health insurance policy, or certificate shall file with the superintendent of insurance an annual statement that includes the full selection criteria, standards, practices, procedures, and programs that measure or tier health care provider performance with respect to quality, cost, or cost-efficiency.

A “provider profiling program” means a program utilizing provider data in order to rate or rank provider quality, cost or efficiency of care by the use of a grade, star, tier, rating or any other form of designation that provides an enrollee with an incentive to use a designated provider based on quality, cost or efficiency of care. At least sixty days prior to using or publicly disclosing the result of the provider program, a carrier shall notify the provider to what criteria they will evaluate them on. Results of such provider-profiling program shall be shared with the provider sixty days before publicly disclosing or utilizing the result of the provider-profiling program. Upon request, a provider is entitled to a copy of the results of any provider-profiling program. Moreover, a carrier shall establish an appeals process and a method of review dispute.

Individual Provider Contracts With Insurance Companies
If a contract for a preferred provider arrangement includes a reference to policies or procedures to which a contracting provider would be bound, such policies and procedures must be provided to the provider for review.
Upon the provider’s request, such policies and procedures must be made in an easily accessible manner and at the time the contract is offered. Moreover, at the time of contract, and at the provider’s request, the following must be made available: (1) the fee schedule or, if there is not a fee schedule for one or more of the services covered under the contract, the terms under which payment is determined. A carrier may request a provider to execute a nondisclosure agreement. (2) The identity of all carriers for which the provider is agreeing to provide services to health plan enrols.

As a condition of participation one of the carrier’s preferred provider arrangements, a contract offered by a carrier may not require a provider to participate in any other carrier’s network subsequently offered by the carrier or by a carrier’s preferred provider arrangement.

Without the providers’ prior written consent, a provider’s contractual participation in a carrier’s preferred provider arrangement may not: (1) subject the provider to health plan payer requirements or fee schedules that materially different from the terms of the provider’s contract with the carrier, unless those materially different terms are set out in writing in a separate section of the contract, such as an exhibit or an amendment; or (2) Permit the terms of the provider’s existing preferred provider arrangement contract to be superseded by a carrier’s subsequent contract with a health plan payer.

A preferred provider arrangement contract may not require a provider providing a service to an enrollee under a health plan included in the provider’s contract to obtain preauthorization as a condition of coverage.

Explanation of remittance advices or comparable documents, whether in paper or electronic form, that accompany and identify payment of a provider’s claims under a carrier’s contract, including contracts offered through a preferred provider arrangement, must identify the administrator and payer of the providers’ claims and include contact information.

24-A MRSA §4303 (18)

Plan requirements
A carrier offering or renewing a health plan in this State must meet a number of requirements regarding:

1. Demonstration of adequate access to providers
2. Credentialing
   2-A. Payment to provider for services rendered during pendency of credentialing.
3. Provider's right to advocate for medically appropriate care
4. Termination of participating providers
5. Prohibition on financial incentives
6. Grievance procedure for enrollees
7. Identification of services provided by certified nurse practitioners and certified nurse midwives
8. Standing referrals to specialists
9. Continuity of care
   7-A Continuity of prescriptions
10. Maximum allowable charges
11. Absolute discretion clauses
12. Protection from balance billing by participating providers
13. Notice of amendments to provider agreements
14. Limits on retrospective denials
15. Uniform explanation of coverage documents and standardized definitions
16. Language and culture
17. Prohibition on "most favored nation" clauses.
18. Provider contract requirements.
19. Information about provider networks
20. Information about prescription drugs
21. Information about estimated costs for the services offered and comparable health care services (health care price transparency tools).
22. Carriers may not deny payment due solely to the fact of an out-or-network provider’s referral.

24-A MRSA § 4303

For more requirements of health insurance carriers, including issues of utilization review, enrollee choice of primary care and other providers, access to clinical trials, and external reviews, see also 24-A MRSA, Chapter 56-A, the Health Plan Improvement Act.

**Incentive Program for comparable services at lower cost**
Beginning January 1, 2019, carriers must establish plans in which enrollees are directly incentivized to shop for low-cost, high-quality participating providers for comparable health care services in the categories of:
1. Physical and occupational therapy;
2. Radiology and imaging;
3. Laboratory services; and
4. Infusion therapy.

24-A MRSA § 4318-A

**Limitations on out-of-network costs**
Beginning January 1, 2019, if an enrollee elects to obtain a covered comparable health care service from an out-of-network provider at a price that is the same or less than the statewide average for the same covered health care service, the carrier must apply enrollee’s payments for that service toward the enrollee’s deductible and out-of-pocket maximum as if the services had been provided by an in-network provider.

24-A MRSA § 4318-B

**Mandated Benefits**
Maine law contains a number of mandated benefits; requiring health insurance plans to cover certain services, providers or individuals. Examples of mandated benefits as of 2018 include maternity and routing newborn care, preventive care, medically necessary infant formula, contraceptives, mental health services, home health services, diabetes supplies and services, mammograms, acupuncture services, breast cancer treatment, prostate cancer screening, colorectal cancer screening, chiropractic services, treatment for HIV/AIDS, children’s early intervention services, orally administered chemotherapeutic drugs and coverage for autism spectrum disorders.

A health policy that provides coverage for the mandate as required may contain provisions for maximum benefits and coinsurance and limitations, deductibles and exclusions to the same extent that these provisions are applicable to all coverage and are not inconsistent with the requirements of the particular mandate.

For a summary of all mandated benefits in Maine and which plans they apply to, see the Bureau of Insurance chart, [http://www.maine.gov/pfr/insurance/consumer/consumer_guides/mandated_benefits.html](http://www.maine.gov/pfr/insurance/consumer/consumer_guides/mandated_benefits.html)

For the full list of requirements that apply to nonprofit hospital service plans, nonprofit medical service plans or nonprofit health care plans, see Title 24, Chapter 19.

For the full list of requirements that apply to individual plans, see Title 24-A, Chapter 33.
For the full list of requirements that apply to group health plans, see Title 24-A, Chapter 35.
For the full list of requirements that apply to HMOs, see Title 24-A, Chapter 56.
For requirements that apply to all plans, see Title 24-A, Chapter 56-A, the Health Plan Improvement Act.

Telemedicine Service Coverage

Telemedicine; definition
For the purposes of this section, “telemedicine,” as it pertains to the delivery of health care services, means the use of interactive audio, video or other electronic media for the purpose of diagnosis, consultation or treatment. “Telemedicine” does not include the use of audio-only telephone, facsimile machine or e-mail.

Coverage
A carrier offering a health plan in this State may not deny coverage on the basis that the coverage is provided through telemedicine if the health care service would be covered were it provided through in-person consultation between the covered person and a health care provider. Coverage for health care services provided through telemedicine must be determined in a manner consistent with coverage for health care services provided through in-person consultation. A carrier may offer a health plan containing a provision for a deductible, copayment or coinsurance requirement for a health care service provided through telemedicine as long as the deductible, copayment or coinsurance does not exceed the deductible, copayment or coinsurance applicable to an in-person consultation.

Interstate practice of telemedicine
A physician not licensed to practice medicine in this State may provide consultative services through interstate telemedicine to a patient located in this State if the physician is registered in accordance with subsection 3. A physician intending to provide consultative services in this State through interstate telemedicine shall provide any information requested by the board and complete information on:
   A. All states and jurisdictions in which the physician is currently licensed;
   B. All states and jurisdictions in which the physician was previously licensed; and
   C. All negative licensing actions taken previously against the physician in any state or jurisdiction.

Interstate practice of telemedicine, Registration
The board may register a physician to practice medicine in this State through interstate telemedicine if the following conditions are met:
   A. The physician is fully licensed without restriction to practice medicine in the state from which the physician provides telemedicine services;
   B. The physician has not had a license to practice medicine revoked or restricted in any state or jurisdiction;
   C. The physician does not open an office in this State, does not meet with patients in this State, does not receive calls in this State from patients and agrees to provide only consultative services as requested by a physician, advanced practice registered nurse or physician assistant licensed in this State and the physician, advanced practice registered nurse or physician assistant licensed in this State retains ultimate authority over the diagnosis, care and treatment of the patient;
   D. The physician registers with the board every 2 years, on a form provided by the board; and
   E. The physician pays a registration fee not to exceed $500.

Mandatory offer to extend coverage for dependent children up to 25 years of age

Dependent child; definition
As used in this section, “dependent child” means the child of a person covered under an individual health insurance policy when that child:
A. Is unmarried;
B. Has no dependent of the child’s own; and
C. Is a resident of this State or is enrolled as a full-time student at an accredited public or private institution of higher education.

Offer to extend coverage
Notwithstanding section 2703, subsection 3, an individual health insurance policy that offers coverage for a dependent child must offer such coverage, at the option of the policyholder, until the dependent child is 25 years of age. An insurer may require, as a condition of eligibility for coverage in accordance with this section, that a person seeking coverage for a dependent child provide written documentation on an annual basis that the dependent child meets the requirements in subsection 1.

24-A MRSA §2742-B (Individual Policies)
24-A MRSA §2833-B (Group Plans)
24-A MRSA §4233-B (HMOs)

Extension of Dependent Coverage
A carrier offering a health plan subject to the requirements of the federal Affordable Care Act that provides dependent coverage of children shall continue to make such coverage available for an adult child until the child turns 26 years of age, consistent with the federal Affordable Care Act

24-A MRSA §4320-B

(52) Interns and Residents
Licensing of Interns and Residents
Residents
An applicant who is qualified under section 3271, subsection 1, may receive a temporary educational certificate from the board to act as a hospital resident. A certificate to a hospital resident may be renewed annually at the discretion of the board for not more than 7 years.

32 MRSA §3279 (2)

Joint-program resident
An applicant who is enrolled in a program of medical and graduate medical training conducted jointly by a medical school accredited by the Liaison Committee on Medical Education and a graduate medical education program approved by the Accreditation Council on Graduate Medical Education may receive a temporary educational certificate from the board to act as a hospital resident as part of that graduate medical education program if the applicant is concurrently enrolled in the final year of medical training and the initial year of graduate medical education. The board may not issue a certificate pursuant to this subsection for a period longer than that required to obtain the M.D. degree. The period during which the certificate is in force may not be considered in determining satisfaction of the requirement for postgraduate medical education under section 3271, subsection 2.

32 MRSA §3279 (2-A)

Conditions of certification
An applicant for a temporary educational certificate may not be certified unless the board finds that the applicant is qualified and that there exists no cause, as set forth in section 3282-A, that would be considered grounds for disciplinary action against a licensed physician or surgeon. The board, in its discretion, may require an examination for applicants for temporary educational certificates. Recipients of these certificates are entitled to all the rights granted to physicians who are licensed to practice medicine and surgery, except that their practice is limited to the training programs in which they are enrolled. A temporary educational certificate may be suspended or revoked, or the board may refuse to renew the certificate, for the reasons stated in section 3282-A, or if the intern or hospital resident has violated the limitations placed upon the intern's temporary educational certificate.

32 MRSA §3279 (3)
Visiting instructors
A physician who has an unrestricted license to practice medicine or surgery in another state may practice medicine or surgery in this State when the physician is performing medical procedures as part of a course of instruction in graduate medical education in a hospital located in this State. The right of a visiting medical instructor to practice medicine in this State may be suspended or revoked for the reasons stated in section 3282-A, or if the visiting medical instructor has performed medical procedures that are not a part of a course of instruction.

32 MRSA §3279 (4)

Contract students
An applicant who is qualified under section 3271, subsection 1, who received a medical education as a contract student as provided in Title 20-A, chapter 421, and who agrees to practice in a primary care or other specialized area as defined in Title 20-A, section 11803, subsection 2, or an underserved area as defined in Title 20-A, section 11802, is considered to have completed the postgraduate training requirements of section 3271, subsection 2, upon satisfactory completion of at least 12 months in a graduate educational program approved as specified in section 3271. The board may make the relicensure of an individual for 4 years after the individual's licensure under this subsection contingent on the individual's continuing to practice in an underserved area. This subsection applies only to individuals entering into a contract under Title 20-A, chapter 421, on or before December 31, 1984.

32 MRSA §3279 (5)

Fees
The board shall set fees for physicians and students licensed pursuant to this section. The amounts set for licenses issued under this section may not be more than $300.

32 MRSA §3279 (6)

(53) Jury Duty
Exemptions
Maine law exempts only the following individuals from jury service:
   A. The Governor;
   B. Those under the age of 18;
   C. Non-residents of the particular county;
   D. Active duty military personnel;
   E. Non-citizens of the U.S.;
   F. Those unable to read, speak and understand the English language.
   (The law changed in 2017 to remove the former exemptions for physicians, dentists, judges, attorneys, and others.)

14 MRSA § 1211

The court also has the authority to excuse individuals from jury service upon a showing of undue hardship, extreme inconvenience, public necessity or inability to render satisfactory jury service because of physical or mental disability.

For more information, visit the State of Maine Judicial Branch website at:
http://www.courts.maine.gov/maine_courts/superior/jury/exemptions.html

(54) Lead Poisoning Prevention
Screening by health care providers
All health care providers shall advise parents of the availability and advisability of screening their children for lead poisoning. A health care program that receives funds from the State and has a child health component shall provide screening of children for lead poisoning in accordance with rules adopted by the department.
For more information:
Maine Lead Poisoning Control Act, 32 MRSA, Chapter 252.
Maine CDC Screening, Testing and Follow Up Guidelines
Maine Lead Poisoning Prevention Program
CDC Childhood Lead Poisoning Prevention Program

Maine Lead Poisoning Prevention Fund

Fund established
The Lead Poisoning Prevention Fund, referred to in this section as "the fund," is established within the department as a nonlapsing fund for the purposes specified in this section.

Sources of fund
The fund is funded from all fees collected under section 1322-F and from other funds accepted by the commissioner or allocated or appropriated by the Legislature

Prevention purposes
Allocations from the fund must be made for the following purposes:
A. Contracts for funding community and worker educational outreach programs to enable the public to identify lead hazards and take precautionary actions to prevent exposure to lead;
B. An ongoing major media campaign to fulfill the purposes of the educational and publicity program required by section 1317-B;
C. Measures to prevent children's exposure to lead, including targeted educational mailings to families with children that occupy dwellings built prior to 1978 with culturally appropriate information on the health hazards of lead, the identification of lead sources, actions to take to prevent lead exposure and the importance of screening children for lead poisoning;
D. Measures to prevent occupational exposures to lead for private and public employees, including improvements in the effectiveness of the occupational disease reporting system required in chapter 259-A in identifying and educating health care providers, employers and lead-exposed adults about occupational lead poisoning prevention strategies; and
E. Funding an assessment of current uses of lead and the availability, effectiveness and affordability of lead-free alternatives.
F. Funding for educational programs and information for owners of rental property used for residential purposes.

Administration
The Bureau of Health shall administer the fund allocations with the review and advice of an advisory board established by the department pursuant to section 1323. Preference must be given to programs that reach high-risk or underserved populations. The bureau may contract for professional services to carry out the purposes of this section.

(55) Limited English Proficiency (LEP)
As part of the federal government’s effort to promote effective communication between health and social service providers and people with limited English proficiency, the Office of Civil Rights has issued revised guidance to help practitioners and organizations meet the federal requirements for LEP. Below are some of the key excerpts from the OCR guidance.
**Who Is Covered**

Department of Health and Human Services regulations, 45 CFR §80.3(b)(2), require all recipients of federal financial assistance from HHS to provide meaningful access to LEP persons. Federal financial assistance includes grants, training, use of equipment, donations of surplus property, and other assistance. Recipients of HHS assistance may include:

A. Hospitals, nursing homes, home health agencies, and managed care organizations  
B. Universities and other entities with health or social service research programs  
C. State, county, and local health agencies  
D. State Medicaid agencies  
E. State, county and local welfare agencies  
F. Programs for families, youth, and children  
G. Head Start programs  
H. Public and private contractors, subcontractors and vendors  
I. Physicians and other providers who receive Federal financial assistance from HHS

Recipients of HHS assistance do not include, for example, providers who only receive Medicare Part B payments. However, MaineCare does require MaineCare “providers [to] ensure that MaineCare members are able to communicate effectively with them regarding their medical needs.” MaineCare will reimburse providers for interpreters required for limited and non-English speaking members and/or deaf/hard of hearing members, when these services are necessary to communicate effectively with members regarding health needs. Interpreter services can only be covered in conjunction with another covered MaineCare service. For more guidance on MaineCare’s policy on paying for interpretation services, see: [http://www.maine.gov/dhhs/oma/MulticulturalResource/appendix_g.html](http://www.maine.gov/dhhs/oma/MulticulturalResource/appendix_g.html). The bottom line is that virtually all physician offices will be required to provide language access services.

**Who is a Limited English Proficient Individual**

Individuals who do not speak English as their primary language and who have a limited ability to read, write, speak, or understand English may be limited English proficient, or "LEP," and may be eligible to receive language assistance with respect to a particular type of service, benefit, or encounter.

**How Does a Recipient Determine the Extent of Its Obligation To Provide LEP Services**

Recipients are required to take reasonable steps to ensure meaningful access to their programs and activities by LEP persons. While designed to be a flexible and fact-dependent standard, the starting point is an individualized assessment that balances the following four factors:

**Four Factor Test:**

A. The number or proportion of LEP persons eligible to be served or likely to be encountered by the program or grantee;  
B. The frequency with which LEP individuals come in contact with the program;  
C. The nature and importance of the program, activity, or service provided by the program to people's lives; and  
D. The resources available to the grantee/recipient and costs. As indicated above, the intent of this guidance is to suggest a balance that ensures meaningful access by LEP persons to critical services while not imposing undue burdens on small business, small local governments, or small nonprofits.

After applying the above four-factor analysis, a recipient may conclude that different language assistance measures are sufficient for the different types of programs or activities in which it engages, or, in fact, that, in certain circumstances, recipient-provided language services are not necessary. (As discussed below, recipients may want to consider documenting their application of the four-factor test to the services they provide.) For instance, some of a recipient's activities will be more important than others
and/or have greater impact on or contact with LEP persons, and thus may require more in the way of language assistance.

Recipients have two main ways to provide language services: **Oral interpretation** either in person or via telephone interpretation service (hereinafter "interpretation") and **written translation** (hereinafter "translation"). Oral interpretation can range from on-site interpreters for critical services provided to a high volume of LEP persons, to access through commercially-available telephonic interpretation services. Written translation, likewise, can range from translation of an entire document to translation of a short description of the document.

For example, if two physicians in the same field, one with a Spanish-speaking assistant and one with a Vietnamese-speaking assistant, practice in the same geographic area and have a custom/practice of referring patients between each other, it may be appropriate for the first doctor to refer LEP Vietnamese patients to the second doctor and for the second doctor to refer LEP Spanish patients to the first doctor. In certain circumstances, a referral would not be appropriate: for example, a Korean speaking LEP woman comes to a battered women's shelter requesting assistance. Although the shelter has space, it has no arrangement to provide language assistance for LEP persons. Instead, as with all LEP persons, the staff only offer her a prepared list of three shelters in the neighborhood that generally provide language assistance. The staff does not check to assure that any of the three alternative shelters can actually provide the Korean language assistance she needs, or that any have space available for her.

The correct mix should be based on what is both necessary and reasonable in light of the four-factor analysis.

**Oral Language Services (Interpretation)**

Interpretation is the act of listening to something in one language (source language) and orally translating it into another language (target language). Where interpretation is needed and is reasonable, recipients should consider some or all of the following options for providing competent interpreters in a timely manner:

A. Hiring Bilingual Staff
B. Hiring Staff Interpreters
C. Contracting for Interpreters
D. Using Telephone Interpreter Lines
E. Using Community Volunteers
F. Use of Family Members or Friends as Interpreters

**Written Language Services (Translation)**

**What Documents Should be Translated**

- Consent and complaint forms.
- Intake forms with the potential for important consequences
- Written notices of eligibility criteria, rights, denial, loss, or decreases in benefits or services, actions affecting parental custody or child support, and other hearings
- Notices advising LEP persons of free language assistance
- Written tests that do not assess English language competency, but test competency for a particular license, job, or skill for which knowing English is not required
- Applications to participate in a recipient's program or activity or to receive recipient benefits or services

**Nonvital written materials could include:**

- Hospital menus
- Third party documents, forms, or pamphlets distributed by a recipient as a public service
C. For a non-governmental recipient, government documents and forms
D. Large documents such as enrollment handbooks (although vital information contained in large documents may need to be translated)
E. General information about the program intended for informational purposes only

Elements of Effective Plan on Language Assistance for LEP Persons
If, after completing the four-factor analysis, a recipient determines that it should provide language assistance services, a recipient may develop an implementation plan to address the identified needs of the LEP populations it serves. Such recipients have considerable flexibility in developing this plan.

(http://www.hhs.gov/ocr/civilrights/resources/specialtopics/lep/policyguidancedocument.html)

(56) MaineCare (see also Agency Rules Links)
MaineCare is the state’s medical assistance program and is administered by the Bureau of Medical Service, Department of Human Services. In 2009, the Legislature adopted a compromise on proposed cuts in reimbursement to hospital-based physicians that increased the MaineCare physician fee schedule from approximately 57% to approximately 70% of Medicare rates effective February 1, 2010.

Useful Links
Office of MaineCare Services (formerly Bureau of Medical Services) Home Page
Office of MaineCare Services Provider Page
MaineCare Benefits Manual, Department of Health and Human Services Rule Chapter 101

(57) Maine Health Data Organization (see also Agency Rules Links)

Purposes
The objective of the organization is to create and maintain a useful, objective, reliable and comprehensive health information database that is used to improve the health of Maine citizens and to issue reports, as provided in section 8712. This database must be publicly accessible while protecting patient confidentiality and respecting providers of care. The organization shall collect, process, analyze and report clinical, financial, quality and restructuring data. The organization shall promote public transparency of the quality and cost of health care in the State in conjunction with the Maine Quality Forum. It is also required by statute to create an interactive website displaying prices paid for specific procedures performed at all Maine hospitals and impacted surgical, diagnostic or other nonhospital facilities.

22 MRSA § 8703
22 MRSA § 8712

The organization is governed by Title 22 MRSA, Chapter 1683 and rules. For more information, view visit the MHDO Website.

Clinical data
Information required
Pursuant to rules adopted by the board for form, medium, content and time for filing, each health care facility shall file with the organization the following information:
A. A completed uniform hospital discharge data set, or comparable information, for each patient discharged from the facility after June 30, 1983 and for each hospital outpatient service occurring after June 30, 1996; and
B. In addition to any other requirements applicable to specific categories of health care facilities, the organization may require the filing of data as set forth in this chapter or in rules adopted pursuant to this chapter.

Additional information on ambulatory services and surgery
Pursuant to rules adopted by the board for form, medium, content and time for filing, each provider shall file with the organization a completed data set, comparable to data filed by health care facilities under subsection 1, paragraph B. This subsection may not be construed to require duplication of information required to be filed under subsection 1.

More than one licensed health care facility or location
When more than one licensed health care facility is operated by the reporting organization, the information required by this chapter must be reported for each health care facility separately. When a provider of health care operates in more than one location, the organization may require that information be reported separately for each location.

Medical record abstract data
In addition to the information required to be filed under subsections 1 and 2 and pursuant to rules adopted by the organization for form, medium, content and time of filing, each health care facility shall file with the organization such medical record abstract data as the organization may require.

Merged data
The board may require the discharge data submitted pursuant to subsection 1 and any medical record abstract data required pursuant to subsection 5 to be merged with associated billing data.

Additional data
Subject to the limitations of section 8704, subsection 1, the board may adopt rules requiring the filing of additional clinical data from other providers and payors as long as the submission of data to the organization is consistent with federal law. Data filed by payors must be provided in a format that does not directly identify the patient.

Authority to obtain information
Nothing in this section may be construed to limit the board's authority to obtain information that it considers necessary to carry out its duties.

Financial data; scope of service data
Financial data and scope of service data must be filed, stored and managed as follows.

Financial data
Each health care facility shall file with the organization, in a form specified by rule pursuant to section 8704, financial information including costs of operation, revenues, assets, liabilities, fund balances, other income, rates, charges and units of services, except to the extent that the board specifies by rule that portions of this information are unnecessary.

Hospitals; standardized accounting template
When filing the financial information required under subsection 1, a hospital also shall file information using the standardized accounting template published in the report of the Commission to Study Maine’s Community Hospitals in February 2005. The hospital shall file this information using an electronic version of the template provided to the hospital by the organization. If in succeeding years the template needs to be modified, the board shall adopt rules specifying the filing requirements. Rules adopted pursuant to this subsection are routine technical rules as defined in Title 5, chapter 375, subchapter 2-A.
Certification required
The board may require certification of such financial reports and attestation from responsible officials of
the health care facility that such reports have to the best of their knowledge and belief been prepared in
accordance with the requirements of the board.

Scope of service data
Each health care facility shall file with the organization scope of service information, including bed
capacity by service provided, special services, ancillary services, physician profiles in the aggregate by
clinical specialties, nursing services and such other scope of service information as the organization
determines necessary for the performance of its duties.

Other health care information
Development of health care information systems
In addition to its authority to obtain information to carry out the specific provisions of this chapter, the
organization may require providers and payors to furnish information with respect to the nature and
quantity of services or coverage provided to the extent necessary to develop proposals for the
modification, refinement or expansion of the systems of information disclosure established under this
chapter. The organization's authority under this subsection includes the design and implementation of
pilot information reporting systems affecting selected categories or representative samples of providers
and payors.

Reports
The organization shall produce clearly labeled and easy-to-understand reports as follows. Unless
otherwise specified, the organization shall distribute the reports on a publicly accessible site on the
Internet or via mail or e-mail, through the creation of a list of interested parties. The organization shall
publish a notice of the availability of these reports at least once per year in the 3 daily newspapers of the
greatest general circulation published in the State. The organization shall make reports available to
members of the public upon request.

Quality
At a minimum, the organization, in conjunction with the Maine Quality Forum, established in Title 24-A,
section 6951, shall develop and produce annual quality reports.

Average Payments
At a minimum, the organization, with advice from the Maine Health Data Processing Center as authorized
in Title 10, section 681, shall develop and produce annual reports on average private-payer payments for
services provided by health care facilities and health care practitioners, excluding emergency services.
For health care facilities, the reports must include, but are not limited to, the average private-payer
payments per service per facility and total number of services per facility.

Comparison report
At a minimum, the organization shall develop and produce an annual report that compares the 15 most
common diagnosis-related groups and the 15 most common outpatient procedures for all hospitals in the
State and the 15 most common procedures for nonhospital health care facilities in the State to similar data
for medical care rendered in other states, when such data are available.

Physician services
The organization shall provide an annual report of the 10 services and procedures most often provided by
osteopathic and allopathic physicians in the private office setting in this State. The organization shall
distribute this report to all physician practices in the State.
Website
The organization must have a publicly accessible interactive website that present the reports on payments for services rendered to Maine residents. The services presented must include at least imaging, preventative health, radiology, surgical services, comparable health services as defined in 24-A MRSA §4318-A(1)(A), and other services that are predominantly elective and may be provided to many patients who are uninsured or underinsured. The website must include data submitted by payors with regard to all health care facilities and practitioners providing comparable health care services as so defined.

22 MRSA §8712

(58) Managed Care Organizations, Select Statutes
Health Plan Improvement Act (Patient Bill of Rights)

Definitions – see 24-A MRSA §4301-A

HMOs and the Practice of Medicine
Any health maintenance organization authorized under this chapter is not deemed to be practicing medicine and is exempt from provisions of law relating to the practice of medicine, except that this subsection may not be asserted by a health maintenance organization as a defense to any action brought by an enrollee pursuant to section 4313.

Confidentiality; liability; access to records
Confidentiality
Any data or information pertaining to the diagnosis, treatment or health of an enrollee or applicant obtained from that enrollee or applicant or a provider by a health maintenance organization must be held in confidence and may not be disclosed to any person except: to the extent that it may be necessary to carry out the purposes of this chapter; upon the express consent of the enrollee or applicant; pursuant to statute or court order for the production of evidence or the discovery of evidence; or in the event of claim or litigation between that enrollee or applicant and the health maintenance organization when such data or information is pertinent. A health maintenance organization is entitled to claim any statutory privileges against such disclosure that the provider who furnished such information to the health maintenance organization is entitled to claim.

24-A MRSA §4222 (3)

Liability
A person who, in good faith and without malice, as a member, agent or employee of a quality assurance committee, assists in the origination, investigation or preparation of a report or information related to treatment previously rendered, submits that report or information to a health maintenance organization or appropriate state licensing board, or assists the committee in carrying out any of its duties under this chapter is not subject to civil liability for damages as a consequence of those actions, nor is the health maintenance organization that established that committee or the officers, directors, employees or agents of that health maintenance organization liable for the activities of that person. This section may not be construed to relieve any person of liability arising from treatment of a patient.

24-A MRSA §4224 (1)

A. The information considered by a quality assurance committee and the records of its actions and proceedings are confidential and not subject to subpoena or order to produce except in proceedings before the appropriate state licensing or certifying agency or in an appeal, if permitted, from the findings or recommendations of the committee. A member of a quality assurance committee or an officer, director, staff person or other member of a health maintenance organization engaged in assisting the committee or any person assisting or furnishing information to the committee may not be subpoenaed to testify in any judicial or quasi-judicial proceeding if the subpoena is based solely on these activities.
B. Information considered by a quality assurance committee and the records and proceedings of that committee used pursuant to paragraph A by a state licensing or certifying agency or in an appeal must be kept confidential and are subject to the same provisions concerning discovery and use in legal actions as are the original information and records in the possession and control of the health care review committee.

Access to records
To fulfill the obligations of a health maintenance organization under section 4204, subsection 2-A, paragraph B, a health maintenance organization must have access to treatment records and other information pertaining to the diagnosis, treatment and health status of any enrollee.

Enrollee choice of primary care provider
A carrier offering a managed care plan shall allow enrollees to choose their own primary care providers, as allowed under the managed care plan's rules, from among the panel of participating providers made available to enrollees under the managed care plan's rules. A carrier shall allow physicians, and certified nurse practitioners who have been approved by the State Board of Nursing to practice advanced practice registered nursing without the supervision of a physician pursuant to Title 32, section 2102, subsection 2-A, to serve as primary care providers for managed care plans. A carrier is not required to contract with certified nurse practitioners or physicians as primary care providers in any manner that exceeds the access and provider network standards required in this chapter or chapter 56, or any rules adopted pursuant to those chapters. A carrier must allow enrollees in a managed care plan to change primary care providers without good cause at least once annually and to change with good cause as necessary. When an enrollee fails to choose a primary care provider, the carrier may assign the enrollee a primary care provider located in the same geographic area in which the enrollee resides.

Indemnification
A contract between a carrier offering a health plan and a provider for the provision of services to enrollees may not require the provider to indemnify the carrier for any expenses and liabilities, including, without limitation, judgments, settlements, attorney's fees, court costs and any associated charges incurred in connection with any claim or action brought against the health plan based on the carrier's own fault. Nothing in this section may be construed to remove responsibility of a carrier or provider for expenses or liabilities caused by the carrier's or provider's own negligent acts or omissions or intentional misconduct.

Limit on retrospective denials of previously paid health insurance claims
The time that has elapsed since the date of payment of the previously paid claim does not exceed 12 months. The retrospective denial of a previously paid claim may be permitted beyond 12 months from the date of payment only for the following reasons:
  A. The claim was submitted fraudulently;
  B. The claim payment was incorrect because the provider or the insured was already paid for the health care services identified in the claim;
  C. The health care services identified in the claim were not delivered by the provider;
  D. The claim payment was for services covered by Title XVIII, Title XIX or Title XXI of the Social Security Act;
  E. The claim payment is the subject of adjustment with another insurer, administrator or payor; or
  F. The claim payment is the subject of legal action.

Other Statutory References of Interest:
Reporting Requirement: 24-A MRSA §4302
Plan Requirements: 24-A MRSA §4302; 24-A MRSA §4303
Utilizations Review: 24-A MRSA §4304
Quality of Care: 24-A MRSA §4305
Access to Clinical Trials: 24-A MRSA §4310
Access to prescription drugs: 24-A MRSA §4311
Independent external review: 24-A MRSA §4312
Carrier liability; cause of action: 24-A MRSA §4313

(59) Mandatory Reports
Maine law requires the mandatory reporting of many diseases, conditions, situations, etc. Many of these are covered in the sections of the same title. For purposes of efficiency, a list of major title mandatory reports is listed below.

List of Mandatory Reports
A. Abortions and Miscarriages
B. Abuse (Adult, Child and Elder)
C. AIDS and HIV
D. Births
E. Burns
F. Cancer
G. Communicable Diseases (Public Health)
H. Crime, Failure to report
I. Deaths
J. Gunshot Wounds
   A person is guilty of failure to report treatment of a gunshot wound if, being a health care practitioner or emergency medical services person, that person treats a human being for a wound apparently caused by the discharge of a firearm and knowingly fails to report the same to a law enforcement agency immediately by the quickest means of communication.
K. Head Injury
L. Impaired Physician
M. Lead Poisonings
N. Occupational Diseases (Public Health)
O. Sentinel Events
P. New Born Hearing Evaluations

17-A MRSA §512 (1)

(60) Medical Education Program
Doctors for Maine’s Future Scholarship Program
There is established the Doctors for Maine’s Future Scholarship Program, referred to in this section as “the scholarship program,” to provide a tuition subsidy of 50% of the cost of attendance annually, up to a maximum of $25,000 per student annually, for eligible students who enter qualifying Maine-based medical school programs for the purpose of increasing the number of physicians in this State who practice in primary care, underserved specialties or underserved areas of the State. For the purposes of this section, “cost of attendance” means the tuition and fees applicable to an eligible student, together with estimated other expenses reasonably related to cost of attendance at a qualifying Maine-based medical school program.

20-A MRSA §12103-A

Qualifying Maine-based medical school program
“Qualifying Maine-based medical school program” means an allopathic or osteopathic medical school program affiliated with a medical school accredited by the Liaison Committee on Medical Education or its successor or the American Osteopathic Association or its successor in which:
A. An educational or health care institution located in this State participates in curriculum development and the selection of students for admission;
B. No fewer than 10 students per class year are enrolled and in which these students are required to complete not less than one academic year of the medical school curriculum at facilities located in this State;
C. Funds are raised through philanthropic resources and the private sector to match 100% of those funds appropriated or allocated by the State for scholarships under section 12103; and
D. The program curriculum includes required clerkship experiences in and training and course completion in rural health care and primary care.

20-A MRSA §12101 (8-A)

For more on Doctors for Maine’s Future Scholarship Program and Eligibility see 20-A MRSA § 12101 (8-A), 20-A MRSA § 12103-A, or Finance Authority of Maine.

(61) Medical Examiner Act
The Medical Examiner Act consists of many sections related to the Office of the Medical Examiner. Below are several select sections as they relate to medical examiners and and/or medical examiner cases.

Certain information confidential
The following records in the possession or custody of a medical examiner or the Office of the Chief Medical Examiner are not public records within the meaning of Title 1, section 402, subsection 3 and are confidential:
A. Medical records relating to a medical examiner case;
B. Law enforcement agency reports or records relating to a medical examiner case;
C. Communications with the Department of the Attorney General relating to a medical examiner case;
D. Communications with the office of a district attorney relating to a medical examiner case;
E. Death certificates and amendments made to the certificates, except for the information for which the medical examiner is responsible, as listed in section 2842, subsection 3, and not ordered withheld by the Attorney General relating to a medical examiner case or missing person;
F. Photographs and transparencies, histological slides, videotapes and other like items relating to a medical examiner case; and
G. Written or otherwise recorded communications that express or are evidence of suicidal intent obtained under section 3028, subsections 4 and 5.

22 MRSA §3022 (8)

Cooperation with research requests
The Office of Chief Medical Examiner shall cooperate with research requests by supplying abstracted data to interested persons consistent with the available resources of the office.

22 MRSA §3022 (10)

Access to or dissemination of confidential records
Access to or dissemination of records made confidential under subsection 8 is limited to:
A. A criminal justice agency for the purpose of the administration of criminal or juvenile justice;
B. A person for whom the Chief Medical Examiner determines access is necessary or desirable to carry out a duty under this Act;
C. A person for whom the Chief Medical Examiner determines access is necessary or desirable to allow for the harvesting of a decedent's organs and other tissues;
D. A person when authorized or required under any state or federal law, rule or regulation; and
E. A person pursuant to a court order.
Access to or dissemination of records as provided under paragraphs A to C can be done as a matter of course by the Chief Medical Examiner unless the Attorney General directs otherwise.

22 MRSA §3022 (12)
Access to certain information by certain persons

Unless a medical examiner case is under investigation by the Department of the Attorney General or the office of a district attorney and the Attorney General or the district attorney determines that there is a reasonable possibility that release or inspection interferes with a criminal investigation or prosecution by the disclosure:

A. Items identified in subsection 8, paragraphs F and G may be inspected and copies obtained, upon payment of any required fee under section 3035, by:
   1. A next of kin of the deceased, as defined under section 2843-A. The Chief Medical Examiner may provide the original of the items described in subsection 8, paragraph G to the next of kin or other person to whom that item is addressed or directed;
   2. An insurer that may be responsible for payment of benefits as a result of a death if relevant to the payment obligation;
   3. An attorney representing the estate of the decedent or the decedent's property if relevant to the representation; and
   4. An attorney representing a person or a person's estate and exploring a possible civil action against the estate of the decedent if relevant to the representation; and

B. A person may inspect and obtain a copy of communications identified in subsection 8, paragraphs C and D, except work product as defined in Rule 16(b)(3) of the Maine Rules of Criminal Procedure, as long as the communications would otherwise be open to inspection and release if in the possession or custody of the Department of the Attorney General or the office of a district attorney.

Testing for HIV

Notwithstanding Title 5, chapter 501, the Chief Medical Examiner in a medical examiner case may test for the human immunodeficiency virus and may disclose the test result as authorized under subsection 12. As used in subsections 10, 12, 13 and 14, "person" means a natural person, including a public servant, or a corporation, partnership, unincorporated association or other legal entity, including a governmental unit.

Medical examiner case

Circumstances of death that must be reported

A medical examiner case may exist and must be reported as provided in section 3026 when remains are found that may be human and raise suspicion that death has occurred under any of the following circumstances:

A. Death is suspected of having been caused by any type of physical injury, including poisoning, regardless of whether the suspected manner of death is homicide, suicide or accident. This circumstance must be reported irrespective of whether the deceased had been attended by a physician, was a patient in a hospital, survived for a considerable time following the physical injury or died from terminal natural causes consequent to and following the physical injury;

B. Suddenly when the person is in apparent good health and has no specific natural disease sufficient to explain death

C. During diagnostic or therapeutic procedures under circumstances indicating gross negligence or when clearly due to trauma or poisoning unrelated to the ordinary risks of those procedures;

D. Death when the person is in custody pursuant to an arrest, confined in a state correctional facility, county institution, facility or local lockup, unless clearly certifiable by an attending physician as due to specific natural causes;

E. Death while the person is a patient or resident of a facility of the Department of Behavioral and Developmental Services or residential care facility maintained or licensed by the Department of Human Services, unless clearly certifiable by an attending physician as due to specific natural causes;
F. Death suspected of being due to a threat to the public health when the authority of the medical examiner is needed to adequately study the case for the protection of the public health;

G. Death suspected of not having been certified, including, but not limited to, bodies brought into the State and any buried remains uncovered other than by legal exhumation;

H. Deaths suspected of being medical examiner cases which may have been improperly certified or inadequately examined, including, but not limited to, bodies brought into the State under those circumstances;

I. Sudden infant death syndrome deaths and all other deaths of children under the age of 18 unless clearly certifiable by an attending physician as due to specific natural causes unrelated to abuse or neglect;

J. Whenever human or possibly human remains are discovered not properly interred or disposed of, for which the responsibility to do so cannot be readily determined; or

K. Any cause when there is no attending physician capable of certifying the death as due to natural causes. When a person dies who is under the care of a religious practitioner who uses prayer and spiritual means of healing, the fact that the deceased has been under such religious care does not warrant suspicion of foul play or investigation beyond that warranted by the other facts of the case.

*In any case in which the necessity of a report is questionable, a report must be made.  

22 MRSA §3025 (1)

Medical examiner case determination

Notwithstanding that a case must be reported under subsection 1, the acceptance of any reported death as a medical examiner case is to be determined by the Chief Medical Examiner unless acceptance is specifically ordered by the Attorney General or district attorney having jurisdiction.

The following deaths that must be reported need not be accepted by the Chief Medical Examiner as a medical examiner case:

A. Deaths due to the consequences of long-term alcohol use, long-term exposure to environmental or occupational toxins or long-term exposure to carcinogens;

B. Deaths in the elderly who have sustained limb or axial fractures, excluding the head, for which they are or have been hospitalized; or

C. Sudden natural deaths in the elderly who have not had previous specific symptoms or who were not under treatment by a physician for the specific natural cause that is considered to be the cause of death

These reportable deaths may be referred back to the attending physician by the Chief Medical Examiner for certification of the death, even though the attending physician has not treated the patient for the specific natural disease that the attending physician will enter as the physician's diagnosis.

22 MRSA §3025 (1-A)

Transplant operations

No operation for the transplant of an organ or a portion of any organ may take place, when the donor's death occurs under circumstances indicating a medical examiner case, without approval of the medical examiner. Any doctor performing a transplant operation when the donor has died under these circumstances shall note the condition of the vital organs in the region of surgery and shall include this notation in a written report of the operation and manner in which death was pronounced, with the report to be given to the medical examiner upon his request. The medical examiner may choose to be present during the removal of the donated organ.

22 MRSA §3025 (3)

Delayed reports

When a death has occurred that falls under this law as a medical examiner case and the body has already been released for final disposition, the case may be accepted and the body ordered held for examination by a medical examiner, but no exhumation may take place when the body has been finally interred, except pursuant to section 3029.
Reports of death

Electronic death registration system
Death certificates must be submitted using the State Registrar of Vital Statistics electronic system. Does not apply to “authorized person” under section 2846 (family member).

Persons suspecting medical examiner case
Any person who becomes aware of a suspected medical examiner case shall immediately notify a law enforcement officer or the Office of Chief Medical Examiner. As used in this subsection, "person" means a natural person, including a public servant, and a corporation, partnership, unincorporated association or any other nonhuman legal entity, including any governmental unit.

Law enforcement officers suspecting medical examiner case
Any law enforcement officer who becomes aware of a suspected medical examiner case shall immediately notify the Office of Chief Medical Examiner.

Medical examiners suspecting medical examiner case
Any medical examiner who becomes aware of a death caused by physical injury, or in which physical injury is the suspected cause, shall immediately notify the Office of Chief Medical Examiner and the appropriate law enforcement agency. The agency shall notify the district attorney for the district in which the body is located.

Cases involving or suspected of involving physical injury attributable to criminal conduct
Any law enforcement officer or medical examiner who becomes aware of a death involving physical injury attributable to criminal conduct, or in which physical injury attributable to criminal conduct is suspected, other than vehicular manslaughter, in addition to complying with the notification requirements in subsection 3, shall immediately notify the Attorney General.

Procedure at scene of death
Movement or alteration of body prohibited
Except as otherwise provided in this section:
A. In any medical examiner case a person may not move or alter the body or any objects at the scene of death prior to the arrival, or without the express authorization, of the medical examiner or Office of Chief Medical Examiner;
B. In any medical examiner case in which physical injury attributable to noncriminal conduct is suspected or in which any physical injury by motor vehicle, including vehicular manslaughter, is suspected, a person may not move or alter the body or any objects at the scene of death prior to the arrival, or without the express authorization, of the district attorney for the district in which the body is located or the district attorney's authorized representative; and
C. In any medical examiner case in which physical injury attributable to criminal conduct other than vehicular manslaughter is suspected, a person may not move or alter the body or any objects at the scene of death prior to the arrival, or without the express authorization, of the Attorney General or the Attorney General's authorized representative.

Preservation or removal of body
In any medical examiner case in which the body is in danger of being destroyed or lost or the location of the body renders it a serious threat to the safety or health of others, a person may take whatever steps are reasonably necessary for the retention or preservation of the body prior to the arrival or authorization of
the medical examiner or the Office of Chief Medical Examiner. The person shall first, if practicable, exactly mark the location and position of the body.

In any medical examiner case in which physical injury attributable to criminal conduct other than vehicular manslaughter is not suspected and the presence of the body is likely to cause hardship or outrage, and a medical examiner or the Office of Chief Medical Examiner can not be reached in a reasonable period of time, the district attorney for the district in which the body is located or the district attorney's authorized representative may authorize removal of the body by the law enforcement officer in charge of the scene. The officer shall first, if practicable, exactly mark the location and position of the body.

A. When death occurs in a medical facility such as a hospital or an ambulance, the body may be removed to a mortuary under the following conditions:
   1. The incident causing the death did not occur in the medical facility;
   2. The body is transported to a secure place in the same condition as when death occurred; and
   3. The only alterations are the disconnecting of fixed medical equipment.

Procedures
A. Before removal of the body as provided in subsection 2, the law enforcement officer shall whenever possible arrange for photographs, measurements and a record of the location and position of the body.
B. When the death is suspected of involving physical injury attributable to criminal conduct other than vehicular manslaughter, the procedure in this subsection must be undertaken with the supervision of an authorized representative of the Attorney General.
C. In all medical examiner cases in which physical injury attributable to criminal conduct other than vehicular manslaughter is suspected, the procedure in this subsection may be waived concurrently by the Chief Medical Examiner and the Attorney General or the Attorney General's authorized representative.
D. In all other medical examiner cases the procedure in this subsection may be waived concurrently by the medical examiner and the district attorney for the district in which the body is located or the district attorney's authorized representative.

Investigation; autopsy
Authority to conduct investigation
The medical examiner or the person expressly authorized by the Chief Medical Examiner has authority to conduct an investigation and inquiry into the cause, manner and circumstances of death in a medical examiner case. The medical examiner or authorized person shall, if it is determined necessary, immediately proceed to the scene and, subject to the authority of the Attorney General, assume custody of the body for the purposes of the investigation, and shall retain custody until the investigation has been completed or until the Chief Medical Examiner has assumed charge of the case.

Investigation by law enforcement officer
When death is not suspected to be the result of physical injury attributable to criminal conduct, the medical examiner may elect not to proceed to the scene, or the Chief Medical Examiner may elect not to dispatch a medical examiner or the person expressly authorized by the Chief Medical Examiner under subsection 1 to the scene. If the medical examiner elects not to proceed to the scene, or the Chief Medical Examiner elects not to dispatch a medical examiner or authorized person to the scene, the law enforcement officer in charge of the scene shall:
   A. Investigate, take photographs and take possession of useful objects as directed by the medical examiner, authorized person or the Office of Chief Medical Examiner pursuant to subsection 4;
B. Remove the body in accordance with the instructions of the medical examiner, authorized person or the Office of Chief Medical Examiner; and
C. Make a report of the investigation available to the medical examiner, authorized person or the Office of Chief Medical Examiner.

22 MRSA §3028 (2)

**Assistance of law enforcement agency**
The medical examiner, the person expressly authorized by the Chief Medical Examiner or the pathologist as described in subsection 8, may request the assistance and use of the facilities of the law enforcement agency having jurisdiction over the case for the purposes of photographing, fingerprinting or otherwise identifying the body. That agency shall provide the medical examiner, authorized person or pathologist with a written report of the steps taken in providing the assistance.

22 MRSA §3028 (3)

**Possession of useful objects**
Except as otherwise directed by the Attorney General, the Attorney General's deputies or assistants, the medical examiner, the person expressly authorized by the Chief Medical Examiner or the Office of Chief Medical Examiner may direct that a law enforcement officer at the scene make measurements, take photographs and take possession of all objects that in the opinion of the medical examiner, authorized person or the Office of Chief Medical Examiner may be useful in establishing the cause, manner and circumstances of death. For these same purposes, the medical examiner, authorized person or the Office of Chief Medical Examiner may direct that a law enforcement officer take possession of any objects or specimens that have been removed from the victim at the scene or elsewhere while under medical care.

22 MRSA §3028 (4)

**Requests for objects**
Any person having possession of any object or objects, as described in subsection 4, shall at the request of the medical examiner or the person expressly authorized by the Chief Medical Examiner give that object or objects to a law enforcement officer, to the medical examiner, to the authorized person or to the Office of Chief Medical Examiner. Medical personnel and institutions turning over any objects or specimens that have been removed from the victim while under medical care are immune from civil or criminal liability when complying with this subsection. Original written or recorded material that might express suicidal intent must be sent to the Office of the Chief Medical Examiner. The Chief Medical Examiner may elect to accept copies in place of originals.

22 MRSA §3028 (5)

**Examination of body**
In all cases except those requiring a report on a body already disposed of and not to be exhumed for examination, the medical examiner or the person expressly authorized by the Chief Medical Examiner shall conduct a thorough examination of the body.

22 MRSA §3028 (6)

**Written report**
Upon completing an investigation, the medical examiner shall submit a written report of findings to the Chief Medical Examiner on forms provided for that purpose. The medical examiner shall retain one copy of the report.

If a medical examiner reports suspected abuse, neglect or exploitation to the Chief Medical Examiner, the Chief Medical Examiner, by reporting that information to the department on behalf of the medical examiner, fulfills the medical examiner's mandatory reporting requirement under section 3477 or 4011-A.

22 MRSA §3028 (7)

**Autopsy**
If, in any medical examiner case, in the opinion of the medical examiner, the Chief Medical Examiner, the district attorney for the district in which the death has occurred or the Attorney General, it is advisable and in the public interest that an autopsy be made, the autopsy must be conducted by the Chief Medical Examiner or by a physician that the medical examiner, with the approval of the Chief Medical Examiner,
may designate. The medical examiner, with the approval of the Chief Medical Examiner, may elect to perform the autopsy. The person who performs the autopsy shall make a complete report of the findings of the autopsy and shall transmit the report to the medical examiner and the Office of Chief Medical Examiner, retaining one copy of the report.

**Autopsy of child**
In the case of a child under the age of 3 years, when death occurs without medical attendance or, if attended, without a specific natural cause, the medical examiner shall order an autopsy. The autopsy may be waived by the Chief Medical Examiner, as long as the Chief Medical Examiner includes the reason for the waiver in the record.

**Chief Medical Examiner; jurisdiction**
The Chief Medical Examiner may assume jurisdiction over a medical examiner case and may recertify the death when the Chief Medical Examiner finds that it is in the public interest to do so. The Chief Medical Examiner shall include the reasons for so doing in the record.

**Final release of body**
In any medical examiner case the body shall not be finally released for embalming or burial except by order of the medical examiner in charge of the case, or by the Chief Medical Examiner. No medical examiner may release a body without first ensuring that the case has been reported to the Office of Chief Medical Examiner.

**Report to domestic abuse panel**
If the Chief Medical Examiner determines that a death resulted from criminal conduct and that the victim was pregnant at the time of death, the Chief Medical Examiner shall send a copy of any report prepared under this section to the Domestic Abuse Homicide Review Panel created pursuant to Title 19-A, section 4013.

**Disposal of unidentified or abandoned human remains**
Whenever unidentified human remains are recovered, the Chief Medical Examiner may store the remains, release them to an educational institution, inter them in an appropriate resting place or have them cremated. Ashes of remains cremated may be disposed of in any appropriate manner. Human remains uncovered in a cared-for cemetery or known to be Indian remains are excluded from the operation of this section.

The Chief Medical Examiner may assume responsibility for the disposal of identified human remains of a deceased resident of this State that are the subject of a medical examiner case if no one takes custody and control of the human remains for a period of 30 days after the Chief Medical Examiner has both completed an autopsy or necessary examination of the human remains and made reasonable inquiry under section 3028-D, subsection 1. Such abandoned remains may be interred or cremated. The Chief Medical Examiner shall file or cause to be filed a certificate of abandonment in the municipality where the human remains were recovered that indicates the means of disposal.

In the absence of a responsible party, payment of expenses incurred by the Chief Medical Examiner pursuant to this section must be made pursuant to section 3028-D, subsection 2 as if the remains were unidentified. The Chief Medical Examiner may seek to recover costs from the estate or municipality of residence of the deceased.

**22 MRSA §3028 (8)**

**22 MRSA §3028 (9)**

**22 MRSA §3028 (10)**

**22 MRSA §3028 (11)**

**22 MRSA §3028 (12)**

**22 MRSA §3028-A**
(62) Medical-Legal Cooperation Code

Members of the Maine State Bar Association (MSBA), the Maine Medical Association (MMA), the Maine Osteopathic Association (MOA) and the Maine Chiropractic Association (MCA) drafted the Medical-Legal Cooperation Code in recognition of the need for co-operation between the medical and legal professions and the duties of both professions to the public and to the administration of justice. The code outlines the duties of both physicians and lawyers in civil proceedings in situations such as providing medical reports and records; providing court testimony; depositions; and billing. In case of disputes regarding the applicability or interpretation of the code, a Medical-Legal Cooperation Committee shall be formed made up of the parties to the Code to mediate the dispute.

The full code can be found here.

(63) Medical Malpractice (see Professional Liability)

(64) Medical Marijuana

Maine Medical Marijuana Act

Maine first enacted a medical marijuana law at referendum in 1999. This law was repealed and replaced by a citizen referendum in 2009 and legislation enacted by the 124th Legislature in 2010. It has been amended several times since.

You can find the Maine Medical Marijuana Act at Title 22, Chapter 558-C. The Department of Health & Human Services has information about the Maine Medical Marijuana Act on the web at: http://www.maine.gov/dhhs/dlrs/mmm/.

The rules governing the program have been updated effective September 2013. They include more information about qualifying conditions, physicians’ written certifications and other details of the program.

Qualifying medical conditions

There is no longer any statutory limitation on the conditions for which a patient may receive a written certification for the use of medical marijuana. A medical provider may issue such a certification to a “qualifying patient” if the provider believes marijuana will have a therapeutic or palliative effect on the patient’s diagnosis.

22 MSRA § 2423-B

DHHS is required to adopt rules regarding the consideration of petitions from the public to add medical conditions or treatments to the list of debilitating medical conditions.

22 MSRA § 2424(2)

Written certification

"Written certification" means a document on tamper-resistant paper signed by a medical provider, that expires within one year and that states that in the medical provider’s professional opinion a patient is likely to receive therapeutic or palliative benefit from the medical use of marijuana to treat or alleviate the patient's medical diagnosis or symptoms associated with the medical diagnosis.

22 MSRA § 2422(16)

Authorized conduct by a physician or nurse practitioner

As of 2014, both physicians and nurse practitioners may issue medical marijuana certificates to qualifying patients.

A physician may provide a written certification for the medical use of marijuana and, after having done so, may state that in the physician's professional opinion a qualifying patient is likely to receive
therapeutic or palliative benefit from the medical use of marijuana to treat or alleviate the patient's medical diagnosis.

22 MSRA § 2423-B

Adult qualifying patient
Prior to providing written certification for the medical use of marijuana under this section, a physician shall inform an adult qualifying patient or the patient’s legal guardian or representative of the risks and benefits of the medical use of marijuana and that the patient may benefit from the medical use of marijuana.

22 MSRA § 2423-B(1)

Minor qualifying patient
Prior to providing written certification for the medical use of marijuana by a minor qualifying patient under this section, a medical provider shall inform the minor qualifying patient and the parent or legal guardian or person having legal custody of the patient of the risks and benefits of the medical use of marijuana and that the patient may benefit from the medical use of marijuana. Except with regard to a minor qualifying patient who is eligible for hospice care or has a diagnosis of epilepsy, cancer, a developmental disability or an intellectual disability, prior to providing a written certification under this section, the treating medical provider shall consult with a qualified physician, referred to in this paragraph as "the consulting physician," from a list of physicians who may be willing to act as consulting physicians maintained by the department that is compiled by the department after consultation with statewide associations representing licensed medical professionals. The consultation between the treating medical provider and the consulting physician may consist of examination of the patient or review of the patient's medical file. The consulting physician shall provide an advisory opinion to the treating medical provider and the parent or legal guardian or person having legal custody of the minor qualifying patient concerning whether the patient is likely to receive therapeutic or palliative benefit from the medical use of marijuana to treat or alleviate the patient's medical diagnosis. If the department or the consulting physician does not respond to a request by a treating medical provider within 10 days of receipt of the request, the treating medical provider may provide written certification for treatment without consultation with another physician.

22 MSRA § 2423-B(2-A)

Patient with substance use disorder
Prior to providing certification for therapeutic or palliative use for substance use disorder, the provider shall develop a recovery plan with the patient.

22 MSRA § 2423-B(2-B)

Bona fide provider-patient relationship
A written certification may be made only in the course of a bona fide medical provider-patient relationship after the medical provider has completed a full assessment of the patient's medical history. If a patient has not provided a medical provider who is not the patient's primary care provider with the name and contact information of the patient's primary care provider, a medical provider shall conduct an in-person consultation with the patient prior to providing a written certification.

22 MSRA § 2423-B(2-C)

Expiration
A written certification form for the medical use of marijuana under this section is valid for the term provided by the qualifying patient's medical provider.

22 MSRA § 2423-B(3)

Form; content
A written certification under this section must be in the form required by rule adopted by the department and may not require a qualifying patient's medical provider to state the patient's specific medical condition.

22 MSRA § 2423-B(4)
Possible sanctions
Nothing in this chapter prevents a professional licensing board from sanctioning a medical provider for failing to properly evaluate or treat a patient's medical diagnosis or otherwise violating the applicable standard of care for evaluating or treating medical diagnoses.  

22 MSRA § 2423-B(5)

Certification issued based on debilitating condition
A medical provider may not condition the issuance of a certification for the medical use of marijuana on any requirements other than that the patient's medical diagnosis may be alleviated by the therapeutic or palliative medical use of marijuana. Nothing in this section may be construed to prevent a medical provider from exercising professional judgment in declining to issue a certification for the medical use of marijuana.

22 MRSA §2423-B(6)

Patient referral disclosure of interest
Prior to providing a referral to a qualifying patient for goods and services associated with a certification for the medical use of marijuana to an entity in which the medical provider has a direct or indirect financial interest, a medical provider shall provide written disclosure to the qualifying patient regarding any direct or indirect financial interest the medical provider has or may have in the resulting referral and shall maintain a copy of this disclosure in the qualifying patient's record.

22 MRSA §2423-B(7)

Continuing medical education
A medical provider who has not previously provided a written certification to a qualifying patient for the medical use of marijuana shall, prior to providing a written certification to a qualifying patient, submit evidence, satisfactory to the department, of successful completion of a one-hour course of continuing medical education relating to medical marijuana within the preceding 24 months.

22 MRSA §2423-B(8)

Medical Marijuana in Schools
A caregiver, parent, guardian or person with legal custody as defined in statute may possess and administer marijuana in a school bus and on the grounds of the preschool or primary or secondary school in which a qualifying patient is enrolled only if:

A. A medical provider has provided the qualifying patient with a current written certification for the medical use of marijuana under this chapter; and
B. Possession of harvested marijuana is for the purpose of administering marijuana to the qualifying patient;
C. That person has notified the school that a caregiver has been designated on behalf of the patient to possess and administer harvested marijuana to the qualifying patient.

Harvested marijuana possessed or administered under this subsection may not be in a form that permits the patient to engage in smoking as defined in Section 1541, except that “smoking” does not include use of a nebulizer.

22 MRSA §2426 (1-A)

General Information
Some risks remain for physicians in treating medical marijuana patients, even in the 31 states (and the District of Columbia) that have enacted medical marijuana legislation:

A. The uncertain status of marijuana under federal law. Use of marijuana remains illegal under federal law. During the Bush Administration, it was never clear how aggressive federal authorities would be towards so-called "medical marijuana," although their focus seemed to be on dispensaries in California that were perceived by some to be pushing the limits of the law. During the Obama administration, Attorney General Eric Holder issued a statement indicating that the Obama Administration had no desire to pursue those involved in legitimate use of medical marijuana under state law. As of this writing, Attorney General Jeff Sessions has expressed an interest in more aggressive federal action against both medical and recreational marijuana.
B. **Drug regulatory concerns.** The standard for prescribing drugs by practitioners is FDA approval. Because marijuana, even for medical purposes, is not approved by the FDA, the MMA staff recommends that you not use the term "prescribe" with respect to medical marijuana and that you not use a prescription blank with such patients. Should you decide to work with medical marijuana patients, it is better framed as a matter of patient choice - that the physician is willing to work with the medical marijuana patient as he or she uses marijuana for medical purposes.

C. **Liability concerns.** While the degree of risk may be debatable, physicians who work with medical marijuana patients cannot eliminate all risk because they are treating a patient who is using an unregulated drug, a drug that does not meet the standard in this country of FDA approval and that may include unknown amounts of active ingredient and impurities. Again, the MMA staff recommends that you include in your documentation of the relationship with the medical marijuana patient that he or she is aware of and assumes responsibility for these risks.

(65) **Medical Records - Access**

In the last several years, in addition to issues of privacy and confidentiality of medical information, access to medical records has been a huge topic on both the state and federal levels. On the federal side there are compliance requirements under the Health Insurance and Portability and Accounting Act (HIPAA). There are also requirements under state law. Federal and state laws are not mutually exclusive and in many ways the two overlap and complement each other. **This section only covers the medical record access requirements under Maine law.**

**Patient access to hospital medical records**

If a patient of an institution licensed as a hospital by the State, after discharge from such institution, makes written request for copies of the patient's medical records, the copies must, if available, be made available to the patient within a reasonable time unless, in the opinion of the hospital, it would be detrimental to the health of the patient to obtain the records. If the hospital is of the opinion that release of the records to the patient would be detrimental to the health of the patient, the hospital shall advise the patient that copies of the records will be made available to the patient's authorized representative upon presentation of a proper authorization signed by the patient. The hospital may exclude from the copies of medical records released any information related to a clinical trial sponsored, authorized or regulated by the federal Food and Drug Administration.

If an authorized representative for a patient requests, in writing, that a hospital provide the authorized representative with a copy of the patient's medical records and presents a proper authorization from the patient for the release of the information, copies must be provided to the authorized representative within a reasonable time.

A written request or authorization for release of medical records under this section satisfies the requirements of section 1711-C, subsection 3.

A patient or, if the patient is a minor who has not consented to health care treatment in accordance with the laws of this State, the minor's parent, legal guardian or guardian ad litem may submit to a hospital health care information that corrects or clarifies the patient's treatment record, which must be retained with the medical record by the hospital. If the hospital adds to the medical record a statement in response to the submitted correction or clarification, the hospital shall provide a copy to the patient or, if the patient is a minor who has not consented to health care treatment in accordance with the laws of this State, the minor's parent, legal guardian or guardian ad litem.

Reasonable costs incurred by the hospital in making and providing paper copies of medical records and additions to medical records may be assessed as charges to the requesting person and the hospital may
require payment prior to responding to the request. The charge for paper copies of records may not exceed $5 for the first page and 45¢ for each additional page, up to a maximum of $250 for the entire medical record.

If a medical record exists in a digital or electronic format, the hospital shall provide an electronic copy of the medical record if an electronic copy is requested and it is reasonably possible to provide it. The hospital may assess as charges reasonable actual costs of staff time to create or copy the medical record and the costs of necessary supplies and postage. Actual costs may not include a retrieval fee or the costs of new technology, maintenance of the electronic record system, data access or storage infrastructure. Charges assessed under this paragraph may not exceed $150.

Release of a patient’s medical records to a person other than the patient or, if the patient is a minor who has not consented to health care treatment with the laws of this State, the minor’s parent, legal guardian or guardian ad litem is governed by section 1711-C.

22 MRSA §1711

Fees charged for physician records
Whenever health care practitioner defined in section 1711-B furnishes requested copies of a patient’s treatment record or a medical report or an addition to a treatment record or medical report to the patient or the patient’s authorized representative, the charge for the copies or the report may not exceed the reasonable costs incurred by the health care practitioner in making and providing the copies or the report. The charge for copies of records may not exceed $5.00 for the first page and $.45 for each additional page. For paper records charges must not exceed $250.00. Charges include reasonable actual cost of staff time and cannot include retrieval fee, cost of new technology, maintenance, data access or storage fees.

In addition, if a medical record exists in a digital or electronic format, the practitioner shall provide an electric copy of the medical record if an electrical copy is requested and it is reasonably possible to provide it. The practitioner may assess as charges reasonable actual cost of staff time to create or copy the medical record and the cost of necessary supplies and postage. Actual cost may not include a retrieval fee or the cost of new technology, maintenance of the electronic record system, data access or storage infrastructure. Charges assessed for digital and electronic format may not exceed $150.00.

22 MRSA §1711-A

Patient access to treatment records; health care practitioners
Access
Upon written authorization executed in accordance with section 1711-C, subsection 3, a health care practitioner shall release copies of all treatment records of a patient or a narrative containing all relevant information in the treatment records to the patient. The health care practitioner may exclude from the copies of treatment records released any personal notes that are not directly related to the patient's past or future treatment and any information related to a clinical trial sponsored, authorized or regulated by the federal Food and Drug Administration. The copies or narrative must be released to the designated person within a reasonable time.

If the practitioner believes that release of the records to the patient is detrimental to the health of the patient, the practitioner shall advise the patient that copies of the treatment records or a narrative containing all relevant information in the treatment records will be made available to the patient's authorized representative upon presentation of a written authorization signed by the patient. The copies or narrative must be released to the authorized representative within a reasonable time. Except as provided in subsection 3, release of a patient's treatment records to a person other than the patient is governed by section 1711-C.

22 MRSA §1711-B (2)
Person receiving the records
Except as otherwise provided, records must be released to:

A. The person who is the subject of the treatment record, if that person is 18 years of age or older and mentally competent;
B. The parent, guardian ad litem or legal guardian of the person who is the subject of the record if the person is a minor, or the legal guardian if the person who is the subject of the record is mentally incompetent;
C. The designee of a durable health care power of attorney executed by the person who is the subject of the record, at such time as the power of attorney is in effect;
D. The agent, guardian or surrogate pursuant to the Uniform Healthcare Decisions Act; or
E. The lay caregiver designated pursuant to section 1711-G by the person who is the subject of the record.

22 MRSA §1711-B (3)

Corrections and clarifications of treatment records
A patient or, if the patient is a minor who has not consented to health care treatment in accordance with the laws of this State, the minor's parent, legal guardian or guardian ad litem may submit to a health care practitioner health care information that corrects or clarifies the patient's treatment record, which must be retained with the treatment record by the health care practitioner. If the health care practitioner adds to the treatment record a statement in response to the submitted correction or clarification, the health care practitioner shall provide a copy to the patient or, if the patient is a minor who has not consented to health care treatment in accordance with the laws of this State, the minor's parent, legal guardian or guardian ad litem.

22 MRSA §1711-B (3-A)

Minors
This section does not affect the right of minors to have their treatment records treated confidentially pursuant to the provisions of chapter 260.

22 MRSA §1711-B (4)

HIV test
Release of information regarding the HIV infection status of a patient is governed by Title 5, section 19203-D.

22 MRSA §1711-B (5)

Retention of records
This section does not alter the existing law or ethical obligations of a health care practitioner with respect to retaining treatment records.

22 MRSA §1711-B (6)

(66) Medical Records - Confidentiality and Privilege

See Also, Health Insurance Portability and Accountability Act

Maine Confidentiality Law
Maine has a number of statutory provisions governing the confidentiality of health care information. While these laws are largely consistent with federal law (HIPAA and others), there are times when they conflict and state law will govern if it is more protective of patient rights. Please consult with your attorney or contact the Maine Medical Association legal team if you have questions about state and federal confidentiality laws. Maine’s primary law on confidentiality of medical records is found at 22 MRSA §1711-C.

Definitions
Definitions in the statute are found at 22 MRSA §1711-C (1).
Confidentiality of health information; disclosure
An individual's health care information is confidential and may not be disclosed other than to the individual by the health care practitioner or facility except as provided in subsections 3, 3-A, 3-B, 6, or 11 of this section of law. Nothing in this section prohibits a health care practitioner or health care facility from adhering to applicable ethical or professional standards provided that these standards do not decrease the protection of confidentiality granted by this section.

Written authorization to disclose
A health care practitioner or facility may disclose health care information pursuant to a written authorization signed by an individual for the specific purpose stated in the authorization. A written authorization to disclose health care information must be retained with the individual's health care information. A written authorization to disclose is valid whether it is in an original, facsimile or electronic form. A written authorization to disclose must contain the following elements:
A. The name and signature of the individual and the date of signature. If the authorization is in electronic form, a unique identifier of the individual and the date the individual authenticated the electronic authorization must be stated in place of the individual's signature and date of signature;
B. The types of persons authorized to disclose health care information and the nature of the health care information to be disclosed;
C. The identity or description of the 3rd party to whom the information is to be disclosed;
D. The specific purpose or purposes of the disclosure and whether any subsequent disclosures may be made pursuant to the same authorization. An authorization to disclose health care information related to substance abuse treatment or care subject to the requirements of 42 United States Code, Section 290dd-2 (Supplement 1998) is governed by the provisions of that law;
E. The duration of the authorization;
F. A statement that the individual may refuse authorization to disclose all or some health care information but that refusal may result in improper diagnosis or treatment, denial of coverage or a claim for health benefits or other insurance or other adverse consequences;
G. A statement that the authorization may be revoked at any time by the individual by executing a written revocation, subject to the right of any person who acted in reliance on the authorization prior to receiving notice of revocation, instructions on how to revoke an authorization and a statement that revocation may be the basis for denial of health benefits or other insurance coverage or benefits; and
H. A statement that the individual is entitled to a copy of the authorization form.

Oral authorization to disclose
When it is not practical to obtain written authorization under subsection 3 from an individual or person acting pursuant to subsection 3-B or when a person chooses to give oral authorization to disclose, a health care practitioner or facility may disclose health care information pursuant to oral authorization. A health care practitioner or facility shall record with the individual's health care information receipt of oral authorization to disclose, including the name of the authorizing person, the date, the information and purposes for which disclosure is authorized and the identity or description of the 3rd party to whom the information is to be disclosed.

Authorization to disclose provided by a 3rd party
When an individual or an authorized representative is unable to provide authorization to disclose under subsection 3 or 3-A, a health care practitioner or facility may disclose health care information pursuant to authorization to disclose that meets the requirements of subsection 3 or 3-A given by a 3rd party listed in this subsection. A health care practitioner or facility may determine not to obtain authorization from a person listed in this subsection when the practitioner or facility determines it would not be in the best interest of the individual to do so. In making this decision, the health care practitioner or facility shall
respect the safety of the individual and shall consider any indicators, suspicion or substantiation of abuse.

Persons who may authorize disclosure under this subsection include:

A. The spouse of the individual;
B. A parent of the individual;
C. An adult who is a child, grandchild or sibling of the individual;
D. An adult who is an aunt, uncle, niece or nephew of the individual, related by blood or adoption;
E. An adult related to the individual, by blood or adoption, who is familiar with the individual's personal values; and
F. An adult who has exhibited special concern for the individual and who is familiar with the individual's personal values.

**22 MRSA §1711-C (3-B)**

Duration of authorization to disclose
An authorization to disclose may not extend longer than 30 months, except that the duration of an authorization for the purposes of insurance coverage under Title 24, 24-A or 39-A is governed by the provisions of Title 24, 24-A or 39-A, respectively.

**22 MRSA §1711-C (4)**

Revocation of authorization to disclose
A person who may authorize disclosure may revoke authorization to disclose at any time, subject to the rights of any person who acted in reliance on the authorization prior to receiving notice of revocation. A written revocation of authorization must be signed and dated. If the revocation is in electronic form, a unique identifier of the individual and the date the individual authenticated the electronic authorization must be stated in place of the individual's signature and date of signature. A health care practitioner or facility shall record receipt of oral revocation of authorization, including the name of the person revoking authorization and the date. A revocation of authorization must be retained with the authorization and the individual's health care information.

**22 MRSA §1711-C (5)**

Disclosure without authorization to disclose
A health care practitioner or facility may disclose, or when required by law must disclose, health care information without authorization to disclose under the circumstances stated in this subsection or as provided in subsection 11. Disclosure may be made without authorization as follows:

A. To another health care practitioner or facility for diagnosis, treatment or care of individuals or to complete the responsibilities of a health care practitioner or facility that provided diagnosis, treatment or care of individuals, as provided in this paragraph.

1. For a disclosure within the office, practice or organizational affiliate of the health care practitioner or facility, no authorization is required.
   For a disclosure outside of the office, practice or organizational affiliate of the health care practitioner or facility, authorization is not required, except that in nonemergency circumstances authorization is required for health care information derived from mental health services provided by:
   a. A clinical nurse specialist licensed under the provisions of Title 32, chapter 31;
   b. A psychologist licensed under the provisions of Title 32, chapter 56;
   c. A social worker licensed under the provisions of Title 32, chapter 83;
   d. A counseling professional licensed under the provisions of Title 32, chapter 119; or
   e. A physician specializing in psychiatry licensed under the provisions of Title 32, chapter 36 or 48.

This subparagraph does not prohibit the disclosure of health care information between a licensed pharmacist and a health care practitioner or facility providing mental health services for the purpose of dispensing medication to an individual;
This subparagraph does not prohibit the disclosure without authorization of health care information covered under this section to a state-designated statewide health information exchange that satisfies the requirement in subsection 18, paragraph C of providing a general opt-out provision to an individual at all times and that provides and maintains an individual protection mechanism by which an individual may choose to opt in to allow the state-designated statewide health information exchange to disclose that individual's health care information covered under Title 34-B, section 1207;

This subparagraph does not prohibit the disclosure without authorization of health care information covered under this paragraph to a health care practitioner or health care facility, or to a payor or person engaged in payment for health care, for purposes of care management or coordination of care. Disclosure of psychotherapy notes is governed by 45 Code of Federal Regulations, Section 164.508(a)(2). A person who has made a disclosure under this subparagraph shall make a reasonable effort to notify the individual or the authorized representative of the individual of the disclosure.

B. To an agent, employee, independent contractor or successor in interest of the health care practitioner or facility including a state-designated statewide health information exchange that makes health care information available electronically to health care practitioners and facilities or to a member of a quality assurance, utilization review or peer review team to the extent necessary to carry out the usual and customary activities relating to the delivery of health care and for the practitioner's or facility's lawful purposes in diagnosing, treating or caring for individuals, including billing and collection, risk management, quality assurance, utilization review and peer review. Disclosure for a purpose listed in this paragraph is not a disclosure for the purpose of marketing or sales;

C. To a family or household member unless expressly prohibited by the individual or a person acting pursuant to subsection 3-B;

D. To appropriate persons when a health care practitioner or facility that is providing or has provided diagnosis, treatment or care to the individual in good faith believes that disclosure is made to avert a serious threat to health or safety and meets the conditions, as applicable, described in 45 Code of Federal Regulations, Section 164.512(j) (2012). A disclosure pursuant to this paragraph must protect the confidentiality of the health care information consistent with sound professional judgment;

E. To federal, state or local governmental entities in order to protect the public health and welfare when reporting is required or authorized by law, to report a suspected crime against the health care practitioner or facility or to report information that the health care facility's officials or health care practitioner in good faith believes constitutes evidence of criminal conduct that occurred on the premises of the health care facility or health care practitioner;

E-1. To federal, state or local governmental entities if the health care practitioner or facility that is providing diagnosis, treatment or care to an individual has determined in the exercise of sound professional judgment that the following requirements, as applicable, are satisfied:

1. With regard to a disclosure for public health activities, for law enforcement purposes or that pertains to victims of abuse, neglect or domestic violence, the provisions of 45 Code of Federal Regulations, Section 164.512(b), (c) or (f) (2012) must be met; and

2. With regard to a disclosure that pertains to a victim of domestic violence or a victim of sexual assault, the provisions of 45 Code of Federal Regulations, Section 164.512(c)(1)(iii)(A) (2012) and Section 164.512(c)(1)(iii)(B)(2012) must be met.

E-2. To federal, state or local governmental entities if the health care practitioner or facility that is providing diagnosis, treatment or care to an individual has determined in the exercise of sound professional judgment that the disclosure is required by section 1726;

F-1. As directed by order of a court or as authorized or required by statute;
F-2. To a governmental entity pursuant to a lawful subpoena requesting health care information to which the governmental entity is entitled according to statute or rules of court.

G. To a person when necessary to conduct scientific research approved by an institutional review board or by the board of a nonprofit health research organization or when necessary for a clinical trial sponsored, authorized or regulated by the federal Food and Drug Administration. A person conducting research or a clinical trial may not identify any individual patient in any report arising from the research or clinical trial. For the purposes of this paragraph, “institutional review board” means any board, committee or other group formally designated by a health care facility and authorized under federal law to review, approve or conduct periodic review of research programs. Health care information disclosed pursuant to this paragraph that identifies an individual must be returned to the health care practitioner or facility from which it was obtained or must be destroyed when it is no longer required for the research or clinical trial. Disclosure for a purpose listed in this paragraph is not a disclosure for the purpose of marketing or sales;

H. To a person engaged in the assessment, evaluation or investigation of the provision of or payment for health care or the practices of a health care practitioner or facility or to an agent, employee or contractor of such a person, pursuant to statutory or professional standards or requirements. Disclosure for a purpose listed in this paragraph is not a disclosure for the purpose of marketing or sales;

I. To a person engaged in the regulation, accreditation, licensure or certification of a health care practitioner or facility or to an agent, employee or contractor of such a person, pursuant to standards or requirements for regulation, accreditation, licensure or certification;

J. To a person engaged in the review of the provision of health care by a health care practitioner or facility or payment for such health care under Title 24, 24-A or 39-A or under a public program for the payment of health care or professional liability insurance for a health care practitioner or facility or to an agent, employee or contractor of such a person;

K. To attorneys for the health care practitioner or facility that is disclosing the health care information or to a person as required in the context of legal proceedings or in disclosure to a court or governmental entity, as determined by the practitioner or facility to be required for the practitioner's or facility's own legal representation;

L. To a person outside the office of the health care practitioner or facility engaged in payment activities, including but not limited to submission to payors for the purposes of billing, payment, claims management, medical data processing, determination of coverage or adjudication of health benefit or subrogation claims, review of health care services with respect to coverage or justification of charges or other administrative services. Payment activities also include but are not limited to:
   1. Activities necessary to determine responsibility for coverage;
   2. Activities undertaken to obtain payment for health care provided to an individual; and
   3. Quality assessment and utilization review activities, including precertification and preauthorization of services and operations or services audits relating to diagnosis, treatment or care rendered to individuals by the health care practitioner or facility and covered by a health plan or other payor;

M. To schools, educational institutions, youth camps licensed under section 2495, correctional facilities, health care practitioners and facilities, providers of emergency services or a branch of federal or state military forces, information regarding immunization of an individual;

N. To a person when disclosure is needed to set or confirm the date and time of an appointment or test or to make arrangements for the individual to receive those services;

O. To a person when disclosure is needed to obtain or convey information about prescription medication or supplies or to provide medication or supplies under a prescription;

P. To a person representing emergency services, health care and relief agencies, corrections facilities or a branch of federal or state military forces, of brief confirmation of general health status;
Q. To a member of the clergy, of information about the presence of an individual in a health care facility, including the person's room number, place of residence and religious affiliation unless expressly prohibited by the individual or a person acting pursuant to subsection 3-B;

R. To a member of the media who asks a health care facility about an individual by name, of brief confirmation of general health status unless expressly prohibited by the individual or a person acting pursuant to subsection 3-B;

S. To a member of the public who asks a health care facility about an individual by name, of the room number of the individual and brief confirmation of general health status unless expressly prohibited by the individual or a person acting pursuant to subsection 3-B; and

T. To a lay caregiver designated by an individual pursuant to section 1711–G.  

Confidentiality policies
A health care practitioner, facility or state-designated statewide health information exchange shall develop and implement policies, standards and procedures to protect the confidentiality, security and integrity of health care information to ensure that information is not negligently, inappropriately or unlawfully disclosed. The policies of health care facilities must provide that an individual being admitted for inpatient care be given notice of the right of the individual to control the disclosure of health care information. The policies must provide that routine admission forms include clear written notice of the individual's ability to direct that that individual's name be removed from the directory listing of persons cared for at the facility and notice that removal may result in the inability of the facility to direct visitors and telephone calls to the individual.

Prohibited disclosure
A health care practitioner, facility or state-designated statewide health information exchange may not disclose health care information for the purpose of marketing or sales without written or oral authorization for the disclosure.

Disclosures of corrections or clarifications to health care information
A health care practitioner or facility shall provide to a 3rd party a copy of an addition submitted by an individual to the individual's health care information if:

A. The health care practitioner or facility provided a copy of the original health care record to the 3rd party on or after February 1, 2000;

B. The correction or clarification was submitted by the individual pursuant to section 1711 or 1711-B and relates to diagnosis, treatment or care;

C. The individual requests that a copy be sent to the 3rd party and provides an authorization that meets the requirements of subsection 3, 3-A or 3-B; and

D. If requested by the health care practitioner or facility, the individual pays to the health care practitioner or facility all reasonable costs requested by that practitioner or facility.

Requirements for disclosures
Except as otherwise provided by law, disclosures of health care information pursuant to this section are subject to the professional judgment of the health care practitioner and to the following requirements:

A. A health care practitioner or facility that discloses health care information pursuant to subsection 3, 3-A or 3-B may not disclose information in excess of the information requested in the authorization.

B. A health care practitioner or facility that discloses health care information pursuant to subsections 3, 3-A, 3-B or 6 may not disclose information in excess of the information reasonably required for the purpose for which it is disclosed.

C. If a health care practitioner or facility believes that release of health care information to the individual would be detrimental to the health of the individual, the health care practitioner or...
facility shall advise the individual and make copies of the records available to the individual's authorized representative upon receipt of a written authorization.

D. If a health care practitioner or facility discloses partial or incomplete health care information, as compared to the request or directive to disclose under subsection 3, 3-A, 3-B or 6, the disclosure must expressly indicate that the information disclosed is partial or incomplete.

22 MRSA §1711-C (10)

Falsification prohibited
A person is guilty of falsifying health care records if, with intent to deceive any person or governmental entity, the person:

A. Makes, or causes to be made, a false material entry in the health care records maintained by a health care provider;

B. Alters, erases, obliterates, deletes, removes or destroys a true material entry in the health care records maintained by a health care provider;

C. Knowingly omits to make a true material entry in the health care records maintained by a health care provider in violation of a duty to do so that is imposed by statute, standard of care or regulatory provision; or

D. Prevents the making of a true material entry or causes the omission of a true material entry in the health care records maintained by a health care provider.

Supplementation of information or correction of an error in health care records in a manner that reasonably discloses that the supplementation or correction was performed and that does not conceal or alter prior entries is not a violation.

17-A MRSA §707-A

Minors
If a minor has consented to health care in accordance with the laws of this State, authorization to disclose health care information pursuant to this section must be given by the minor unless otherwise provided by law.

22 MRSA §1711-C (12)

Enforcement
This section may be enforced within 2 years of the date a disclosure in violation of this section was or should reasonably have been discovered.

A. When the Attorney General has reason to believe that a person has intentionally violated a provision of this section, the Attorney General may bring an action to enjoin unlawful disclosure of health care information.

B. An individual who is aggrieved by conduct in violation of this section may bring a civil action against a person who has intentionally unlawfully disclosed health care information in the Superior Court in the county in which the individual resides or the disclosure occurred. The action may seek to enjoin unlawful disclosure and may seek costs and a forfeiture or penalty under paragraph C. An applicant for injunctive relief under this paragraph may not be required to give security as a condition of the issuance of the injunction.

C. A person who intentionally violates this section is subject to a civil penalty not to exceed $5,000, payable to the State, plus costs. If a court finds that intentional violations of this section have occurred after due notice of the violating conduct with sufficient frequency to constitute a general business practice, the person is subject to a civil penalty not to exceed $10,000 for health care practitioners and $50,000 for health care facilities, payable to the State. A civil penalty under this subsection is recoverable in a civil action.
D. Nothing in this section may be construed to prohibit a person aggrieved by conduct in violation of this section from pursuing all available common law remedies, including but not limited to an action.  

**Waiver prohibited**  
Any agreement to waive the provisions of this section is against public policy and void.  

**Immunity**  
A cause of action in the nature of defamation, invasion of privacy or negligence does not arise against any person for disclosing health care information in accordance with this section. This section provides no immunity for disclosing information with malice or willful intent to injure any person.  

**Application**  
This section applies to all requests, directives and authorizations to disclose health care information executed on or after February 1, 2000. An authorization to disclose health care information executed prior to February 1, 2000 that does not meet the standards of this section is deemed to comply with the requirements of this section until the next health care encounter between the individual and the health care practitioner or facility.  

**Participation in a state-designated statewide health information exchange**  
The following provisions apply to participation in a state-designated statewide health information exchange.  

A. A health care practitioner may not deny a patient health care treatment and a health insurer may not deny a patient a health insurance benefit based solely on the provider's or patient's decision not to participate in a state-designated statewide health information exchange. Except when otherwise required by federal law, a payor of health care benefits may not require participation in a state-designated statewide health information exchange as a condition of participating in the payor's provider network.  

B. Recovery for professional negligence is not allowed against any health care practitioner or health care facility on the grounds of a health care practitioner's or a health care facility's nonparticipation in a state-designated statewide health information exchange arising out of or in connection with the provision of or failure to provide health care services. In any civil action for professional negligence or in any proceeding related to such a civil action or in any arbitration, proof of a health care practitioner's, a health care facility's or a patient's participation or nonparticipation in a state-designated statewide health information exchange is inadmissible as evidence of liability or nonliability arising out of or in connection with the provision of or failure to provide health care services. This paragraph does not prohibit recovery or the admission of evidence of reliance on information in a state-designated statewide electronic health information exchange when there was participation by both the patient and the patient's health care practitioner.  

C. A state-designated statewide health information exchange to which health care information is disclosed under this section shall provide an individual protection mechanism by which an individual may opt out from participation to prohibit the state-designated statewide health information exchange from disclosing the individual's health care information to a health care practitioner or health care facility.  

D. At point of initial contact, a health care practitioner, health care facility or other entity participating in a state-designated statewide health information exchange shall provide to each patient, on a separate form, at minimum:  
   1. Information about the state-designated statewide health information exchange, including a description of benefits and risks of participation in the state-designated statewide health information exchange;
2. A description of how and where to obtain more information about or contact the state-designated statewide health information exchange;

3. An opportunity for the patient to decline participation in the state-designated statewide health information exchange; and

4. A declaration that a health care practitioner, health care facility or other entity may not deny a patient health care treatment based solely on the provider's or patient's decision not to participate in a state-designated statewide health information exchange.

The state-designated statewide health information exchange shall develop the form for use under this paragraph, with input from consumers and providers. The form must be approved by the office of the state coordinator for health information technology within the Governor's office of health policy and finance.

E. A health care practitioner, health care facility or other entity participating in a state-designated statewide health information exchange shall communicate to the exchange the decision of each patient who has declined participation and shall do so within a reasonable time frame, but not more than 2 business days following the receipt of a signed form, as described in paragraph D, from the patient, or shall establish a mechanism by which the patient may decline participation in the state-designated statewide health information exchange at no cost to the patient.

F. A state-designated statewide health information exchange shall process the request of a patient who has decided not to participate in the state-designated statewide health information exchange within 2 business days of receiving the patient's decision to decline, unless additional time is needed to verify the identity of the patient. A signed authorization from the patient is required before a patient is newly entered or reentered into the system if the patient chooses to begin participation at a later date.

Except as otherwise required by applicable law, regulation or rule or state or federal contract, or when the state-designated statewide health information exchange is acting as the agent of a health care practitioner, health care facility or other entity, the state-designated statewide health information exchange shall remove health information of individuals who have declined participation in the exchange. In no event may health information retained in the state-designated statewide health information exchange as set forth in this paragraph be made available to health care practitioners, health care facilities or other entities except as otherwise required by applicable law, regulation or rule or state or federal contract, or when the health care practitioner, health care facility or other entity is the originator of the information.

G. A state-designated statewide health information exchange shall establish a secure website accessible to patients. This website must:

1. Permit a patient to request a report of who has accessed that patient's records and when the access occurred. This report must be delivered to the patient within 2 business days upon verification of the patient's identity by the state-designated statewide health information exchange;

2. Provide a mechanism for a patient to decline participation in the state-designated statewide health information exchange; and

3. Provide a mechanism for the patient to consent to participation in the state-designated statewide health information exchange if the patient had previously declined participation.

H. A state-designated statewide health information exchange shall establish for patients an alternate procedure to that provided for in paragraph F that does not require Internet access. A health care practitioner, health care facility or other entity participating in the state-designated statewide health information exchange shall provide information about this alternate procedure to all patients. The information must be included on the form identified in paragraph D.

I. A state-designated statewide health information exchange shall maintain records regarding all disclosures of health care information by and through the state-designated statewide health
information exchange, including the requesting party and the dates and times of the requests and disclosures.

J. A state-designated statewide health information exchange may not charge a patient or an authorized representative of a patient any fee for access or communication as provided in this subsection.

K. Notwithstanding any provision of this subsection to the contrary, a health care practitioner, health care facility or other entity shall provide the form and communication required by paragraphs D and F to all existing patients following the effective date of this subsection.

L. A state-designated statewide health information exchange shall meet or exceed all applicable federal laws and regulations pertaining to privacy, security and breach notification regarding personally identifiable protected health information, as defined in 45 Code of Federal Regulations, Part 160. If a breach occurs, the state-designated statewide health information exchange shall arrange with its participants for notification of each individual whose protected health information has been, or is reasonably believed by the exchange to have been, breached. For purposes of this paragraph, "breach" has the same meaning as in 45 Code of Federal Regulations, Part 164, as amended.

M. The state-designated statewide health information exchange shall develop a quality management plan, including auditing mechanisms, in consultation with the office of the state coordinator for health information technology within the department, who shall review the plan and results.

**Physician and Psychotherapist Privilege**

A communication is "confidential" if not intended to be disclosed to third persons other than those present to further the interest of the patient in the consultation, examination, or interview, or persons reasonably necessary for the transmission of the communication, or persons who are participating in the diagnosis and treatment under the direction of the physician or psychotherapist, including members of the patient's family.

**Maine Rules of Evidence 503(a)(4)**

**General rule of privilege**

A patient has a privilege to refuse to disclose and to prevent any other person from disclosing confidential communications made for the purpose of diagnosis or treatment of the patient's physical, mental or emotional condition, including alcohol or drug addiction, among the patient, the patient's physician or psychotherapist, and persons who are participating in the diagnosis or treatment under the direction of the physician or psychotherapist, including members of the patient's family.

**Maine Rules of Evidence 503(b)**

**Who may claim the privilege**

The privilege may be claimed by the patient, by the patient's guardian or conservator, or by the personal representative of a deceased patient. The person who was the physician or psychotherapist at the time of the communication is presumed to have authority to claim the privilege but only on behalf of the patient.

**Maine Rules of Evidence 503(d)**

**American Medical Association requirement**

A physician shall…safeguard patient confidence within the constraints of the law. Physicians have an ethical obligation to preserve the confidentiality of information gathered in association with the care of the patient. When disclosing patients’ personal health information, physicians should restrict disclosure to the minimum necessary information and notify the patient of the disclosure, when feasible.

Physicians may disclose personal health information without the specific consent of the patient (or authorized surrogate when the patient lacks decision-making capacity):

- a. To other health care personnel for purposes of providing care or for health care operations;
- b. To appropriate authorities when disclosure is required by law;
c. to other third parties situated to mitigate the threat when in the physician’s judgment there is a reasonable probability that the patient will seriously harm himself or herself or the patient will inflict serious physical harm on an identifiable individual or individuals.

For any other disclosures, physicians should obtain the consent of the patient (or authorized surrogate) before disclosing personal health information.

From AMA Code of Medical Ethics 
Opinion 3.2.1, Confidentiality

(67) Medical Records - Retention

Maine does not currently have a law that governs the length of time that you must retain patient records if you are a private medical practice. Accordingly, the principal guidance is the AMA's ethics opinions and Maine's statute of limitations for bringing lawsuits. The minimum length of time the MMA recommends for record retention is six years. However, Maine hospital licensing regulations specify a seven (7) year retention period and this likely would apply to hospital-based practices. See DHS Rule Chapter 112, Regulations for Licensure of General & Specialty Hospitals, §3.5.5. It is common for physicians to keep records for as long as ten years, and some malpractice carriers recommend this length. For minors, records should be maintained until the age until majority, plus the statute of limitations (3 years), or until the minor turns 21. Maine's hospital licensing regulations require the records of minors to be retained for 6 years beyond the age of majority or age 24. AMA Ethics Opinion 3.3.1, Management of Medical Records states that medical considerations are the primary basis for deciding how long to retain medical records and the Board of Licensure in Medicine has a similar policy. Therefore, physicians should keep a permanent record of information such as chemotherapy, operative notes, and immunization records. Note that some HMO and MCO agreements require departing physicians to retain medical records for a longer period of time than the recommended minimums. To ensure compliance with your contractual obligations, please refer back to your HMO or MCO agreements. Please contact the MMA or view the MMA Physician’s Guide to Closing a Practice for more detailed information on record retention periods.

Record Disposal

As the owner of the medical record, the physician has the obligation to properly store or dispose of the record. The physician has an ethical obligation to notify patients of a move or closure so that they may obtain a copy of their medical records or have them transferred to another practice. Ideally, the physician will notify each patient by letter to the last known address. Often, a written notice is supplemented by publishing a notice in the newspaper on three or more occasions giving patients a reasonable period of time to request records or a transfer of records. This will, hopefully, reach patients for whom the physician does not have an accurate address. See AMA Ethics Opinion 3.3.1, Management of Medical Records and MMA’s Physician’s Guide to Closing a Practice.

(68) Medical Staff

Hospital duties

The governing body of every licensed hospital shall assure that:

A. **Organization of medical staff.** Its medical staff is organized pursuant to written bylaws that have been approved by the governing body

B. **Provider privileges.** Provider privileges extended or subsequently renewed to any physician are in accordance with those recommended by the medical staff as being consistent with that physician's training, experience and professional competence;

C. **Program for identification and prevention of medical injury.** It has a program for the identification and prevention of medical injury which shall include at least the following:

1. One or more professional competence committees with responsibility effectively to review the professional services rendered in the facility for the purpose of insuring quality of medical care of patients therein. Such responsibility shall include a review of
the quality and necessity of medical care provided and the preventability of medical complications and deaths;

2. A grievance or complaint mechanism designed to process and resolve as promptly and effectively as possible grievances by patients or their representatives related to incidents, billing, inadequacies in treatment and other factors known to influence malpractice claims and suits;

3. A system for the continuous collection of data with respect to the provider's experience with negative health care outcomes and incidents injurious to patients, whether or not they give rise to claims, patient grievances, claims, suits, professional liability premiums, settlements, awards, allocated and administrative costs of claims handling, costs of patient injury prevention and safety engineering activities, and other relevant statistics and information; and

4. Education programs for the provider's staff personnel engaged in patient care activities dealing with patient safety, medical injury prevention, the legal aspects of patient care, problems of communication and rapport with patients and other relevant factors known to influence malpractice claims and suits; and

D. **External professional competence committee.** Where the nature, size or location of the health care provider makes it advisable, the provider may, upon recommendation of its medical staff, utilize the services of an external professional competence committee or one formed jointly by 2 or more providers.

**24 MRSA §2503**

**Provider, entity and carrier reports**

A health care provider or health care entity shall, within 60 days, report in writing to the disciplined practitioner's board or authority the name of any licensed, certified or registered employee or person privileged by the provider or entity whose employment or privileges have been revoked, suspended, limited or terminated or who resigned while under investigation or to avoid investigation for reasons related to clinical competence or unprofessional conduct, together with pertinent information relating to that action. Pertinent information includes a description of the adverse action, the date, the location and a description of the event or events giving rise to the adverse action. Upon request, the following information must be released to the board or authority: medical records relating to the event or events; written statements signed or prepared by any witness or complainant to the event; and related correspondence between the practitioner and the provider or entity. The report must include situations in which employment or privileges have been revoked, suspended, limited or otherwise adversely affected by action of the health care practitioner while the health care practitioner was the subject of disciplinary proceedings, and it also must include situations where employment or privileges have been revoked, suspended, limited or otherwise adversely affected by act of the health care practitioner in return for the health care provider or health care entity terminating such proceeding. Any reversal, modification or change of action reported pursuant to this section must be reported immediately to the practitioner's board or authority, together with a brief statement of the reasons for that reversal, modification or change. The failure of any health care provider or health care entity to report as required is a civil violation for which a fine of not more than $5,000 may be adjudged.

Carriers providing managed care plans are subject to the reporting requirements of this section when they take adverse actions against a practitioner's credentials or employment for reasons related to clinical competence or unprofessional conduct that may adversely affect the health or welfare of the patient.

**24 MRSA §2506**

**Effect of filing**

The filing of a report with the board pursuant to this chapter, investigation by the board or any disposition by the board may not, in and of itself, preclude any action by a hospital or other health care facility or health care entity or professional society comprised primarily of physicians to suspend, restrict or revoke the privileges or membership of the physician.
Review committee member immunity
A physician licensed under this chapter who is a member of a utilization review committee, medical
review committee, surgical review committee, peer review committee or disciplinary committee that is a
requirement of accreditation by the Joint Commission on Accreditation of Hospitals or is established and
operated under the auspices of the physician's respective state or county professional society or the Board
of Licensure in Medicine is immune from civil liability for undertaking or failing to undertake an act
within the scope of the function of the committee.

Records of proceedings of medical staff review committees confidential
All proceedings and records of proceedings concerning medical staff reviews, hospital reviews and other
reviews of medical care conducted by committees of physicians and other health care personnel on behalf
of hospitals located within the State or on behalf of individual physicians, when the reviews are required
by state or federal law, rule or as a condition of accreditation by the Joint Commission on Accreditation
of Hospitals or the American Osteopathic Association Committee on Hospital Accreditation or are
conducted under the auspices of the state or county professional society to which the physician belongs,
are confidential and are exempt from discovery.

Provision of information protected by this section to the board pursuant to Title 24, section 2506 does not
waive or otherwise affect the confidentiality of the records or the exemption from discovery provided by
this section for any other purpose.

(69) Medicare Assignment, Posting Acceptance of
An allopathic physician licensed pursuant to chapter 48, an osteopathic physician licensed pursuant to
chapter 36, a chiropractor licensed pursuant to chapter 9 and a podiatrist licensed pursuant to chapter 51
who treats Medicare-eligible individuals shall post in a conspicuous place that professional's policy
regarding the acceptance of Medicare assignment.

This posting must state the policy on accepting assignment and name the individual with whom the
patient should communicate regarding the policy.

The Board of Licensure in Medicine, the Board of Osteopathic Licensure, the Board of Licensure of
Podiatric Medicine and the Board of Chiropractic Licensure shall enforce the provisions of this section
and inform each licensee of the licensee's obligation under this law. Each board may discipline a licensee
under its jurisdiction for failing to comply with this section and impose a monetary penalty of not less
than $100 and not more than $1,000 for each violation.

(70) Mental and Behavioral Health (see also Agency Rules Links)
A complex set of statutes and regulation apply to mental health treatment and records. The State of
Maine Office of Substance Abuse and Mental Health Services provides useful guidance on the rights and
legal issues involved in mental health care, including guardianship, involuntary commitment and
confidentiality. See http://www.state.me.us/dhhs/samhs/mentalhealth/rights-legal/index.html Generally,
Title 34-B of the Maine Statutes governs mental health services.
**Confidentiality of information**

See also, Section on Medical Records – Confidentiality

**Generally**

The provisions dealing with confidentiality of mental health records apply to a person receiving services from the department, from any state institution or from any agency licensed or funded to provide services falling under the jurisdiction of the department. All orders of commitment, medical and administrative records, applications and reports, and facts contained in them, pertaining to any client shall be kept confidential and may not be disclosed by any person.

34-B MRSA §1001 (2)

34-B MRSA § 1207

For exceptions, such as when providers can disclose to family members, to avert harm, to coordinate care or to law enforcement, see: 34-B MRSA §1207.

**Violation**

Disclosure of client information in violation of this section is an offense under the licensing standards of the mental health professional committing the violation and must be promptly reported to the licensing board with jurisdiction for review, hearing and disciplinary action.

34-B MRSA §1207 (4-A)

**Client Rights**

For the rights applying to any resident of a state institutions or in a hospital or residential care facility, see 34-B MRSA §1430 and 34-B MRSA §3803.

**Hospitalization**

Chapter 3 of Title 34-B deals with hospitalization. See 34-B MRSA § 3801-3805 for the general provisions, 34-B MRSA §3831-3832 for the provisions applying to voluntary admissions and 34-B § 3861-3873-A for the provisions applying to involuntary admissions, such as emergency procedures, judicial procedures, discharge and progressive treatment programs.

**Intellectual Disabilities and Autism**

Chapter 5 of Title 34-B deals with services provided to those with intellectual disabilities and autism.

**Olmstead and the “Least Restrictive” Policy**

Olmstead v. L.C. was a landmark US Supreme Court case that that established to commonly known principle of the “least restrictive treatment environment.” The Court held that “states are required to provide community based treatment for persons with mental disabilities when the state’s treatment professions determine that such placement is appropriate, affected persons do not oppose such treatment and placement can be reasonably accommodated, taking into account the resources available to the state and the needs of others with mental disabilities.” Further, undue institutionalization of persons with mental disabilities qualifies as discrimination by reason of disability under the Americans with Disabilities Act.


**Maine policy concerning least restrictive treatment environments**

It is the policy of the State to provide education, training and habilitative services to mentally retarded persons who need those services, except that nothing in this chapter may replace or limit the right of any mentally retarded person to treatment by spiritual means alone, through prayer, if that treatment is requested by the person or by his next of kin or guardian.
It is the policy of the State that the setting for the services described in subsection 1 must, consistent with adequate care and treatment:

A. Impose the fewest possible restrictions on the liberty of persons with intellectual disabilities or autism; and

B. Be as close as possible to the patterns and norms of the mainstream of society.

See 34-B MRSA, Chapter 5 for more information on the provision of services to those with intellectual disabilities, including community based services, and the rights of those with intellectual disabilities. 34-B MRSA, Chapter 6 also applies to services for autism, including services for children.

**Sterilization**

34-B MRSA, Chapter 7 contains the Due Process in Sterilization Act of 1982, which containes the informed consent requirements for sterilization as well as the requirements for obtaining consent by the court for sterilization for someone unable to give informed consent.

**Declaration Directing Medical Treatment of Psychotic Disorders**

Any person 18 years of age or older who suffers from a psychotic condition but is competent and in a state of remission at the time of execution may execute a declaration directing that medical treatment, including the administration of psychotropic drugs, be provided at a time when the person has lapsed and is not able to make decisions regarding medical treatment. The provisions detailing the requirements of the declaration are found in 34-B MRSA, Chapter 11.

**Guardianship**

Maine Probate Code governs the guardianship process in Maine for both minors and incapacitated adults. See Title 18-A, Article 5 for the provisions related to guardianship.

**Midwifery**

**Midwifery Licensing Requirements and Scope of Practice**

**License required**

Beginning January 1, 2020, a person may not practice, offer to practice or profess to be authorized to practice midwifery, or hold oneself out to the public, as a midwife licensed in this State or use the words "certified professional midwife" or "certified midwife" or the letters "C.P.M." or "C.M." or other words or letters to indicate that the person using the words or letters is a licensed certified midwife or licensed certified professional midwife or that may misrepresent to the public that the person is authorized to practice midwifery in this State, unless that person is licensed in accordance with this subchapter.

See 32 MRSA §12531 (1)

**Scope of practice, medical testing**

The scope of practice of a certified professional midwife includes authorization to order and interpret medical laboratory tests and ultrasound scanning and to obtain equipment and supplies necessary for the safe practice of midwifery.

See 32 MRSA §12535 (3)

**Scope of practice, administration of drugs**

The scope of practice of a certified professional midwife includes the authority to obtain and administer certain drugs as determined by board rule. The board shall limit the drug formulary for certified professional midwives to only those medications that are indicated for the safe conduct of pregnancy, labor and birth and care of women and newborns and that a midwife is educationally prepared to...
administer and monitor. These may not include schedule II, III or IV drugs as defined in the federal Controlled Substances Act of 1970, 21 United States Code, Section 812.

32 MRSA §12535 (4)

Limitations on scope of practice
Certified professional midwives must refer clients to a hospital-based perinatal care provider and may not provide birth services to parents in a home or freestanding birth center setting when there is a reasonable likelihood that any of the following conditions exist:
A. Multifetal gestation;
B. Breech presentation;
C. Vaginal birth after a cesarean section; and
D. Conditions that present a moderate or high risk of harm to parent or child as defined in board rules.

Notwithstanding the above limitations, the board and the Board of Licensure in Medicine, jointly, prior to January 1, 2021 or the board beginning January 1, 2021 may adopt rules relating to the provision of birth services by certified professional midwives in cases in which there is a reasonable likelihood that any condition identified above exists.

32 MRSA §12536

For additional information regarding the qualifications for licensure as a certified midwife or a certified professional midwife, reporting requirements, and other information see MRSA 32 Ch. 113-B Subchapter 4.

Certified Midwives Access to Certain Medications

Drug administration by certified midwives under certain conditions
A midwife who can verify to a licensed pharmacist by certification card that the midwife has met the certification standards of an international certification agency whose mission is to establish and administer certification for the credential of certified professional midwife or other certifying body recognized by the board may:

Possess, in the course of the practice of midwifery, only the noncontrolled prescription drugs and substances set out in this subsection:
A. Oxygen;
B. Oxytocin, excluding the oxytocic drug methergine, for the sole purpose of postpartum control of maternal hemorrhage;
C. Vitamin K;
D. Eye prophylaxis; and
E. Local anesthetics or numbing agents for repair of lacerations; and

32 MRSA §13811 (1)

Administer, in the course of the practice of midwifery, those drugs that are listed in subsection 1. When administering oxytocin, a certified midwife may not administer more than 20 units of oxytocin to a single patient. Oxytocin may be administered only for postpartum purposes in order to treat hemorrhaging and specifically may not be used to induce labor. When a certified midwife administers oxytocin in accordance with this subsection, the certified midwife shall report that use to the maternal and child health division of the Department of Health and Human Services, the Maine Center for Disease Control and Prevention within 7 days of the use of oxytocin.

32 MRSA §13811 (2)

Dispensing of medication by pharmacist
A pharmacist, who in good faith relies upon a certification card presented by a midwife identifying that the midwife has met the certification standards described under section 13811, may sell and dispense to the midwife the noncontrolled prescription drugs and substances identified in section 13811.

32 MRSA §13812 (1)
Minors, Treatment and Consent of Minor’s Rights to Health Care Treatment

Usually, a physician must obtain consent to treatment of a minor from a parent or guardian, although the minor generally is involved in the process. Most of the time, the minor and both parents are involved in a family discussion about the treatment, but situations involving divorced parents can present challenges to the physician that are more practical than legal. In most cases of divorce, the parents have “shared parental rights and responsibilities,” meaning that both parents have the right to be involved in the major decisions of the child’s life, including decisions about health care treatment. 19-A MRSA §1501 (5). If a parent is claiming exclusive rights to make decisions about a minor’s medical treatment, the physician should as for documentation such as a divorce judgment awarding “sole parental rights and responsibilities” to one parent, an order terminating the parental rights of one parent, or other court order limiting a parent’s rights to participate in the medical decision-making process.

While the general rule is that physicians must obtain informed consent to treatment for minors from their parents, there are exceptions to the rule. Some minors may provide consent to all types of health care treatment and all minors may provide consent to some types of health care treatment. A minor may give consent to all types of health care treatment if the minor:

- Has been living separately from parents or legal guardians for at least 60 days and is independent of parental support;
- Is or was legally married;
- Is or was a member of the Armed Forces of the United States;
- Has been emancipated by the court pursuant to Title 15, section 3506-A. (22 MRSA §1503)

All minors may give consent to certain sensitive types of treatment where an obligation of parental consent may be an obstacle to treatment and, therefore, may not be in the best interest of the minor. These types of treatment include:

- Family planning services, including contraception, pregnancy testing, and emergency contraception (22 MRSA §1908);
- Treatment of venereal disease or drug alcohol abuse in the hospital setting, but parental consent is required if the hospitalization continues for more than 16 hours (22 MRSA §1823);
- Collection of sexual assault evidence through a sexual assault forensic examination (22 MRSA §1507);
- Treatment of venereal disease or drug or alcohol abuse by a physician (32 MRSA §§2595);
- Treatment of drug or alcohol abuse or for emotional or psychological problems (22 MRSA §1502);
- Consent to give blood by a 17-year-old (22 MRSA §1502-A);
- Certain services provided by alcohol and drug counselors, social workers, or psychologists (32 MRSA §§6221, 7004, 3817).

A minor may consent to an abortion if she accomplishes one of the following:

- Provides the physician performing the abortion with her informed written consent and the written consent of a parent or another adult family member such as an aunt or grandmother;
- Provides the physician performing the abortion with her informed written consent and receives abortion counseling. The counseling may be provided by a physician or from an approved counselor, who may be a psychiatrist, a psychologist, a social worker, an ordained clergy member, a physician’s assistant, a nurse practitioner, a guidance counselor, a registered nurse, or a licensed practical nurse; or
- Provides the physician performing the abortion with her informed written consent and a court order. (22 MRSA §1597-A).
The physician retains discretion to notify the parents if he or she believes that failure to do so would “seriously jeopardize the health of the minor or would seriously limit the practitioner’s or provider’s ability to provide treatment.” 22 MRSA §1505 (2). In general, a minor who consents to health care treatment is entitled to the same confidentiality rights as adults and is financially responsible to the physician for that treatment. 22 MRSA §§1505(1) and 1506. A physician who takes reasonable steps to determine that a minor is entitled to consent to health care treatment is immune from liability for a parent’s claim that the physician provided care without parental consent. 22 MRSA §1504.

**Definition**

**Minor:** means a person under 18 years of age. 22 MRSA §1501 (3)

**Consent, General**

In addition to the ability to consent to treatment for health services as provided in sections 1823 and 1908 and Title 32, sections 2595, 3292, 3817, 6221 and 7004, a minor may consent to treatment for abuse of alcohol or drugs or for emotional or psychological problems. 22 MRSA §1502

**Consent to give blood**

A minor may consent to give blood if the minor is at least 17 years of age, notwithstanding any other provision of law. 22 MRSA §1502-A

**Authority (to Consent)**

A minor may give consent to all medical, mental, dental and other health counseling and services if the minor:

A. **Living separately; independent of parental support**- Has been living separately from parents or legal guardians for at least 60 days and is independent of parental support;
B. **Married**- Is or was legally married;
C. **Armed Forces**- Is or was a member of the Armed Forces of the United States; or
D. **Emancipated**- Has been emancipated by the court pursuant to Title 15, section 3506-A.

22 MRSA §1503

**Good faith reliance on consent**

A health care practitioner or health care provider who takes reasonable steps to ascertain that a minor is authorized to consent to health treatment as authorized in section 1503 and who subsequently renders treatment in reliance on that consent is not liable for failing to have secured consent of the minor's parent or guardian prior to providing health care services to the minor. 22 MRSA §1504

**Confidentiality; notification**

A. **Confidentiality**- Except as otherwise provided by law, a minor who may consent to health care services, as provided in this chapter or by other provision of law, is entitled to the same confidentiality afforded to adults.
B. **Parental notification**- A health care practitioner or health care provider may notify the parent or guardian of a minor who has sought health care under this chapter if, in the judgment of the practitioner or provider, failure to inform the parent or guardian would seriously jeopardize the health of the minor or would seriously limit the practitioner's or provider's ability to provide treatment.

22 MRSA §1505

**Financial responsibility**

Unless the parent or guardian expressly agrees to assume full or partial responsibility, a minor who consents to health care services as provided in this chapter is responsible for the costs of those services. A minor may not be denied benefits or services to which the minor is entitled from a health care...
practitioner, health care provider, insurer or public agency because the minor has given the consent for
those services as provided in this chapter.

22 MRSA §1506

**Consent for sexual assault forensic examination**
Notwithstanding the limitations set forth in section 1503, a minor may consent to health services
associated with a sexual assault forensic examination to collect evidence after an alleged sexual assault.

22 MRSA §1507

**Consent to a minor's decision to have an abortion**
Except as otherwise provided by law, no person may knowingly perform an abortion upon a pregnant
minor unless:

A. The attending physician has received and will make part of the medical record the informed
written consent of the minor and one parent, guardian or adult family member;
B. The attending physician has secured the informed written consent of the minor as prescribed in
subsection 3 and the minor, under all the surrounding circumstances, is mentally and physically
competent to give consent;
C. The minor has received the information and counseling required under subsection 4, has secured
written verification of receiving the information and counseling and the attending physician has
received and will make part of the medical record the informed written consent of the minor and
the written verification of receiving information and counseling required under subsection 4; or
D. The Probate Court or District Court issues an order under subsection 6 on petition of the minor or
the next friend of the minor for purposes of filing a petition for the minor, granting:
   1. To the minor majority rights for the sole purpose of consenting to the abortion and the
      attending physician has received the informed written consent of the minor; or
   2. To the minor consent to the abortion, when the court has given its informed written
      consent and the minor is having the abortion willingly, in compliance with subsection 7.

22 MRSA §1597-A (2)

**Treatment of minors, Consent**

**Hospitals**
Any hospital licensed under this chapter or alcohol or drug treatment facility licensed pursuant to section
7801 that provides facilities to a minor in connection with the treatment of that minor for:

A. venereal disease;
B. abuse of drugs or alcohol; or
C. for the collection of sexual assault evidence through a sexual assault forensic examination
is under no obligation to obtain the consent of that minor's parent or guardian or to inform that parent or
guardian of the provision of such facilities so long as such facilities have been provided at the direction of
the person or persons referred to in Title 32, sections 2595 (repealed), 3292, 3817, 6221 or 7004 (social
workers).

**The 16-Hour Rule and Parental Consent**
The hospital shall notify and obtain the consent of that minor's parent or guardian if that hospitalization
continues for more than 16 hours.

22 MRSA §1823

**Medical Providers (Physicians)**
An individual licensed under this chapter who renders medical care to a minor for:

A. treatment of venereal disease;
B. abuse of drugs or alcohol; or
C. for the collection of sexual assault evidence through a sexual assault forensic examination
is under no obligation to obtain the consent of the minor's parent or guardian or to inform the parent or
guardian of the treatment. This section may not be construed to prohibit the licensed individual rendering
the treatment from informing the parent or guardian. For purposes of this section, "abuse of drugs" means the use of drugs solely for their stimulant, depressant or hallucinogenic effect upon the higher functions of the central nervous system and not as a therapeutic agent recommended by a practitioner in the course of medical treatment.

32 MRSA §3292

Psychological Services, Drug or Alcohol Abuse
Any person licensed under this chapter who renders psychological services to a minor for problems associated with the abuse of drugs or alcohol is under no obligation to obtain the consent of said minor's parent or guardian or to inform such parent or guardian of such services. Nothing in this section shall be construed so as to prohibit the licensed person rendering such services from informing such parent or guardian. For purposes of this section "abuse of drugs" means the use of drugs solely for their stimulant, depressant or hallucinogenic effect upon the higher functions of the central nervous system and not as a therapeutic agent recommended by a practitioner in the course of medical treatment.

32 MRSA §3817

Counseling Services, Drug or Alcohol Abuse
Any person licensed under this chapter who renders counseling services to a minor for the treatment of problems associated with the abuse of drugs or alcohol is under no obligation to obtain the consent of that minor's parent or guardian or to inform that parent or guardian of that treatment. Nothing in this section may be construed so as to prohibit the licensed person rendering that treatment from informing that parent or guardian. For purposes of this section "abuse of drugs" means the use of drugs solely for their stimulant, depressant or hallucinogenic effect upon the higher functions of the central nervous system and not as a therapeutic agent recommended by a practitioner in the course of medical treatment.

32 MRSA §6221

Medical treatment order
Who may Petition
The department (of Human Services), a physician or a chief medical administrator of a hospital may petition for a medical treatment order.

22 MRSA §4071 (1)

Contents of petition
A petition shall be sworn and shall include at least the following:

A. Name, date of birth and municipal residence, if known, of the child;
B. The name and address of the petitioner and his professional position;
C. Name and municipal residence, if known, of each parent and custodian;
D. A summary of the medical diagnosis and treatment alternatives;
E. A request for the court to order specific treatment; and
F. A statement that attempts to notify and secure consent from the custodians have been unsuccessful, either because they cannot be located or they have refused consent.

22 MRSA §4071 (2)

Notice to parents and custodians
The petitioner shall, by any reasonable means, attempt to notify the parents and custodians of his intent to request the order and of the time and place he will make the request, unless the petitioner believes that the child would suffer increased serious injury during the time needed to notify them.

22 MRSA §4071 (3)

Order
On the basis of the petition or other evidence, the court may order medical treatment for the child if the custodians are unable or unwilling to consent to it, and the treatment is necessary to treat or prevent an immediate risk of serious injury. The order shall include a notice to the parents and custodians of their right to counsel, as required under section 4032, subsection 2, paragraph G, and notice of the date and time of the hearing.

22 MRSA §4071 (4)
Service of order
If a hearing has not been held prior to issuing the order, a copy of the order and petition shall be served on the parents and custodians by:

A. In-hand delivery by the judge or court clerk to any parent, custodian or their counsel who is present when the order is issued;
B. Service in accordance with the District Court Civil Rules. Notwithstanding the civil rules, service by publication of an order and petition shall be complete 5 days after a single publication; or
C. Another manner ordered by the court.

Hearing
If a hearing has not been held prior to issuing the order, then it shall be held within 10 days of its issuance, unless all parties agree to a later date. If, after the hearing, the court finds, by a preponderance of the evidence, that the medical treatment ordered is necessary to treat or prevent the immediate risk of serious injury to the child, then it may continue the order.

Authorization for Certain Campers To Self-administer Emergency Medication
A recreational camp for boys or girls must have a written policy authorizing campers to self-administer emergency medication, including, but not limited to, an asthma inhaler or an epinephrine pen.

Nursing (see also Agency Rules Links)
For Maine statues governing the practice of nursing, see Title 32, Chapter 31 of the Maine Revised Statutes. This Chapter also applies to advanced practice registered nurses, including certified nurse practitioners, certified nurse midwives, certified clinical nurse specialists and certified nurse anesthetists. For information regarding certified nursing assistants and unlicensed assistive persons in facilities, see also Title 22, Sections 1812-G and 1812-J.

Nurse health program
The board may establish protocols for the operation of a professional review committee as defined in Title 24, section 2502, subsection 4-A. The protocols must include the committee's reporting information the board considers appropriate regarding reports received, contracts or investigations made and the disposition of each report, as long as the committee is not required to disclose any personally identifiable information. The protocols may not prohibit an impaired nurse from seeking alternative forms of treatment.

The board may contract with other agencies, individuals, firms or associations for the conduct and operation of a nurse health program operated by a professional review committee as that term is defined in Title 24, section 2502, subsection 4-A. (At present those functions are fulfilled by the Maine Medical Association’s Medical Professionals’ Health Program.)

Occupational Health (see also Workers’ Compensation)
Duties of physicians and hospitals
All physicians or hospitals shall report to the Department of Human Services all persons diagnosed as having an occupational disease no later than 30 days from the date of diagnosis or from discharge from a hospital. The report shall include any factor known to the physician which is suspected of being a contributing factor to the disease, including, but not limited to, whether or not the person smokes and, if so, the frequency of smoking.
A physician, upon notification by the Department of Human Services, shall report to the department any further information requested by the department concerning any person now or formerly under his care, diagnosed as having or having had an occupational disease.

No physician or hospital complying with the reporting requirements of this section may be liable for any civil damages as a result of those acts.

Confidentiality
The names and related information which may identify individuals having an occupational disease shall be confidential and may be released only to other public health officials, agents or agencies, or by court order or by written authorization of the individual being reported on. All other information submitted pursuant to this chapter may be made available to the public.

Medical Allowance for Licensed Activity
A person may fish for lobster and crab under another person’s license if the licensee is unable to use that license due to a substantial illness or medical condition documented by a physician. Such temporary medical allowance is for one year and is renewable for one additional year.

(75) Opioid Medication Prescribing

Definitions
A. Controlled substance means a controlled substance included in schedules II, III or IV of 21 United States Code, Section 812 or 21 Code of Federal Regulations, Section 1308.
B. Acute pain means pain that is the normal, predicted physiological response to a noxious chemical or thermal or mechanical stimulus. "Acute pain" typically is associated with invasive procedures, trauma and disease and is usually time-limited.
C. Administer means an action to apply a prescription drug directly to a person by any means by a licensed or certified health care professional acting within that professional's scope of practice. "Administer" does not include the delivery, dispensing or distribution of a prescription drug for later use.
D. Chronic pain means pain that persists beyond the usual course of an acute disease or healing of an injury. "Chronic pain" may or may not be associated with an acute or chronic pathologic process that causes continuous or intermittent pain over months or years.
E. Dispenser means a pharmacist who is licensed or registered under Title 32 or a licensed health care professional with authority to dispense or administer prescription drugs.
F. Prescriber means a licensed health care professional with authority to prescribe controlled substances and a veterinarian licensed under Title 32, chapter 71-A with authority to prescribe controlled substances.

Dispensing by Nurses
Certified nurse practitioners, registered nurses, and licensed practical nurses may dispense opioid medication for substance abuse treatment purposes to patients within an opioid treatment program under the direction of the medical director of the program.
Required Check of Prescription Monitoring Information

Prescribers
Upon initial prescription of a benzodiazepine or an opioid medication to a person and every 90 days for as long as that prescription is renewed, a prescriber shall check prescription monitoring information for records related to that person.

Dispensers
A dispenser shall check prescription monitoring information prior to dispensing a benzodiazepine or an opioid medication to a person under any of the following circumstances:

A. The person is not a resident of this State;
B. The prescription is from a prescriber with an address outside of this State;
C. The person is paying cash when the person has prescription insurance on file; or
D. According to the pharmacy prescription record, the person has not had a prescription for a benzodiazepine or an opioid medication in the previous 12-month period.

A dispenser shall withhold a prescription until the dispenser is able to contact the prescriber of that prescription if the dispenser has reason to believe that the prescription is fraudulent or duplicative.

Exception; hospital setting and facilities
When a licensed or certified health care professional directly orders or administers a benzodiazepine or opioid medication to a person in an emergency room setting, an inpatient hospital setting, a long-term care facility or a residential care facility, or in connection with a surgical procedure, the requirements to check prescription monitoring information established in this section do not apply.

Violation
A person who violates this section commits a civil violation for which a fine of $250 per incident, not to exceed $5,000 per calendar year, may be adjudged.

Requirements regarding prescription of opioid medication
Limits on opioid medication prescribing
Except as provided in the “exceptions” section below, an individual licensed under this chapter and whose scope of practice includes prescribing opioid medication may not prescribe:

A. To a patient any combination of opioid medication in an aggregate amount in excess of 100 morphine milligram equivalents of opioid medication per day;
B. To a patient who, on the effective date of this section, has an active prescription for opioid medication in excess of 100 morphine milligram equivalents of an opioid medication per day, an opioid medication in an amount that would cause that patient's total amount of opioid medication to exceed 300 morphine milligram equivalents of opioid medication per day; except that, on or after July 1, 2017, the aggregate amount of opioid medication prescribed may not be in excess of 100 morphine milligram equivalents of opioid medication per day;
C. On or after January 1, 2017, within a 30-day period, more than a 30-day supply of an opioid medication to a patient under treatment for chronic pain. "Chronic pain" has the same meaning as in Title 22, section 7246, subsection 1-C; or
D. On or after January 1, 2017, within a 7-day period, more than a 7-day supply of an opioid medication to a patient under treatment for acute pain.

Exceptions
An individual licensed under this chapter whose scope of practice includes prescribing opioid medication is exempt from the limits on opioid medication prescribing established in subsection 1 only:

A. When prescribing opioid medication to a patient for:
1. Pain associated with active and aftercare cancer treatment;
2. Palliative care in conjunction with a serious illness
3. End-of-life and hospice care;
4. Medication-assisted treatment for substance use disorder; or
5. Other circumstances determined in rule by the Department of Health and Human Services. Those rules establish the following additional exceptions:
   i. A pregnant person;
   ii. A prescription for acute pain for a patient taking opioids for chronic pain;
   iii. A patient being actively tapered from high doses of opioids;
   iv. A patient who is intolerant to previously prescribed opioids.

B. When directly ordering or administering a benzodiazepine or opioid medication to a person in an emergency room setting, an inpatient hospital setting, a long-term care facility or a residential care facility, or in connection with a surgical procedure.

Electronic prescribing
An individual licensed under this chapter and whose scope of practice includes prescribing opioid medication with the capability to electronically prescribe shall prescribe all opioid medication electronically. An individual who does not have the capability to electronically prescribe must request a waiver from this requirement from the Commissioner of Health and Human Services stating the reasons for the lack of capability, the availability of broadband infrastructure, and a plan for developing the ability to electronically prescribe opioid medication. The commissioner may grant a waiver including circumstances in which exceptions are appropriate, including prescribing outside of the individual's usual place of business and technological failures.

Continuing education
An individual licensed under this chapter must successfully complete 3 hours of continuing education every 2 years on the prescription of opioid medication as a condition of prescribing opioid medication. The board shall adopt rules to implement this subsection. (Note: The Board of Licensure in Medicine requires this CME of all its licensees, regardless of whether they prescribe opioids.)

Penalties
An individual who violates this section commits a civil violation for which a fine of $250 per violation, not to exceed $5,000 per calendar year, may be adjudged. The Department of Health and Human Services is responsible for the enforcement of this section.

Partial dispensing of prescription for opioid medication
Partial dispensing authorized
Notwithstanding any law or rule to the contrary, a pharmacist may partially dispense a prescription for an opioid medication in a lesser quantity than the recommended full quantity indicated on the prescription if requested by the patient for whom the prescription is written. The remaining quantity of the prescription in excess of the recommended full quantity is void and may not be dispensed without a new prescription.

Notice to practitioner
If a pharmacist partially dispenses a prescription for an opioid medication as permitted under this section, the pharmacist or the pharmacist's designee shall, within a reasonable time following the partial dispensing but not more than 7 days, notify the practitioner of the quantity of the opioid medication actually dispensed. The notice may be conveyed by a notation on the patient's electronic health record or by electronic transmission, by facsimile or by telephone to the practitioner.
Optometry (see also Agency Rules Links)
Scope of Practice
An optometrist may not administer therapeutic drugs by injection, other than for emergency treatment of anaphylaxis. They may, however, dispense or sell contact lenses that contain and deliver pharmaceutical agents that are authorized under the Maine Revised Statutes, Title 32, chapter 34-A.

An optometrist who graduated from optometric college in the year 1996 or thereafter and who is an advanced therapeutic licensee is authorized to independently treat glaucoma. In order to be authorized to independently treat glaucoma, an advanced therapeutic licensee who graduated from optometric college prior to 1996 must provide evidence to the board of no more than 30 glaucoma-related consultations with a physician.

Licensure Requirements
The following are the requirements applicants must meet before licensure:
A. Be at least 18 years of age;
B. Be a graduate of a recognized school of optometry; and
C. Have succeeded in an examination as described in section 2422.

For more on Optometrists’ Scope of Practice see Title 32, chapter 34-A and Agency Rules links.

Peer Review
Review committee member immunity
A physician licensed under this chapter who is a member of a utilization review committee, medical review committee, surgical review committee, peer review committee or disciplinary committee that is a requirement of accreditation by the Joint Commission on Accreditation of Hospitals or is established and operated under the auspices of the physician's respective state or county professional society or the Board of Licensure in Medicine is immune from civil liability for undertaking or failing to undertake an act within the scope of the function of the committee.

Records of proceedings of medical staff review committees confidential
All proceedings and records of proceedings concerning medical staff reviews, hospital reviews and other reviews of medical care conducted by committees of physicians and other health care personnel on behalf of hospitals located within the State or on behalf of individual physicians, when the reviews are required by state or federal law, rule or as a condition of accreditation by the Joint Commission on Accreditation of Hospitals or the American Osteopathic Association Committee on Hospital Accreditation or are conducted under the auspices of the state or county professional society to which the physician belongs, are confidential and are exempt from discovery.

Pharmacy Practice
Scope of Practice
Vaccine administration
Pharmacists’ scope of practice is expanded to allow them to administer all vaccines licensed by the United States Food and Drug Administration that are recommended by the United States Centers for
Disease Control and Prevention Advisory Committee on Immunization Practices, or successor organization, for administration to adults to a person 18 years of age or older according to a valid prescription when the person has an existing primary care physician or other existing relationship with a nurse practitioner or an authorized practitioner in this State. A pharmacist may administer vaccines licensed by the FDA that are outside the guidelines recommended by the Centers for Disease Control and Prevention Advisory Committee on Immunization Practices when it is deemed medically necessary by the practitioner. When the person does not have an existing relationship with a primary care physician, nurse practitioner or other practitioner in this State, the pharmacist may proceed to administer according to a treatment protocol established by an authorized practitioner or a written standing order. Licensed pharmacists can administer all forms of influenza vaccines to a person 7 years of age or older without a prescription.

Collaborative Drug Therapy

A pharmacist licensed in this State who meets the qualifications and requirements of section 13842 and rules adopted by the board may engage in collaborative drug therapy management pursuant to a collaborative practice agreement with a practitioner.

A pharmacist engaging in collaborative drug therapy management pursuant to subsection 1 is entitled to adequate access to a patient's history, disease status, drug therapy and laboratory and procedure results and may:

A. Collect and review a patient's history;
B. Obtain and check vital signs;
C. Order and evaluate the results of laboratory tests directly related to drug therapy under the supervision of, or in direct consultation with, a practitioner and in accordance with approved protocols applicable to the practice setting and when the evaluation does not include a diagnostic component; and
D. Initiate, monitor, modify and discontinue drug therapy for a particular patient pursuant to the collaborative practice agreement with a practitioner who is treating the patient, as long as the action is reported to the practitioner in a timely manner as determined by rules adopted pursuant to section 13846.

Naloxone Prescribing

Pharmacists may prescribe and dispense naloxone hydrochloride to any person, regardless of age, in accordance with rules established by the Board of Pharmacy.

Nicotine Replacement Products

“Practice of pharmacy” includes the ordering and dispensing of over-the-counter nicotine replacement products approved by the FDA.

For more on Pharmacists’ Scope of Practice see the Maine Pharmacy Act, 32 MRSA Chapter 117.

Pharmacist Health Program

The Maine Board of Pharmacy may establish protocols for the operation of a professional review committee as defined in Title 24, section 2502, subsection 4-A. The protocols must include the committee’s reporting information the board considers appropriate regarding reports received, contracts or investigations made and the disposition of each report, as long as the committee is not required to disclose any personally identifiable information. The protocols may not prohibit an impaired pharmacist or pharmacy technician from seeking alternative forms of treatment.
The board has the power to contract with other agencies, individuals, firms or associations for the conduct and operation of a pharmacist health program operated by a professional review committee as that terms is defined in Title 24, section 2502, subsection 4-A.

**Professional review committee defined**

“Professional review committee” means a committee of physicians, dentists, pharmacists, nurses or a combination of members of all 3 professions formed by a professional society for the purpose of identifying and working with physicians, dentists and other licensees of the Board of Dental Examiners, physician assistants, pharmacists and pharmacy technicians and nurses who are disabled or impaired by virtue of physical or mental infirmity or by the misuse of alcohol or drugs, as long as the committee operates pursuant to protocols approved by the Board of Licensure in Medicine, the Board of Dental Examiners, the Board of Osteopathic Licensure, the Maine Board of Pharmacy, and the State Board of Nursing.

**Patient information regulation**

**Explanation by pharmacist**

With each new prescription dispensed, the pharmacist, in addition to labeling the prescription in accordance with the requirements of the State, must orally explain to the patient or the patient's agent the directions for use and any additional information, in writing if necessary, to assure the proper utilization of the medication or device prescribed. For those prescriptions delivered outside the confines of the pharmacy, the explanation shall be by telephone or in writing. This section does not apply to those prescriptions for patients in hospitals or institutions where the medication is to be administered by a nurse or other individual licensed to administer medications or to those prescriptions for patients who are to be discharged from a hospital or institution.

**Maintenance of current reference material**

To ensure that proper information is available to each pharmacist, each pharmacy or pharmacist shall maintain current reference material on drug interactions.

**Retail price**

With each prescription dispensed, the pharmacist shall disclose to the patient in writing the usual and customary price of the prescription and the cost of any payment toward the price required of the patient.

**Physician Assistants**

Physician Assistants in Maine historically were required to receive a license from either the Board of Licensure in Medicine or the Board of Osteopathic Licensure, depending on the licensure of the supervising physician. As of 2014, the Boards are creating one set of consistent rules for licensure. Physician Assistants are now also advisory members of both the medical and osteopathic boards.

The relevant laws and rules can be found here:

Board of Licensure in Medicine laws: 32 MRSA § 3270-A, §3270-C, §3270-E

Board of Licensure in Medicine rules

Board of Osteopathic Licensure laws: 32 MRSA § 2594-A, § 2594-D, §2594-E

A Physician Assistant may delegate medical acts to a medical assistant employed by the physician assistant or by an employer of the physician assistant. However, such delegation must be permitted in the plan of supervision established by the physician assistant and the supervisory physician.

32 MRSA §2594-A
The sections of law authorizing allopathic and osteopathic physicians to delegate to physician assistants are found at 32 MRSA §2594-A and 32 MRSA §3270-A.

**Podiatric assistants**

For the law governing podiatric assistants, see 32 MRSA §3552-A.

**(80) Physician Licensing (see also Agency Rules Links)**

**Allopathic Physicians (MDs)**

The Board of Licensure in Medicine is the entity in Maine responsible for licensing allopathic physicians (MDs) and physician assistants supervised by allopathic physicians.

The statutes governing the dealings of the Board and the licensure/practice of medicine in Maine are found at 32 MRSA, Chapter 48.

32 MRSA § 3263-3269 apply to the functions of the Board of Medicine.

32 MRSA § 3270-3289 deal with the process and requirements for licensure and discipline of physicians. It is also important to review the Board of Licensure in Medicine Rules, including Chapter 1 dealing with licensure and Board procedures, Chapter 10 applying to sexual misconduct and Chapter 21 covering the use of controlled substances for pain.

32 MRSA § 3280 deals with license renewal and board notification for MD physicians. The requirements were changed in 2017 to provide for immediate and automatic expiration of a medical license that is not renewed by the expiration date, with provisions for reinstatement within 90 days. Until it is reinstated, the physician is prohibited from practicing medicine.

For more information, visit the [website of the Board of Licensure in Medicine](#).

**Osteopathic Physicians (DOs)**

The Board of Osteopathic Licensure is the entity in Maine responsible for licensing osteopathic physicians (DOs) and physician assistants supervised by osteopathic physicians.

The statutes governing the dealings of the Board and the licensure/practice of medicine by osteopathic physicians in Maine are found at 32 MRSA, Chapter 36.

32 MRSA § 2561-2563-A apply to the functions of the Board of Osteopathic Licensure.

32 MRSA § 2571-2592-A deal with the process and requirements for licensure and discipline of physicians.

It is also important to review the Board of Osteopathic Licensure Rules, including Chapter 21 covering the use of controlled substances for pain.

For more information, visit the [website of the Board of Osteopathic Licensure](#).

**Physicians licensed in other states**

Physicians holding a current, unrestricted license in another state who have a written agreement with an athletic team located in the state of licensure may provide medical services to the following persons while the team is traveling to or from or participating in a sporting event in this State:

1. A member of the athletic team;
2. A member of the team’s coaching, communications, equipment, or sports medicine staff;
3. A member of a band or cheerleading squad accompanying the team; or
4. The team’s mascot.
This does not include services at a health care facility, including a hospital, ambulatory surgical facility or any other facility where medical care, diagnosis or treatment is provided on an inpatient or outpatient basis.

**Maintenance of Certification**
The licensing boards for physicians (MD and DO) may not require an applicant for initial licensure or license renewal to obtain specialty board certification, osteopathic continuous certification, or maintenance of such certification as a condition of licensure.

**Interstate Medical Licensure Compact**
A physician licensed to practice in a Compact member state may seek licensure in other member states through a streamlined process. The statute sets out rules for eligibility, application, renewal, enforcement and oversight.

**(81) Physician, Mental Health Professional, and Licensed Counseling Professional-Patient Privilege**

**Definitions**

A. **Patient:** A person who consults or is examined or interviewed by a health care professional, a mental health professional, or a licensed counseling professional.

B. **Health care professional:**
   1. A person authorized to practice as a physician;
   2. A licensed physician’s assistant; or
   3. A licensed nurse practitioner;
   Under Maine law or under substantially similar law of any state or nation, while that person is practicing the health care profession for which he or she is licensed.

C. **Mental health professional:**
   1. A health care professional engaged in the diagnosis or treatment of a mental or emotional condition, including alcohol or drug addiction;
   A person licensed or certified as a psychologist or psychological examiner under Maine state law or under substantially similar law of any state or nation while practicing as such;
   A person licensed as a clinical social worker under Maine state law or under substantially similar law of any state or nation while practicing as such.

D. **Licensed counseling professional:**
   1. A “licensed professional counselor”;
   2. A “licensed clinical professional counselor”;
   3. A “licensed marriage and family therapist” or;
   4. A “licensed pastoral counselor”;
   Who is licensed to diagnose and treat mental health disorders, intra and inter-personal problems, or other dysfunctional behavior of a social and spiritual nature under 32 M.R.S. §13858, or under a substantially similar law of any other state or nation, while that person is practicing the counseling profession for which he or she is licensed.

**Confidential Communication**
A communication is "confidential" if not intended to be disclosed to third persons other than those present to further the interest of the patient in the consultation, examination, or interview, or persons reasonably necessary for the transmission of the communication, or persons who are participating in the diagnosis
and treatment under the direction of the health care, mental health, or licensed counseling professional, including members of the patient's family.

Maine Rules of Evidence, Rule 503 (a) (4)

General rule of privilege
A patient has a privilege to refuse to disclose and to prevent any other person from disclosing confidential communications made for the purpose of diagnosis or treatment of the patient's physical, mental or emotional condition, including alcohol or drug addiction, between or among the patient, the patient's health care professional, mental health professional, or licensed counseling professional, and persons who are participating in the diagnosis or treatment under the direction of the health care professional, mental health professional, or licensed counseling professional, including members of the patient's family.

Maine Rules of Evidence, Rule 503 (b)

Privilege of accused
When an examination of the mental condition of an accused in a criminal proceeding is ordered by the court for the purpose of determining criminal responsibility, the accused has a privilege to refuse to disclose and to prevent any other person from disclosing any communication concerning the offense charged, made in the course of the examination.

Maine Rules of Evidence, Rule 503 (c)

Who may claim the privilege
The privilege may be claimed by the patient, by the patient's guardian or conservator, or by the personal representative of a deceased patient. The person who was the health care, mental health, or licensed counseling professional at the time of the communication is presumed to have authority to claim the privilege but only on behalf of the patient.

Maine Rules of Evidence, Rule 503 (d)

Exceptions
Proceedings for hospitalization
There is no privilege under this rule for communications relevant to an issue in proceedings to hospitalize the patient for mental illness, if the professional in the course of diagnosis or treatment has determined that the patient is in need of hospitalization.

Examination by order of court
If the court orders an evaluation of a patient’s physical, mental, or emotional condition, whether the patient is a party or a witness, the privilege does not apply to communications made during the course of that evaluation, unless the court orders otherwise. However, a criminal defendant’s communications during the course of a court-ordered evaluation or examination are still privileged to the extent provided by section (c) of this rule.

Condition an element of claim or defense
The privilege under this rule does not apply to communications relevant to an issue of a physical, mental, or emotional condition of the patient if:

A. The condition is an element of the patient’s claim or defense; or
B. The condition is an element of the claim or defense of:
   1. Any party claiming through or under the patient;
   2. Any party claiming because of the patient’s condition;
   3. Any party claiming as a beneficiary of the patient; or

Any party claiming through a contract to which the patient is or was a party.

After the patient’s death
The privilege does not apply after the patient’s death in any proceeding in which any party puts the patient’s physical, mental, or emotional condition in issue.  

**Maine Rules of Evidence, Rule 503 (e)**

**Exception for information communicated to acquire drugs by deception.**
Information communicated to a prescribing health care provider, or a person acting under the direction or supervision of a prescribing health care provider, in an effort to violate this section, including a violation by procuring the administration of a scheduled drug by deception, may not be deemed a privileged communication. 

**17-A MRSA §1108 (3)**

(82) **Power of Attorney for Healthcare (see also Advanced Healthcare Directives)**

Under Maine law, the term “advance directive” means any spoken or written instructions someone gives about the health care someone wants if a time comes when they are too ill to decide. A health care power of attorney is an example of an advance directive that must be in writing. A written power of attorney allows patients to appoint agents to make health care decisions, choose treatments they want or do not want, name a primary care provider, state wishes about donating their body or body parts and state wishes about funeral and burial arrangements.

Anyone 18 or older may execute a Maine Health Care Advance Directive in whole or in part. Those younger than 18 may also be able to use an advance directive under certain limited circumstances. For Maine’s sample Advanced Healthcare Directive form, see http://www.maine.gov/dhhs/oads/aps-guardianship/documents/advdirectivesform.pdf

For frequently asked questions about Advanced Healthcare Directives in Maine, see: http://www.themha.org/policy-advocacy/Issues/End-of-Life-Care/Facts-About-Health-Care-Advance-Directives.aspx

For Maine’s law on Advanced Healthcare Directives, including obligations of health care providers, see the Uniform Health-Care Decisions Act, Title 18-A, Part 8 (§5-801 and following).

**Obligations of the health care provider**
Health care providers must comply with an individual instruction of the patient and with a reasonable interpretation of that instruction made by a person then authorized to make health-care decisions for the patient.

**18-A MRSA §5-807(d)**

**Decisions by surrogate**
In absence of an agent named in an Advanced Healthcare Directive (or court-appointed guardian), physicians may turn to other surrogates to make health care decisions for patients who lack capacity, in this order:

1. Spouse (unless legally separated);
2. Someone with whom the patient share an emotional, physical and financial bond similar to a spouse;
3. Adult children;
4. Parents;
5. Adult brothers and sisters;
6. Adult grandchildren;
7. Adult nieces and nephews;
8. Adult aunts and uncles; or
8. Another adult relative of the patient who is familiar with the patient's personal values and is reasonably available for consultation.

**18-A MRSA §5-805.**

Maine law also allows for financial power of attorneys, see **18-A MRSA Part 9.**

### (83) Professional Liability

#### Medical Malpractice

Medical malpractice is an aspect of the common law “tort” of “negligence.” Black’s Law Dictionary (5th Edition) defines “tort” as “a private or civil wrong or injury, other than a breach of contract, for which the court will provide a remedy in the form of an action for damages.” The term “damages” means an award of money. Each action for negligence requires the existence of a legal duty from the defendant to the plaintiff, breach of that duty, and harm as a proximate result of that breach.

The following are types of damages that may be awarded in actions for professional negligence:

A. **Compensatory damages:** means damages to compensate an injured party. There are two types of compensatory damages:
   1. Economic damages, which consist of:
      a. Current and future medical expenses, including the cost of medical care, medicines and medical supplies; and
      b. Employment-related damages, including lost wages, lost or diminished earning capacity and lost earning opportunity; and
   2. Noneconomic damages, which are damages for physical and emotional pain, suffering, inconvenience, physical impairment, mental anguish, disfigurement, loss of enjoyment of life, loss of society and companionship, loss of consortium, hedonic damages, injury to reputation and all other nonpecuniary losses of any kind or nature.

B. **Punitive damages:** means damages intended to punish willful, malicious or fraudulent behavior and to discourage similar behavior by others.

#### Notice of claim before suit

No action for professional negligence may be commenced until the plaintiff has:

A. Served and filed written notice of claim in accordance with section 2853;
B. Complied with the provisions of subchapter IV-A; and
C. Determined that the time periods provided in section 2859 have expired.

**24 MRSA §2903 (1)**

#### Tort Reforms

Maine has seven provisions of law that may be referred to as Maine’s “tort reforms.” These provisions have contributed to a medical malpractice insurance market that offers some of the lowest liability premium rates in the country and the second lowest to Vermont in New England according to a 2005 study on the subject by the Maine Bureau of Insurance. These provisions include:

A. Mandatory pre-litigation screening and mediation panels;
B. A favorable statute of limitations;
C. A provision addressing evidence of payments to a plaintiff from a collateral source;
D. A provision regarding the apportioning of fault among a claimant and one or more defendants;
E. Limits on attorney contingent fees;
F. A provision providing for periodic payment of damage awards; and
G. A provision protecting statements of benevolence or sympathy (“I’m sorry” law).

#### Mandatory Pre-Litigation Screening & Mediation Panels

Maine’s *Mandatory prelitigation screening and mediation panels, 24 M.R.S.A. §§2851-2859,* were adopted in the mid-1980s to identify and encourage the early resolution of meritorious malpractice claims
and to encourage the early withdrawal or dismissal of claims that lack merit. The 3-member panel is composed of a chairperson who is a retired judge or someone with “judicial experience,” an attorney, and a health care practitioner (preferably the same type of practitioner in the same specialty as the practitioner alleged to have committed malpractice). The panel conducts a hearing, generally lasting no more than a day, within an administrative structure that is less formal than a judicial hearing. The panel applies the same standard of proof that a court would apply in a civil matter. The plaintiff must prove negligence and proximate causation by a preponderance of the evidence (more likely than not) and the defendant must prove comparative negligence by the same standard. The panel makes its findings in writing within 30 days of the close of the hearing. The findings must address the following questions about negligence and causation:

A. Whether the acts or omissions complained of constitute a deviation from the applicable standard of care by the health care practitioner or provider charged with that care;
B. Whether the acts or omissions complained of proximately caused the injury complained of; and
C. If negligence on the part of the health care practitioner or provider is found, whether any negligence on the part of the patient was equal to or greater than the negligence on the part of the practitioner or provider.

The proceedings before the panel, including deliberations, testimony, and findings, are confidential, but the panel findings are admissible in later court action according to guidelines set forth in the statute and in a series of decisions by the Maine Supreme Judicial Court. These Law Court decisions have provided some procedural and evidentiary guidelines for the panel process, but they have upheld the constitutionality of the panel’s role in resolving medical malpractice cases in Maine.

While the statute is entitled “screening and mediation panels,” in practice the panels conduct a screening hearing, and mediation does not take place in that context.

The Maine Bureau of Insurance has commissioned the only independent third party review of the panel process and that review is favorable. In its 1997 report entitled, Analysis of the Effectiveness of the Medical Professional Liability Prescreening Panels, AMI Risk Consultants, Inc. found that the panels have been successful in promoting quicker recovery for those who receive awards and promoting earlier dismissal of claims that conclude with no award while not reducing the overall average size of awards. While several other states have a screening panel system in their medical liability laws, none seem to have a system that is as effective as Maine’s. Maine’s system is the national standard against which other states’ systems are judged. The majority of medical malpractice claims in Maine are resolved at or before the panel stage in the case. Each year, only a few medical malpractice cases in Maine are tried to a jury. Accordingly, Maine’s screening panels probably are the most important “reform” of our medical liability laws.

Statute of Limitations

The Statute of limitations for health care providers and health care practitioners, 24 M.R.S.A. §2902, provides that actions for medical negligence must be brought within 3 years from occurrence for adults. For minors, the action must be brought within 6 years from occurrence or 3 years from the age of majority whichever is shorter. The limitation of actions involving a foreign object is measured from discovery of the object. Compared to other states, the Maine statute of limitations for medical malpractice is quite favorable to the health care practitioner because the general civil statute of limitations is 6 years from occurrence and because of the absence of a “discovery rule.” Most other states’ statute of limitations for medical malpractice permit the period of litigation exposure to be expanded by measuring the period from the time the negligent act or omission is or should have been discovered, rather than from the occurrence of the negligent act or omission.

24 MRSA §§2851-2859

24 MRSA §2902
Certain Negligence Suits
For professional negligence actions against certain health care providers the statute of limitations on professional negligence actions is six years when the action is based on a sexual act or sexual contact. This applies to psychiatrists, psychologists, social workers, professional counselors, pastoral counselors, marriage and family therapist and clinical professional counselors.

Evidence of Payments to a Plaintiff from a Collateral Source
In 24 M.R.S.A. §2906, Collateral sources, the Maine legislature has permitted evidence of expenses for medical care, rehabilitation services, loss of earnings, loss of earning capacity, or other economic loss paid or payable by a collateral source to be admissible after the verdict but before judgment. The statute requires the court to reduce the award accordingly, thereby mitigating the impact on the defendant(s) and the liability insurance carrier(s).

Comparative Negligence: Apportioning Fault
This section describes how Maine law addresses situations in which multiple defendants and the claimant may have been negligent. In 14 M.R.S.A. §156, Comparative negligence, the Maine legislature has blended the concepts of joint and several liability, modified comparative negligence, and contributory negligence. It provides that a plaintiff’s recovery may be reduced to the extent the jury determines is “just and equitable” having regard to the claimant’s share of responsibility for the damage. Recovery is barred if the claimant is found to be equally or more at fault. Finally, the statute states that each defendant is jointly and severally liable to the claimant for the full amount of the damages, but any defendant has the right to request that the jury determine the percentage of fault contributed by each defendant.

Limits on Attorney Contingent Fees
In 24 M.R.S.A. §2961, Contingent Fees, the Maine legislature has limited the contingency fees that plaintiffs’ attorneys may charge in order to mitigate the incentive to inflate fee recovery by pushing for higher damages. The limits, excluding litigation expenses, are:
   A. 33 1/3% of the first $100,000 in damages;
   B. 25% of the next $100,000 in damages; and
   C. 20% of any amount over $200,000.
A plaintiff’s attorney may, however, petition the court for a higher award and based upon anecdotal reports, it is common practice for attorneys to make such requests of the court. According to AMA data, approximately 16 states have limits on attorney contingent fees.

Periodic Payment of Damage Awards
Maine’s tort laws recognize that requiring a health care practitioner to pay a large damage award in a lump sum could pose a significant financial hardship that could threaten the viability of his or her practice. Maine law permits a court to order periodic payment rather than lump-sum payment if future damages equal or exceed $250,000. See 24 M.R.S.A. §2951, Provision for structured awards.

Communications of Sympathy or Benevolence (“I’m Sorry” Provision)
Maine’s tort laws recognize that patients who have experienced an unanticipated outcome during an encounter with the health care system are less likely to take legal action if the health care practitioner(s) show empathy in the circumstances and communicate directly with them about the unanticipated outcome. The so-called “I’m sorry” provision prevents any statement or gesture of apology, sympathy, or condolence to a patient, or to his or her relative or representative, from being used in an action for professional negligence. See 24 M.R.S.A. §2907, Communications of sympathy or benevolence.
Liability Claims Reports
Liability insurance carriers are required to report claims involving a healthcare provider to the appropriate board or state licensing authority within 30 days of their receipt. Such claims must include a copy or summary of reports received. Further information may be found at 24 MRSA §2601 or §2602.

Cancellation or nonrenewal
Any insurer required to report claims information shall also notify the Superintendent of Insurance of the cancellation or nonrenewal of any insured occasioned by either the number of claims against that insured or by the insured's failure to conform to appropriate standards of the medical profession. The information is entitled to the confidentiality protection of 24 MRSA §2604. The superintendent must file a copy of the report within 30 days of its receipt, with the applicable licensing board or authority.

(84) Public Health (see also Agency Rules Links)
Maine’s Public Health laws can be found in Title 22 of the Maine Revised Statutes.

Public Health Infrastructure
Part 2 of Title 22 establishes the State’s public health infrastructure including district and local health officers.

Statewide Coordination in Public Health Activities and the Universal Wellness Initiative
In 2009, the Legislature enacted a bill the coordinates and streamlines the public health system in this State. It prepares the state public health system for national federally recognized public health accreditation and ensures the effective, efficient, and evidence-based delivery of essential public health services. It recognizes and formally establishes Healthy Maine Partnerships, district coordinating councils for public health, and the Statewide Coordinating Council for Public Health. It also establishes a universal wellness initiative using the existing resources of the public health infrastructure. The initiative requires the development and distribution of a resource toolkit for the uninsured and a health risk assessment for all people of the State with a focus on the uninsured and those facing health disparities. It also requires the Department of Health and Human Services, Maine Center for Disease Control and Prevention to issue an annual report card on health for each public health district in the State and for the state health plan to publish the report cards.

For more on the Public Health Infrastructure see Title 22, MRSA, Chapter 152.

Fund for a Healthy Maine
The Fund for a Healthy Maine (FHM) was created by the Maine Legislature in 1999 to receive and disburse Maine’s share of payments from a settlement of major lawsuits involving tobacco companies. Maine participated in the national tobacco settlement because many Maine people have suffered disease and death as a result of tobacco use. The Fund remains Maine's primary investment in public health and preventative healthcare. For more information about the fund and approved uses of the funds, see 22 MRSA §1511.

Control of Notifiable Diseases and Conditions
Maine’s statutes on responding to notifiable diseases and conditions is contained in Title 22 MRSA, Chapter 250.

Health emergency
In the event of an actual or threatened epidemic or public health threat, the department may declare that a health emergency exists and may adopt emergency rules for the protection of the public health relating to:
A. Procedures for the isolation and placement of infected persons for purposes of care and treatment or infection control;
B. Procedures for the disinfection, seizure or destruction of contaminated property; and
C. The establishment of temporary facilities for the care and treatment of infected or exposed persons, which are subject to the supervision and regulations of the department and to the limitations set forth in section 807.

22 MRSA §802 (2)

Inspection
If the department has reasonable grounds to believe that there exists a public health threat, either on public or private property, a duly authorized agent of the department may enter any place, building, vessel, aircraft or common carrier with the permission of the owner, agent or occupant where the public health threat is reasonably believed to exist and may inspect and examine the same. If entry is refused, that agent shall apply for an inspection warrant from the District Court pursuant to Title 4, section 179, prior to conducting the inspection.

22 MRSA §803

Control of communicable diseases
The department may establish procedures for agents of the department to use in the detection, contacting, education, counseling and treatment of individuals having or reasonably believed to have a communicable disease. The procedures shall be adopted in accordance with the requirements of this chapter and with the rules adopted under section 802.

For purposes of carrying out this chapter, the department may designate facilities and private homes for the confinement and treatment of infected persons posing a public health threat. The department may designate any such facility in any hospital or other public or private institution, other than a jail or correctional facility. Designated institutions must have necessary clinic, hospital or confinement facilities as may be required by the department. The department may enter into arrangements for the conduct of these facilities with public officials or persons, associations or corporations in charge of or maintaining and operating these institutions.

22 MRSA §807

For more on investigating, examinaning and responding to individuals believed to have a communicable disease, see 22 MRSA §§807-814

Privileged or confidential communications in public health situations
Privileges abrogated: Subject to the limitations imposed by United States Code, Title 42, Sections 290dd-3 and 290ee-3, the physician-patient and psychotherapist-patient privileges under the Maine Rules of Evidence and those confidential communications described under Title 5, section 19203, Title 24-A, section 4224, Title 32, section 7005 and Title 34-B, section 1207, are abrogated to the extent necessary to permit reporting to the Bureau of Health any incidents of notifiable disease or condition; cooperating with the Bureau of Health or an intervention team appointed by the Bureau of Health in investigating a case of a notifiable disease or condition or suspected epidemic, or taking preventive action in such a case; or giving evidence in a proceeding pursuant to this chapter. Information released to the bureau pursuant to this section must be kept confidential and may not be disclosed by the bureau except as provided in section 824 and Title 5, section 19203, subsection 8.

22 MRSA §815 (1)

Limitation: Statements made to a licensed mental health or medical professional in the course of counseling, diagnosis, therapy, treatment or evaluation when the privilege is abrogated under this section may not be used against the client in a criminal proceeding.

22 MRSA §815 (2)
Immunity
For private institutions
Any private institution, its employees or agents are immune from civil liability to the extent provided in
Title 14, chapter 741, as if that institution were a state agency and its employees and agents were state
employees, for any acts taken to provide for the confinement or restraint of a person committed pursuant
to this chapter or for participating in reporting under this chapter, or for engaging in any prescribed care
within the meaning of this chapter in support of the State's response to a declared extreme public health
emergency in accordance with the provisions of this chapter and Title 37-B, chapter 13, subchapter 2.

Reporting and proceedings
Any person participating in reporting under this chapter or participating in a related communicable
disease investigation or proceeding, including, but not limited to, any person serving on or assisting a
multidisciplinary intervention team or other investigating or treatment team, is immune from civil liability
for the act of reporting or participating in the investigation or proceeding in good faith. Good faith does
not include instances when a false report is made and the reporting person knows or should know the
report is false.

Reporting communicable diseases
General
Whenever any physician knows or has reason to believe that any person whom the physician examines or
cares for has or is afflicted with any disease or condition designated as notifiable, that physician shall
notify the department and make such a report as may be required by the rules of the department. Reports
must be in the form and content prescribed by the department and the department shall provide forms for
making required reports.

Time requirements
The reporting of a notifiable disease or condition must be made by telephone to the department
immediately upon determination that a person has that disease and must be followed by a written report
mailed to the department within 48 hours.

Confidentiality
Any person who receives information pursuant to this chapter shall treat as confidential the names of
individuals having or suspected of having a notifiable disease or condition, as well as any other
information that may identify those individuals. This information may be released to the department for
adult or child protection purposes in accordance with chapters 958-A and 1071, or to other public health
officials, agents or agencies or to officials of a school where a child is enrolled, for public health
purposes, but that release of information must be made in accordance with Title 5, chapter 501, where
applicable. In a the event of an actual or threatened epidemic or outbreak or public health threat or
emergency, as declared by the Director of the Bureau of Health, the information may also be released to
private health care providers and health and human services agencies for the purpose of carrying out
public health functions as authorized by this chapter. Information not reasonably required for the purposes
of this section may not be released. All information submitted pursuant to this chapter that does not name
or otherwise identify individuals having or suspected of having a notifiable disease or condition may be
made available to the public at the sole discretion of the department.

Penalties
Any person who knowingly and willfully fails to comply with reporting requirements for notifiable
diseases or conditions commits a civil violation for which a fine of not more than $250 may be adjudged.
A person who knowingly or recklessly makes a false report under section 822 or who knowingly violates
section 824, is civilly liable for actual damages suffered by a person reported upon and for punitive
damages and commits a civil violation for which a fine of not more than $500 may be adjudged.
Mandatory blood-borne pathogen test

Definitions

A. Bona fide occupational exposure: means skin, eye, mucous membrane or parenteral contact of a person with the potentially infectious blood or other body fluids of another person that results from the performance of duties by the exposed person in the course of employment.

B. Employer; employer of the person exposed: includes a self-employed person who is exposed to the potentially infectious blood or other body fluids of another person.

C. Informed consent: means consent that is:
   1. Based on an actual understanding by the person to be tested:
      a. That the test is being performed;
      b. Of the nature of the test;
      c. Of the persons to whom the results of that test may be disclosed;
      d. Of the purpose for which the test results may be used; and
      e. Of any reasonably foreseeable risks and benefits resulting from the test; and
      f. Wholly voluntary and free from express or implied coercion.

Judicial consent to blood-borne pathogen test

A. Petition
   Any person who experiences a bona fide occupational exposure may petition the District Court with jurisdiction over the facility or other place where the exposure occurred to require the person whose blood or body fluid is the source of the exposure to submit to a blood-borne pathogen test and to require that the results of the test be provided to the petitioner as long as the following conditions have been met:
   1. The exposure to blood or body fluids creates a significant risk of infection with a blood-borne pathogen, as defined by the Bureau of Health through the adoption of rules;
   2. The authorized representative of the employer of the person exposed has informed the person whose blood or body fluid is the source of the occupational exposure and has sought to obtain written informed consent from the person whose blood or body fluid is the source of the exposure; and
   3. Written informed consent was not given by the person whose blood or body fluid is the source of the exposure and that person has refused to be tested.

B. Pre-hearing duties of the court (omitted)

C. Hearings (omitted)

D. Determination
   The court shall require the person whose blood or body fluid is the source of the exposure to obtain a blood-borne pathogen test and shall require that the results of the test be provided to the petitioner only if the petitioner proves by a preponderance of the evidence that:
   1. The exposure to blood or body fluids of the person created a significant risk of infection with a blood-borne pathogen as defined by the Bureau of Health through the adoption of rules;
   2. An authorized representative of the employer of the person exposed has informed the patient of the occupational exposure and has sought to obtain written informed consent from the person whose blood or body fluid is the source of the exposure; and
   3. Written informed consent was not given by the person whose blood or body fluid is the source of the exposure and that person has refused to be tested.

E. Consent
   The court may not order a person whose blood or body fluid is the source of the exposure to obtain a blood-borne pathogen test unless the employee exposed to the blood or body fluids of that person has consented to and obtained a blood-borne pathogen test immediately following that documented exposure.
F. **Costs**

The employer of the person exposed is responsible for the petitioner's reasonable costs related to obtaining the results of a blood-borne pathogen test pursuant to this section, including the payment of the petitioner's attorney's fees.

22 MRSA §832

**Occupational Diseases (see Occupational Health) (see also Agency Rules Links)**

**Duties of physicians and hospitals**

All physicians or hospitals shall report to the Department of Human Services all persons diagnosed as having an occupational disease no later than 30 days from the date of diagnosis or from discharge from a hospital. The report shall include any factor known to the physician which is suspected of being a contributing factor to the disease, including, but not limited to, whether or not the person smokes and, if so, the frequency of smoking.

A physician, upon notification by the Department of Human Services, shall report to the department any further information requested by the department concerning any person now or formerly under his care, diagnosed as having or having had an occupational disease.

No physician or hospital complying with the reporting requirements of this section may be liable for any civil damages as a result of those acts.

22 MRSA §1493

**Confidentiality**

The names and related information which may identify individuals having an occupational disease shall be confidential and may be released only to other public health officials, agents or agencies, or by court order or by written authorization of the individual being reported on. All other information submitted pursuant to this chapter may be made available to the public.

22 MRSA §1494

**Employment During Extreme Public Health Emergency**

**Required leave**

An employer shall grant reasonable and necessary leave from work, with or without pay, for an employee for the following reasons related to an extreme public health emergency:

A. The employee is unable to work because the employee is under individual public health investigation, supervision or treatment related to an extreme public health emergency;

B. The employee is unable to work because the employee is acting in accordance with an extreme public health emergency order;

C. The employee is unable to work because the employee is in quarantine or isolation or is subject to a control measure in accordance with extreme public health emergency information or directions issued to the public, a part of the public or one or more individuals;

D. The employee is unable to work because of a direction given by the employee's employer in response to a concern of the employer that the employee may expose other individuals in the workplace to the extreme public health emergency threat; or

E. The employee is unable to work because the employee is needed to provide care or assistance to one or more of the following individuals: the employee's spouse or domestic partner as defined under Title 18-A, section 1-201, subsection (10-A); the employee's parent; or the employee's child or child for whom the employee is the legal guardian.

For purposes of this subsection, "extreme public health emergency" has the same meaning as in Title 22, section 801, subsection 4-A.

26 MRSA §875 (1)

**Exceptions**

An employer who fails to grant a leave under subsection 1 is not in violation of subsection 1 if:
A. The employer would sustain undue hardship from the employee's absence, including the need to
downsiz e for legitimate reasons related to the impact of the extreme public health emergency on
the operation of the business;
B. The request for leave is not communicated to the employer within a reasonable time under the
circumstances; or
C. The employee to be granted leave under subsection 1, paragraph E is a state, county or municipal
employee whose responsibilities are related to services necessary for protecting the public's
health and safety in an extreme public health emergency if the employer requires the employee to
work, unless there are no other options or persons able to provide care or assist one or more of the
individuals listed under subsection 1, paragraph E.

26 MRSA §875 (2)

Duration of leave
Leave granted under subsection 1 must be for the duration of an extreme public health emergency and for
a reasonable and necessary time period following the termination of the extreme public health emergency
for diseases or conditions that are contracted or exposures that occurred during the extreme public health
emergency.

26 MRSA §875 (3)

Documentation.
Upon the employee's return to work, the employer has the right to request and receive written
documentation from a physician or public health official supporting the employee's leave.

26 MRSA §875 (4)

Benefits retained
The taking of leave under this subchapter may not result in the loss of any employee benefits accrued
before the date on which the leave commenced and does not affect the employee's right to health
insurance benefits on the same terms and conditions as applicable to similarly situated employees. For
any leave that extends beyond the time described in subsection 3, the employer shall allow an employee
to continue the employee's benefits at the employee's expense. The employer and employee may negotiate
for the employer to maintain benefits at the employer's expense for the duration or any portion of this
extended leave.

26 MRSA §875 (5)

Civil penalties
The Department of Labor may assess civil penalties of up to $200 for each violation of this section if
notice of the violation is given to the employer and the department within 6 months of the occurrence.

26 MRSA §875 (6)

Application
This subchapter applies to all public and private employers, including the State and its political
subdivisions.

26 MRSA §875 (7)

Expedited Partner Therapy
For the full statute regarding expedited partner therapy, see Title 22, Sections 1241 and 1242 of the Maine
Revised Statutes.

General
Notwithstanding any other provision of law, a health care professional who makes a clinical diagnosis of
a sexually transmitted disease may provide expedited partner therapy for the treatment of the sexually
transmitted disease if in the judgment of the health care professional the sexual partner is unlikely or
unable to present for comprehensive health care, including evaluation, testing and treatment for sexually
transmitted diseases. Expedited partner therapy is limited to a sexual partner who may have been exposed
to a sexually transmitted disease within the previous 60 days and who is able to be contacted by the
patient.
"Expedited partner therapy" means prescribing, dispensing, furnishing or otherwise providing prescription antibiotic drugs to the sexual partner or partners of a person clinically diagnosed as infected with a sexually transmitted disease without physical examination of the partner or partners.

Counseling
A health care professional who provides expedited partner therapy shall provide counseling for the patient, including advice that all women and symptomatic persons, and in particular women with symptoms suggestive of pelvic inflammatory disease, are encouraged to seek medical attention. The health care professional shall also provide written materials provided by the department to be given by the patient to the sexual partner that include at a minimum the following:

A. A warning that a woman who is pregnant or might be pregnant should not take certain antibiotics and should immediately contact a health care professional for an examination;
B. Information about the antibiotic and dosage provided or prescribed; clear and explicit allergy and side effect warnings, including a warning that a sexual partner who has a history of allergy to the antibiotic or the pharmaceutical class of antibiotic should not take the antibiotic and should be immediately examined by a health care professional;
C. Information about the treatment and prevention of sexually transmitted diseases;
D. The requirement of abstinence until a period of time after treatment to prevent infecting others;
E. Notification of the importance of the sexual partner's receiving examination and testing for the human immunodeficiency virus and other sexually transmitted diseases and information regarding available resources;
F. Notification of the risk to the sexual partner, others and the public health if the sexually transmitted disease is not completely and successfully treated;
G. The responsibility of the sexual partner to inform that person's sexual partners of the risk of sexually transmitted disease and the importance of prompt examination and treatment;
H. Advice to all women and symptomatic persons, and in particular women with symptoms suggestive of pelvic inflammatory disease, to seek medical attention; and
I. Information other than the information under paragraphs A to H as determined necessary by the department.

Immunity for health care professional
A health care professional who provides expedited partner therapy in good faith without fee or compensation under this section and provides counseling and written materials as required in subsection 1 is not subject to civil or professional liability in connection with the provision of the therapy, counseling and materials, except in the case of willful and wanton misconduct. A health care professional is not subject to civil or professional liability for choosing not to provide expedited partner therapy.

Motorcycle helmet requirement for persons under 18 years of age
Requirement
The following persons must wear protective headgear:
A. If under 18 years of age, a passenger on a motorcycle or moped or in an attached side car;
B. If under 18 years of age, an operator of a motorcycle or moped.

Compliance
An operator of a motorcycle or moped or a parent or guardian may not allow a passenger under the age of 18 years to ride in violation of this section.
School health
School health advisory
Each school board shall appoint one or more physicians or family or pediatric nurse practitioners to serve as a school health advisor.

Duties
The school health advisor shall advise the administrative unit on school health issues, policies and practices and may also perform any other health-related functions assigned by the board.

Other functions
A school health advisor may perform other medical and health-related duties assigned by the school board which may include all or some of the following:
A. Examine and diagnose students referred by teachers and other school employees to protect against the outbreak of contagious diseases in the schools;
B. Examine students for participation in physical education and athletic activities;
C. Advise and serve as medical consultant to the school nurse; or
D. Examine school employees and property if the physician believes it is necessary to protect the health of students.

Prohibition
A school health advisor may not treat any student examined under this subchapter unless the school health is also the student's personal health care provider.

Appointment
Appointment shall be on a yearly basis.

Epinephrine in Schools
Physicians and school health advisors are authorized to enter into collaborative practice agreements with school nurses to delegate the administration of epinephrine to non-licensed school personnel.

Medical Marijuana in Schools
A primary caregiver as defined in statute may possess and administer marijuana in a nonsmokeable form in a school bus and on the grounds of the preschool or primary or secondary school in which a minor qualifying patient is enrolled only if:
A. A medical provider has provided the minor qualifying patient with a current written certification for the medical use of marijuana under this chapter; and
B. Possession of marijuana in a nonsmokeable form is for the purpose of administering marijuana in a nonsmokeable form to the minor qualifying patient.

Tracking Prevalence of Childhood Obesity
A school nurse or trained screener shall collect body mass index data from students in the school administrative unit in accordance with rules of the Department of Health and Human Services. Data may not be collected from a student whose parent or guardian objects on religious or philosophical grounds. A school nurse shall report the data collected to the Department of Health and Human Services, Maine Center for Disease Control and Prevention. Data reported pursuant to this subsection may be reported in the aggregate only and may not identify an individual student.

School Physical Education
The Commissioner of Education shall conduct a statewide assessment, using a survey or sampling methodology, of the current physical education capacities of elementary schools in the State.
The Obesity and Chronic Disease Fund is established as an interest-bearing account administered by the Department of Education. Balances in the fund may be used to pay for new equipment, new staff training, new personnel, new administrative costs and other expenses not related to an existing physical education program and for the implementation of a new physical education program for elementary schools.

For more information on school health, including immunization, screening and nutrition requirements, and bullying prevention see Title 20-A, Chapter 223 of the Maine Statutes.

For information on bullying and head injury policies, see also 20-A MRSA § 254.

The requirement for scoliosis screening of students in schools has been repealed.

**Menu Labeling**
A chain restaurant with locations in the State of Maine shall state on a food display tag, menu or menu board the total amount of calories per serving of each food and beverage item listed for sale on the food display tag, menu or menu board. A menu or menu board or written nutrition information provided to a customer by a chain restaurant must contain the following statement in a clear and conspicuous manner and in a prominent location: "To maintain a healthy weight, a typical adult should consume approximately 2,000 calories per day; however, individual calorie needs may vary.

"Chain restaurant" means an eating establishment that does business under the same trade name in 20 or more locations, at least one of which is located in the State, that offers predominantly the same type of meals, food, beverages or menus, regardless of the type of ownership of an individual location. "Chain restaurant" does not include a grocery store.

For more information, see Public Laws 2009, Chapter 359.

**Lyme Disease**
Every health care provider that orders a laboratory test for the presence of Lyme disease shall provide the patient with a copy of the results of the test.

**Treatment**
A physician may prescribe, administer or dispense long-term antibiotic therapy for a therapeutic purpose to eliminate infection or to control a patient's symptoms upon making a clinical diagnosis that the patient has Lyme disease or displays symptoms consistent with a clinical diagnosis of Lyme disease. The physician shall document the clinical diagnosis and treatment in the patient's medical record. The clinical diagnosis must be based on knowledge obtained through medical history and physical examination only or in conjunction with testing that provides supportive data for the clinical diagnosis.

**Research, Maine Biomedical Research Program**
The Maine Biomedical Research Program is established to promote economic development and jobs in the State primarily by making state investments in organizations with successful results in attracting biomedical research funds from specified grant sources. As a secondary purpose, the Maine Biomedical Research Program is intended to provide incentives for small eligible institutions to grow. The program shall disburse program funds from the Maine Biomedical Research Fund to eligible institutions.

For more information regarding eligibility and the application process. See 5 MRSA §13103.
Research, Human Subjects (Select Sections)

“The Common Rule”

All human subjects research that is either funded or support by the federal government falls under the jurisdiction of Title 45 of the Code of Federal Regulation, Section 46, The Protections of Human Subjects (“The Common Rule”). As stated in 45 CFR § 46:

“…this policy applies to all research involving human subjects conducted, supported or otherwise subject to regulation by any federal department or agency which takes appropriate administrative action to make the policy applicable to such research. This includes research conducted by federal civilian employees or military personnel, except that each department or agency head may adopt such procedural modifications as may be appropriate from an administrative standpoint. It also includes research conducted, supported, or otherwise subject to regulation by the federal government outside the United States.”

45 CFR §46.101

The Common Rule, enforcement and regulatory oversight for HHS funded research is administrated by the Office of Human Research Protections (OHRP). For more information, please click on any of the links below:

- Full Text of the Common Rule
- NIH Required Training for Human Participant Protections
- The Belmont Report
- IRB Registration Information
- OHRP Guidance by Topic
- FDA Compliance Website
- CDC Research Website
- NIH Research Resources

It should be noted that all of the following federal agencies have adopted similar version of the Common Rule governing research support by them:

Department of Agriculture 7 CFR §1e
Department of Energy 10 CFR §745
National Aeronautics and Space Administration 14 CFR §1230
Department of Commerce 15 CFR §27
Consumer Product Safety Commission 16 CFR §1028
Agency for International Development 22 CFR §225
Department of Housing and Urban Development 24 CFR §60
Department of Justice 28 CFR §46
Department of Defense 32 CFR §219
Department of Education 34 CFR §97
Department of Veterans Affairs 38 CFR §16
Environmental Protection Agency 40 CFR §26
National Science Foundation 45 CFR §690
Department of Transportation 49 CFR §11
Central Intelligence Agency By Executive Order 12333
Department of Agriculture By Statute

Select Definitions

- Research: “…a systematic investigation, including research development, testing and evaluation, designed to develop or contribute to generalizable knowledge. Activities which meet this
definition constitute research for purposes of this policy, whether or not they are conducted or supported under a program which is considered research for other purposes.”

B. **Research subject to regulation:** and similar terms are intended to encompass those research activities for which a federal department or agency has specific responsibility for regulating as a research activity, (for example, Investigational New Drug requirements administered by the Food and Drug Administration). It does not include research activities which are incidentally regulated by a federal department or agency solely as part of the department's or agency's broader responsibility to regulate certain types of activities whether research or non-research in nature (for example, Wage and Hour requirements administered by the Department of Labor).

C. **Human subject:** means a living individual about whom an investigator (whether professional or student) conducting research obtains:
   1. Data through intervention or interaction with the individual, or
   2. Identifiable private information.

D. **Intervention:** includes both physical procedures by which data are gathered (for example, venipuncture) and manipulations of the subject or the subject's environment that are performed for research purposes. Interaction includes communication or interpersonal contact between investigator and subject. Private information includes information about behavior that occurs in a context in which an individual can reasonably expect that no observation or recording is taking place, and information which has been provided for specific purposes by an individual and which the individual can reasonably expect will not be made public (for example, a medical record). Private information must be individually identifiable (i.e., the identity of the subject is or may readily be ascertained by the investigator or associated with the information) in order for obtaining the information to constitute research involving human subjects.

E. **Institutional Review Board (IRB):** means an institutional review board established in accord with and for the purposes expressed in this policy.

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(87) **Rural Medical Access Program (see also Agency Rules Links)**

The Rural Medical Access Program is governed by [Title 24-A M.R.S.A., Chapter 75, §6301-6311](http://www.maine.gov/pfr/insurance/regulated/insurance_companies/insurer/rmap/index.html) and [Maine Regulation Chapter 630](#). Please refer to the Statute and Regulation for a complete understanding of the Program.

The purpose of the RMAP is to promote prenatal services in underserved areas in Maine. [Title 24-A, M.R.S.A., §6302](#) The RMAP provides medical malpractice premium assistance to qualified eligible physicians who are licensed and practicing in Maine, who provide prenatal care and delivery services, and practice at least 50% in underserved areas of the state. [§6307](#) Eligibility for premium assistance, as well as underserved areas in Maine, is determined by the state’s Department of Health and Human Services.

For more information, visit the Bureau of Insurance:


(88) **Sentinel Events, Reporting of Definitions**

A. **Division:** means the Department of Health and Human Services, Division of Licensing and Regulatory Services.

B. **Health care facility:** means a state institution as defined under Title 34-B, chapter 1 or a health care facility licensed by the division, except that it does not include a facility licensed as a nursing facility or licensed under chapter 1664. “Health care facility” includes a general and specialty
hospital, an ambulatory surgical facility, an end-stage renal disease facility and an intermediate care facility for persons with mental retardation or developmental disabilities.

C. **Immediate jeopardy:** means a situation in which the provider’s noncompliance with one or more conditions of participation in the federal Medicare program has caused, or is likely to cause, serious injury, harm or impairment to or death of a patient.

D. **Major permanent loss of function:** means sensory, motor, physiological or intellectual impairment that was not present at the time of admission and requires continued treatment or imposes persistent major restrictions in activities of daily living.

E. **Near miss:** means an event or situation that did not produce patient injury, but only because of chance, which may include, but is not limited to, robustness of the patient or a fortuitous, timely intervention.

F. **Root cause analysis:** means a structured process for identifying the causal or contributing factors underlying adverse events. The root cause analysis follows a predefined protocol for identifying these specific factors in causal categories.

G. **Sentinel event:** means:
   1. An unanticipated death, or patient transfer to another health care facility, unrelated to the natural course of the patient’s illness or underlying condition or proper treatment of that illness or underlying condition in a health care facility;
   2. A major permanent loss of function unrelated to the natural course of the patient’s illness or underlying condition or proper treatment of that illness or underlying condition in a health care facility that is present at the time of the discharge of the patient. If within 2 weeks of discharge from the facility, evidence is discovered that the major loss of function was not permanent, the health care facility is not required to submit a report pursuant to section 8753, subsection 2;
   3. An unanticipated perinatal death or major permanent loss of function in an infant with a birth weight over 2,500 grams that is unrelated to the natural course of the infant’s mother’s illness or underlying condition or proper treatment of that illness or underlying condition in a health care facility; and
   4. Other serious and preventable events as identified by a nationally recognized quality forum and determined in rules adopted by the department pursuant to section 8756.

**22 MRSA §8752**

**Mandatory reporting of sentinel events**

A health care facility shall notify the division whenever a sentinel event has occurred, as provided in this chapter.

**Notification**

A health care facility shall notify the division of a sentinel event by the next business day after the event occurred or the next business day after the facility discovers that the event occurred. The notification must include the date and time of notification, the name of the health care facility and the type of sentinel event pursuant to section 8752, subsection 4-A.

**22 MRSA §8753 (1)**

**Reporting**

The health care facility shall file a written report no later than 45 days following the notification of the occurrence of a sentinel event pursuant to subsection 1. The written report must be signed by the chief executive officer of the facility and must contain the following information:

A. Facility name and address;
B. Name, title and phone number of the contact person for the facility;
C. The date and time of the sentinel event;
D. The type of sentinel event and a brief description of the sentinel event; and
E. A thorough and credible root cause analysis. A root cause analysis is thorough and credible only in accordance with the following.

1. A thorough root cause analysis must include: a determination of the human and other factors most directly associated with the sentinel event and the processes and systems related to its occurrence; an analysis of the underlying systems and processes to determine where redesign might reduce risk; an inquiry into all areas appropriate to the specific type of event; an identification of risk points and their potential contributions to the event; a determination of potential improvement in processes or systems that would tend to decrease the likelihood of such an event in the future or a determination, after analysis, that no such improvement opportunities exist; an action plan that identifies changes that can be implemented to reduce risks or formulates a rationale for not undertaking such changes; and, where improvement actions are planned, an identification of who is responsible for implementation, when the action will be implemented and how the effectiveness of the action will be evaluated.

2. A credible root cause analysis must include participation by the leadership of the health care facility and by the individuals most closely involved in the processes and systems under review, is internally consistent without contradictions or unanswered questions, provides an explanation for all findings, including those identified as “not applicable” or “no problem,” and includes the consideration of any relevant literature.

3. The root cause analysis submitted to the division may exclude protected professional competence review information pursuant to the Maine Health Security Act.

Cooperation
A health care facility that has filed a notification or a report of the occurrence of a sentinel event under this section shall cooperate with the division as necessary for the division to fulfill its duties under section 8754.

Immunity
A person who in good faith reports a near miss, a suspected sentinel event or a sentinel event or provides a root cause analysis pursuant to this chapter is immune from any civil or criminal liability for the act of reporting or participating in the review by the division. "Good faith" does not include instances when a false report is made and the person reporting knows the report is false. This subsection may not be construed to bar civil or criminal action regarding perjury or regarding the sentinel event that led to the report.

Near miss notification
A health care facility may notify the division of the occurrence of a near miss. Should a facility report a near miss, the notification must include the date and time of notification, the name of the health care facility and the type of event or situation pursuant to section 8752, subsection 4-A that is related to the near miss.

Compliance
Oversight
The division shall place primary emphasis on ensuring effective corrective action by the facility.

Penalties
When the division determines that a health care facility failed to report a sentinel event pursuant to this chapter, the health care facility is subject to a penalty imposed in conformance with Title 5, chapter 375, subchapter 4 and payable to the State of not more than $10,000 per violation. If the facility in good faith notified the division of a suspected sentinel event and the division later determines it is a sentinel event, the facility is not subject to a penalty for that event. Funds collected pursuant to this section must be
deposited in a dedicated special revenue account to be used to support sentinel event reporting and education.

**Administrative hearing and appeal**

To contest the imposition of a penalty under this section, a health care facility must submit to the division a written request for an administrative hearing within 10 days of notice of imposition of a penalty pursuant to this section. Judicial appeal must be in accordance with Title 5, chapter 375, subchapter 7.

**Injunction**

Notwithstanding any other remedies provided by law, the Office of the Attorney General may seek an injunction to require compliance with the provisions of this chapter.

**Enforcement**

The Office of the Attorney General may file a complaint with the District Court seeking injunctive relief for violations of this chapter.

**Targeted surveillance for methicillin-resistant Staphylococcus aureus (MRSA)**

All hospitals licensed under chapter 405 shall perform targeted surveillance for methicillin-resistant Staphylococcus aureus in high-risk populations, as defined by the Maine Quality Forum established pursuant to Title 24-A, section 6951, consistent with the federal Centers for Disease Control and Prevention guidelines.

**Sexual Misconduct**

**Sexual Misconduct by Physicians and Physician Assistants (see also Agency Rules Links)**

**Definition**

Physician/physician assistant sexual misconduct is behavior that exploits the physician/physician assistant-patient relationship in a sexual way. This behavior is nondiagnostic and/or nontherapeutic, may be verbal or physical, and may include expressions or gestures that have a sexual connotation or that a reasonable person would construe as such. Sexual misconduct is considered incompetence and unprofessional conduct as defined by 32 M.R.S.A 2591-A (2) and 32 M.R.S.A. 3282-A (2).

**Levels of Misconduct**

Under Maine Law there are two levels of sexual misconduct. Behavior listed in both levels may be the basis for disciplinary action.

A. **Sexual violation**

Any conduct by a physician/physician assistant with a patient that is sexual or may be reasonably interpreted as sexual, even when initiated by or consented to by a patient, including but not limited to:

1. Sexual intercourse, genital to genital contact;
2. Oral to genital contact;
3. Oral to anal contact or genital to anal contact;
4. Kissing in a sexual manner (e.g. - French kissing);
5. Any touching of a body part for any purpose other than appropriate examination, treatment, or comfort, or where the patient has refused or has withdrawn consent;
6. Encouraging the patient to masturbate in the presence of the physician/physician assistant or masturbation by the physician/physician assistant while the patient is present; and,
7. Offering to provide practice-related services, such as drugs, in exchange for sexual favors.
B. **Sexual impropriety**

   Behavior, gestures, or expressions by the physician/physician assistant that are seductive, sexually suggestive, or sexually demeaning to a patient, including but not limited to:
   
   1. Kissing;
   2. Disrobing, draping practices or touching of the patient’s clothing that reflect a lack of respect for the patient’s privacy; deliberately watching a patient dress or undress, instead of providing privacy for disrobing;
   3. Subjecting a patient to an examination in the presence of another when the physician/physician assistant has not obtained the verbal or written consent of the patient or when consent has been withdrawn;
   4. Examination or touching of genitals without the use of gloves;
   5. Inappropriate comments about or to the patient, including but not limited to making sexual comments about a patient’s body or underclothing; making sexualized or sexually demeaning comments to a patient, criticizing the patient’s sexual orientation (homosexual, heterosexual, or bisexual); making comments about potential sexual performance during an examination or consultation (except when the examination or consultation is pertinent to the issue of sexual function or dysfunction); requesting details of sexual history or sexual likes or dislikes when not clinically indicated;
   6. Using the physician/physician assistant-patient relationship to solicit a date or initiate romantic relationship;
   7. Initiation by the physician/physician assistant of conversation regarding the sexual problems, preferences, or fantasies of the physician/physician assistant; and,
   8. Examining the patient without verbal or written consent.

**Sanctions**

A. **Sexual violation**

   Findings of sexual violations are egregious enough to warrant revocation of a physician/physician assistant’s medical license. Boards may, at times, find that mitigating circumstances do exist and, may impose a lesser sanction.

B. **Sexual impropriety**

   Findings of sexual impropriety will result in harsh sanction, which may include revocation. Special consideration should be given to at least the following when determining an appropriate sanction:
   
   1. Patient harm;
   2. Severity of impropriety;
   3. Culpability of licensee;
   4. Psychotherapeutic relationship;
   5. Inappropriate termination of physician/physician assistant-patient relationship;
   6. Age of patient;
   7. Physical /mental capacity of patient;
   8. Number of times behavior occurred;
   9. Number of patients involved;
   10. Period of time relationship existed; and,

**Department of Professional and Financial Regulation, Board of Licensure in Medicine/Board of Osteopathic Licensure, Rule Chapter 10**

**Sexual activity with recipient of services prohibited**

A person who owns, operates or is an employee of an organization, program or residence that is operated, administered, licensed or funded by the Department of Behavioral and Developmental Services or the Department of Human Services may not engage in a sexual act, as defined in Title 17-A, section 251, subsection 1, paragraph C, with another person or subject another person to sexual contact, as defined in
Title 17-A, section 251, subsection 1, paragraph D, if the other person, not the actor's spouse, is a person with mental illness who receives therapeutic, residential or habilitative services from the organization, program or residence.

Transportation and storage of forensic examination kits
If an alleged victim of gross sexual assault has a forensic examination and has not reported the alleged offense to a law enforcement agency when the examination is complete, the licensed hospital or licensed health care practitioner that completed the forensic examination shall notify the nearest law enforcement agency. That law enforcement agency shall transport the completed kit, identified only by a tracking number assigned by the kit manufacturer, to its evidence storage facility. The law enforcement agency shall store the kit for at least 90 days from the time of receipt. If during that 90-day period the alleged victim reports the offense to a law enforcement agency, the investigating agency shall take possession of the kit.

In the case of a forensic examination performed under Title 24, section 2986, subsection 5, the law enforcement agency must immediately notify the district attorney for the district in which the hospital or health care practitioner is located that such a forensic examination has been performed and a forensic examination kit has been completed under Title 24, section 2986, subsection 5.

Sexual Assault Support Information
To access hotline advocates for free and confidential support, or to receive more information on sexual assault and available services, please call the toll-free statewide hotline. You will automatically be connected to the sexual assault crisis and support center nearest you.
Statewide Sexual Assault Support Hotline 1-800-871-7741 Or TTY: 1-888-458-5599

For more information, visit the Maine Coalition Against Sexual Assault (MECASA) and see the links to centers that serve each Maine county.

Certain Negligence Suits
For professional negligence actions against certain health care providers the statute of limitations on professional negligence actions is extended six years when the action is based on a sexual act or sexual contact. This applies to psychiatrists, psychologists, social workers, professional counselors, pastoral counselors, marriage and family therapist and clinical professional counselors.

(90) Staffing & Supervision
Criminal background checks
A facility or health care provider subject to the licensing or certification processes of chapter 405, 412 or 419 shall obtain, prior to hiring an individual who will work in direct contact with a consumer, criminal history record information on that individual, including, at a minimum, criminal history record information from the Department of Public Safety, State Bureau of Identification. A facility or provider subject to licensing under chapter 419 shall conduct a comprehensive background check for individuals employed in positions that have direct access to a consumer’s property, personally identifiable information, financial information or resources in accordance with applicable federal and state laws. The comprehensive background check must be conducted in accordance with state law and rules adopted by the department. The facility or health care provider shall pay for the comprehensive or criminal background check required by this section.

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Delegation
For the Maine law on physician delegation to physician assistants and other employees, see 32 MRSA §3270-A and 32 MRSA §2594-A.

Physician Assistants (See Also Physician Assistants Section)
An individual may render medical services if these services are rendered under the supervision and control of a physician or surgeon and if that individual has satisfactorily completed a training program approved by the Board of Licensure in Medicine/Board of Osteopathic Licensure and a competency examination determined by this board. Supervision and control may not be construed as requiring the personal presence of the supervising and controlling physician at the place where these services are rendered, unless a physical presence is necessary to provide patient care of the same quality as provided by the physician.

Support Staff
A physician or surgeon may delegate to the physician's or surgeon's employees or support staff certain activities relating to medical care and treatment carried out by custom and usage when the activities are under the control of the physician or surgeon. The physician delegating these activities to employees or support staff, to program graduates or to participants in an approved training program is legally liable for the activities of those individuals, and any individual in this relationship is considered the physician's agent.

For the law on physician immunity from civil and criminal liability when supervising a physician assistant on active state service in the performance of the physician assistant's duty, see 37-B MRSA §185.

(91) Sterilization
Informed consent required
Except as provided in this chapter, prior to initiating sterilization procedures on any individual, a physician shall obtain and record the informed consent of that individual. 34-B MRSA §7004 (1)

Hearing required to determine ability to give informed consent for sterilization
A hearing to determine ability to give informed consent for sterilization is required when sterilization is sought for:
A. Persons under age 18 years and not married or otherwise emancipated;
B. Persons presently under public or private guardianship or conservatorship;
C. Persons residing in a state institution providing care, treatment or security, or otherwise in state custody; or
D. Persons from whom a physician could not obtain informed consent.
34-B MRSA §7004 (2)

(92) Suicide
See AMA Positions on Physician Assisted Suicide
5.7 Physician-Assisted Suicide
H-140.952 Physician Assisted Suicide

Maine Law
Currently, Maine does not have a specific law covering the issue of physician-assisted suicides. Maine does, however, have a law that makes it a crime for any person to aid or solicit another to commit suicide.
A. A person is guilty of aiding or soliciting suicide if he intentionally aids or solicits another to commit suicide, and the other commits or attempts suicide.
B. Aiding or soliciting suicide is a Class D crime.
(93) Treatment of Terminally Ill Patients, Experimental/ Investigational

This bill authorizes manufacturers of drugs, biological products and devices that have completed Phase I of a United States Food and Drug Administration-approved clinical trial but have not yet been approved for general use and remain under clinical investigation to make them available to eligible terminally ill patients.

Definitions

Eligible patient means a person who has:
A. Received a diagnosis of a terminal illness for which no standard treatment is effective and the diagnosis has been attested by the person's treating physician;
B. Considered all treatment options approved by the United States Food and Drug Administration;
C. Not been accepted into a clinical trial within one week of completion of the clinical trial application process;
D. Received a recommendation from the person's treating physician for an investigational drug, biological product or device;
E. Given written, informed consent for the use of the investigational drug, biological product or device under paragraph D or, if the person is a minor or lacks the mental capacity to provide informed consent, whose parent or legal guardian has given written, informed consent on the person's behalf; and
F. Received documentation from the person's treating physician that the person meets all of the conditions in this subsection.

Investigational drug, biological product or device means a drug, biological product or device that has successfully completed Phase I of a United States Food and Drug Administration-approved clinical trial but has not yet been approved for general use by the United States Food and Drug Administration and remains under investigation in such a clinical trial.

Terminal illness means a disease or condition that, without life-sustaining measures, will soon result in death or in a state of permanent unconsciousness from which recovery is unlikely.

Treating physician means a physician who has primary responsibility for the care of a patient and treatment of that patient's terminal illness.

Written, informed consent means a written document signed by a patient or, if the patient is a minor or lacks the mental capacity to provide informed consent, a parent or legal guardian of the patient. The document must be attested by the patient's treating physician and a witness and include the following information:
A. An explanation of the United States Food and Drug Administration-approved treatments for the disease or condition from which the patient suffers;
B. A statement that the patient concurs with the patient's treating physician that all United States Food and Drug Administration-approved and standard treatments for the disease or condition from which the patient suffers are unlikely to prolong the patient's life;
C. Clear identification of the specific investigational drug, biological product or device that the patient is seeking to use; and
D. A description of the best and worst potential outcomes of using the investigational drug, biological product or device identified under paragraph C with a description of the most likely outcome. The description must include the possibility that new, unanticipated, different or worse symptoms might result and that death could be hastened by the proposed treatment. The
description must be based on the treating physician's knowledge of the proposed treatment in conjunction with the treating physician's knowledge of the patient's overall medical condition.

22 MRSA §2671

**Action against health care practitioner or health care provider license prohibited**

A licensing board may not revoke, refuse to renew or suspend the license of or take any action against a health care practitioner as defined in Title 24, section 2502, subsection 1-A based solely on the health care practitioner's recommendations to an eligible patient regarding access to or treatment with an investigational drug, biological product or device, as long as the recommendations are consistent with medical standards of care.

The licensing agency may not revoke, refuse to renew or suspend the license of or take any action against a health care provider as defined in Title 24, section 2502, subsection 2 based solely on the health care provider’s involvement in the care of an eligible patient using an investigational drug, biological product or device.

22 MRSA §2673

**No private cause of action created**

This chapter does not create a private cause of action against a manufacturer of an investigational drug, biological product or device or against any other person or entity involved in the care of an eligible patient using the investigational drug, biological product or device for any harm done to the eligible patient resulting from the investigational drug, biological product or device if the manufacturer or other person or entity is complying in good faith with the provisions of this chapter and has exercised reasonable care.

22 MRSA §2675

**Optional participation of health care practitioners and providers**

This chapter does not require a health care practitioner who is licensed in the State or a health care provider that is licensed in the State to provide any service related to an investigational drug, biological product or device.

22 MRSA §2677

(94) **Utilization Review (see also Agency Rules Links)**

**Review entities**

**Licensure**

A person, partnership or corporation, other than an insurer, nonprofit service organization, health maintenance organization, preferred provider organization or employee of those exempt organizations, that performs medical utilization review services on behalf of commercial insurers, nonprofit service organizations, 3rd-party administrators, health maintenance organizations, preferred provider organizations or employers shall apply for licensure by the Bureau of Insurance and pay an application fee of not more than $400 and an annual license fee of not more than $100; except that programs of review of medical services for occupational claims compensated under Title 39-A are subject only to the certification requirements of that title and are not subject to licensure under this section. A person, partnership or corporation, other than an insurer or nonprofit service organization, health maintenance organization, preferred provider organization or the employees of exempt organizations, may not perform utilization review services or medical utilization review services unless the person, partnership or corporation has received a license to perform those activities.

24-A MRSA §2771 (1)

**Listing**

The Bureau of Insurance shall compile and maintain a current listing of persons, partnerships or corporations licensed pursuant to this section.

24-A MRSA §2771 (2)
**Information required**

Each person, partnership or corporation licensed pursuant to this section shall, at the time of initial licensure and on or before April 1st of each succeeding year, provide the Bureau of Insurance with the following information:

A. The process by which the entity carries out its utilization review services. The information provided to the bureau must include the categories of health care personnel that perform any activities coming under the definition of utilization review and whether or not these individuals are licensed in the State. The information provided to the bureau also must include copies of any licensure agreements the utilization review entity has in effect with any entity that sells or furnishes the utilization review entity with medical utilization review criteria and the expiration date of any such agreements. If the utilization review entity develops its own medical utilization review criteria, the utilization review entity shall include copies of any policies and procedures or both for the use of the criteria;

B. The process used by the entity for addressing beneficiary or provider complaints;

C. The types of utilization review programs offered by the entity, such as:
   1. Second opinion programs;
   2. Prehospital admission certification;
   3. Preinpatient service eligibility determination; or
   4. Concurrent hospital review to determine appropriate length of stay; and

D. The process chosen by the entity to preserve beneficiary confidentiality of medical information.

As part of its initial application, the entity shall submit copies of all materials to be used to inform beneficiaries and providers of the requirements of its utilization review plans and their rights and responsibilities under the plan.

**Transition for existing entities.**

Notwithstanding subsection 1, persons, partnerships or corporations performing utilization review services on the effective date of this section shall have 90 days from its effective date to submit an application to the superintendent. The superintendent shall act upon those applications within 6 months of the date of receipt of the application, during which time the review entities may continue to perform medical utilization review services.

**Minimum standards**

A utilization review program of the applicant must meet the following minimum standards:

**Notification of adverse decisions**

Notification of an adverse decision by the utilization review agent must be provided to the insured or other party designated by the insured within a time period to be determined by the superintendent through rulemaking and must include the name of the utilization review agent who made the decision.

**Reconsideration of determinations**

All licensees shall maintain a procedure by which insureds, patients or providers may seek reconsideration of determinations of the licensee.

**Accessibility of representatives**

A representative of the licensee must be accessible by telephone to insureds, patients or providers and the superintendent may adopt standards of accessibility by rule.

**Medical utilization review criteria**

The licensee must have written medical utilization review criteria to be employed in the review process. The criteria must be available for review as a part of any review conducted pursuant to section 2774, subsection 1 and a copy of the criteria must be provided to the bureau upon request.
Information materials; confidentiality
A copy of the materials designed to inform applicable patients of the requirements of the utilization plan and the responsibilities and rights of patients under the plan and an acknowledgment that all applicable state and federal laws to protect the confidentiality of individual medical records are followed must be filed with the bureau.

Penalty for noncompliance with utilization review programs
A medical utilization review program may not recommend or implement a penalty of more than $500 for failure to provide notification. This subsection does not limit the right of insurers to deny a claim when appropriate prospective or retroactive review concludes that services or treatment rendered were not medically necessary.

Prohibited activities
A medical utilization review entity shall ensure that an employee does not perform medical utilization review services involving a health care provider or facility in which that employee has a financial interest.

Utilization review services
As used in this chapter, unless the context indicates otherwise, "utilization review services" or "medical utilization review services" means a program or process by which a person, partnership or corporation, on behalf of an insurer, nonprofit service organization, 3rd-party administrator, health maintenance organization, preferred provider organization or employer that is a payor for or that arranges for payment of medical services, seeks to review the utilization, appropriateness or quality of medical services provided to a person whose medical services are paid for, partially or entirely, by that insurer, nonprofit service organization, 3rd-party administrator, health maintenance organization, preferred provider organization or employer. The terms include these programs or processes whether they apply prospectively or retrospectively to medical services. Utilization review services include, but are not limited to, the following:
A. Second opinion programs;
B. Prehospital admission certification;
C. Preinpatient service eligibility certification;
D. Concurrent hospital review.

Enforcement
The following provisions govern enforcement of this chapter.
A. Periodic reviews
The superintendent may conduct periodic reviews of the operations of the entities licensed pursuant to this chapter to ensure that they continue to meet the minimum standards set forth in section 2772 and any applicable rules adopted by the superintendent. The superintendent may perform periodic telephone audits of licensees to determine if representatives of the licensee are reasonably accessible, as required by section 2772.
B. Action against licensee
The superintendent is authorized to take appropriate action against a licensee which fails to meet the standards of this chapter or any rules adopted by the superintendent, or who fails to respond in a timely manner to corrective actions ordered by the superintendent. The superintendent may impose a civil penalty not to exceed $1,000 for each violation, as permitted by section 12-A, or may deny, suspend or revoke the license.
C. Opportunity to provide information and request hearing
Before taking the actions authorized by this section to deny, suspend or revoke the license, the superintendent shall provide the licensee with reasonable time to supply additional information demonstrating compliance with the requirements of this chapter and the opportunity to request a hearing to be held consistent with the provisions of the Maine Administrative Procedure Act, Title 5, chapter 375.

D. Authority to adopt rules
   The superintendent may adopt rules necessary to implement the provisions of this chapter

E. Rulings on appropriateness of medical judgments not authorized
   Nothing in this chapter requires or authorizes the superintendent to rule on the appropriateness of medical decisions or judgments rendered by review entities and their agents.  

Utilization Review Data

Report required
On or before April 1st of each year, any insurer or third-party administrator which issues or administers a program or contract in this State providing coverage for hospital care that contains a provision whereby in nonemergency cases the insured is required to be prospectively evaluated through a prehospital admission certification, prepatient service eligibility program or any similar preutilization review or screening eligibility program or any similar preutilization review or screening procedure prior to the delivery of contemplated hospitalization, inpatient or outpatient health care or medical services which are prescribed or ordered by a duly licensed physician shall file a report on the results of that evaluation for the preceding year with the superintendent which shall contain the following:

A. The number and type of evaluations performed. For the purposes of this section, the term "type of evaluations" means the following preutilization review categories: presurgical inpatient days; setting of medical service, such as inpatient or outpatient services; and the number of days of service;
B. The result of the evaluation, such as whether the medical necessity of the level of service contemplated by the patient's physician was agreed to or whether benefits paid for the service were reduced by the insurer;
C. The number and result of any appeals by the patients or their physicians as a result of initial review decisions to reduce benefits for services as determined through prospective evaluations; and
D. Any complaints filed in a court of competent jurisdiction and served upon an insurer filing under this section stating a cause of action against that insurer on the basis of damages to patients alleged to have been approximately caused by a delay, reduction or denial of medical benefits by the insurer, as determined through prospective evaluations, and the determination of liability or other disposition of the complaint.

Residents
This section is applicable to evaluations, appeals and complaints relating to residents of this State only.

Confidentiality
Any information provided pursuant to this section shall not identify the patients.

Penalty for failure to notify of hospitalization
An insurance policy may not include a provision permitting the insurer to impose a penalty for the failure of any person to notify the insurer of an insured person's hospitalization for emergency treatment. For purposes of this section, "emergency treatment" has the same meaning as defined in Title 22, section 1829.
This section applies to policies and certificates executed, delivered, issued for delivery, continued or renewed in this State after the effective date of this section. For purposes of this section, all policies are deemed to be renewed no later than the next yearly anniversary of the contract date.  

**Notification prior to cancellation**

The superintendent shall, by January 1, 1991, adopt rules to provide for notification of the insured person and another person, if designated by the insured, prior to cancellation of a health insurance certificate for nonpayment of premiums, and to provide restrictions on cancellation when an insured person suffers from organic brain disease.

The rules may include, but are not limited to, definitions, minimum disclosure requirements, notice provisions and cancellation restrictions.

The requirements of this section apply to all policies and certificates executed, delivered, issued for delivery, continued or renewed in this State.

**Penalty for noncompliance with utilization review programs**

A policy or certificate issued or renewed after April 8, 1994 may not contain a provision that permits, upon retroactive review and confirmation of medical necessity, the imposition of a penalty of more than $500 for failure to provide notification under a utilization review program. This section does not limit the right of insurers to deny a claim when appropriate prospective or retroactive review concludes that services or treatment rendered were not medically necessary.

**Workers’ Compensation**

Workers' Compensation in Maine is governed by Title 39-A of the Maine Revised Statutes. You can also download the rules applicable to Workers’ Compensation from the Maine Workers’ Compensation Board.

**Entitlement to compensation and services generally**

Title 39-A MRSA, Chapter 5 includes the general rules of employee eligibility for Workers’ Compensation as well as the role of physicians in providing medical examinations, handling medical information and the medical fee schedule. Chapter 5 of the Workers’ Compensation rules includes more information on medical fee reimbursement levels and physician reporting requirements.

**Entitlement**

If an employee who has not given notice of a claim of common law or statutory rights of action, or who has given the notice and has waived the claim or rights, as provided in section 301, receives a personal injury arising out of and in the course of employment or is disabled by occupational disease, the employee must be paid compensation and furnished medical and other services by the employer who has assented to become subject to this Act.

**Selection of Independent Medical Examiners in Workers' Compensation Cases**

Examiner system

The board shall develop and implement an independent medical examiner system consistent with the requirements of this section. As part of this system, the board shall, in the exercise of its discretion, create, maintain and periodically validate a list of not more than 50 health care providers that it finds to be the most qualified and to be highly experienced and competent in their specific fields of expertise and in the treatment of work-related injuries to serve as independent medical examiners from each of the health care providers.

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care specialties that the board finds most commonly used by injured employees. An independent medical examiner must be certified in the field of practice that treats the type of injury complained of by the employee. Certification must be by a board recognized by the American Board of Medical Specialties or the American Osteopathic Association or their successor organizations. The board shall establish a fee schedule for services rendered by independent medical examiners and adopt any rules considered necessary to effectuate the purposes of this section.

**Duties**

An independent medical examiner shall render medical findings on the medical condition of an employee and related issues as specified under this section. The independent medical examiner in a case may not be the employee's treating health care provider and may not have treated the employee with respect to the injury for which the claim is being made or the benefits are being paid. Nothing in this subsection precludes the selection of a provider authorized to receive reimbursement under section 206 to serve in the capacity of an independent medical examiner. Unless agreed upon by the parties, a physician who has examined an employee at the request of an insurance company, employer or employee in accordance with section 207 during the previous 52 weeks is not eligible to serve as an independent medical examiner. **39-A MRSA §312**

See also, Chapter 4, Maine Workers’ Compensation Board Rules

**Useful Links and Websites**

Maine's Workers' Compensation Board Web Site
Workers' Compensation FAQ's
Facts About Maine's Workers' Compensation Laws

**Workers’ Compensation and HIPAA**

Many physicians and physician practices face this issue on a regular basis. In general, unless restricted by law, a covered entity must disclose protected health information about a patient being examined for a workers’ compensation claim without needing an authorization to:

- An insurer,
- State Workers’ Compensation Administrators or
- Other persons involved in the workers’ compensation system.

Provided that:

- The covered entity complies with all relevant federal and state laws; and
- The minimum necessary standard is maintained between the covered entity and the receiving party(s).

**45 CFR §164.512 (L) and OCR Guidance**

(96) **Wrongful birth; wrongful life**

It is the intent of the Legislature that the birth of a normal, healthy child does not constitute a legally recognizable injury and that it is contrary to public policy to award damages for the birth or rearing of a healthy child.

- **Birth of healthy child; claim for damages prohibited.**
  
  No person may maintain a claim for relief or receive an award for damages based on the claim that the birth and rearing of a healthy child resulted in damages to him. A person may maintain a claim for relief based on a failed sterilization procedure resulting in the birth of a healthy child and receive an award of damages for the hospital and medical expenses incurred for the sterilization procedures and pregnancy, the pain and suffering connected with the pregnancy and the loss of earnings by the mother during pregnancy.

- **Birth of unhealthy child; damages limited.**
  
  Damages for the birth of an unhealthy child born as the result of professional negligence shall be limited to damages associated with the disease, defect or handicap suffered by the child.
C. **Other causes of action.**

   This section shall not preclude causes of action based on claims that, but for a wrongful act or omission, maternal death or injury would not have occurred or handicap, disease, defect or deficiency of an individual prior to birth would have been prevented, cured or ameliorated in a manner that preserved the health and life of the affected individual.

   **24 MRSA §2931**

(97) **Wrongful Death**

   Maine law recognizes a cause of action for wrongful death. Damages for the loss of comfort, society and companionship is limited to $250,000, increased from $75,000. The cap for non-economic damages has been increased from $400,000 to $500,000.

   A. Whenever the death of a person shall be caused by a wrongful act, neglect or default, and the act, neglect or default is such as would, if death had not ensued, have entitled the party injured to maintain an action and recover damages in respect thereof, then the person or the corporation that would have been liable if death had not ensued shall be liable for damages as provided in this section, notwithstanding the death of the person injured and although the death shall have been caused under such circumstances as shall amount to a felony.

   B. Every wrongful death action must be brought by and in the name of the personal representative of the deceased person. The amount recovered in every wrongful death action, except as otherwise provided, is for the exclusive benefit of the surviving spouse if no minor children of the children if no surviving spouse, one-half for the exclusive benefit of the surviving spouse and one-half for the exclusive benefit of the minor children to be divided equally among them if there are both surviving spouse and minor children, and to the deceased's heirs to be distributed as provided in section 2-106 if there is neither surviving spouse nor minor children. The jury may give damages as it determines a fair and just compensation with reference to the pecuniary injuries resulting from the and in addition shall give such damages that will compensate the estate of the deceased person for reasonable expenses of medical, surgical and hospital care and treatment and for reasonable funeral expenses. In addition, the jury may give damages not exceeding $500,000 for the loss of comfort, society and companionship of the deceased, including any damages for emotional distress arising from the same facts as those constituting the underlying claim, to the persons for whose benefit the action is brought. The jury may also give punitive damages not exceeding $250,000. An action under this section must be commenced within 2 years after the decedent's death. If a claim under this section is settled without an action having been commenced, the amount paid in settlement must be distributed as provided in this subsection. A settlement on behalf of minor children is not valid unless approved by the court, as provided in Title 14, section 1605.

   C. Whenever death ensues following a period of conscious suffering, as a result of personal injuries due to the wrongful act, neglect or default of any person, the person who caused the personal injuries resulting in such conscious suffering and death shall, in addition to the action at common law and damages recoverable therein, be liable in damages in a separate count in the same action for such death, brought, commenced and determined and subject to the same limitation as to the amount recoverable for such death and exclusively for the beneficiaries in the manner set forth in subsection (b), separately found, but in such cases there shall be only one recovery for the same injury.

   D. Any action under this section brought against a governmental entity under Title 14, sections 8101 to 8118, shall be limited as provided in those sections.

   **18-A MRSA §2-804**

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