VERMONT GUIDE TO HEALTH CARE LAW

Published by The Vermont Medical Society
February Edition
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Vermont Guide to Health Care Law
INTRODUCTION

The Vermont Medical Society is proud to release this February 2017 edition of the online Vermont Guide to Health Care Law. Work on the Guide was supported by many of the leading law firms and attorneys around the state. It is designed to give physicians and health care facilities a fundamental understanding of legal and regulatory requirements that affect the delivery of health care in Vermont today.

This guide is organized so that users may readily find the information that they seek. Topics are organized in alphabetical order. Many chapters are organized in a question-and-answer format to present material in a manner that directly responds to frequently asked questions.

We are very grateful to the attorneys and other experts whose research, writing, and editing made this guide possible. Despite their busy schedules, they devoted many, many volunteer hours to this project. Their dedication, knowledge and experience made the guide the high quality resource that it is. We couldn’t have done it without them. Thank you also to the authors and editors of prior editions of the Guide who gave us the foundation from which to work, especially Madeleine Mongan of the Vermont Medical Society and Tracy Bach with Vermont Law School, who led the effort to create the prior edition.

Please send your comments about the guide to Jessa Barnard at the Vermont Medical Society, jbarnard@vtmd.org.

DISCLAIMER

Please note that although information contained in the Guide relates to legal issues, this information is intended to be used for informational and educational purposes only. The information in the Guide is not to be used as legal advice. Persons seeking legal advice should consult a lawyer.

The Guide sponsor, authors and editors intend to provide information that is accurate and useful to health care professionals and facilities. Information about legal issues, however, may become outdated and not all information could be included in the Guide. The Guide sponsor, authors and editors disclaim liability for any consequences related to any decisions or results, arising out of or based on, any use or interpretation of information contained in or omitted from the Vermont Guide to Health Care Law.
VERMONT GUIDE TO HEALTH CARE LAW

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BUSINESS ISSUES

Topics Covered in this Chapter:
Structure/Choice of Entity
Sole Proprietorships
Corporations
Partnerships
Limited Liability Companies
Corporate and LLC Formalities
Partner/Shareholder/Member Agreements
Opening A Medical Practice
Closing A Medical Practice
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This chapter discusses the structure of various business entities and other issues to be considered in opening and closing a medical practice.

STRUCTURE/CHOICE OF ENTITY

One of the first decisions to be made as a business owner is how the business should be structured. The decision should be made only after consulting with an accountant and an attorney. There are several choices for creating an entity through which to do business. The most widely used entities for physicians and other health care professionals are professional corporations, partnerships, limited liability partnerships, or professional limited liability companies. Many factors should be considered in determining which entity is the most well suited for health care practitioners.

Regardless of the type of business entity, a physician or health care professional is personally liable for negligence or wrongful conduct while rendering professional services. A professional cannot shield himself from professional liability by creating an entity. However, creating an entity may protect a physician from:

- general business liabilities (e.g. patient or employee accidents);
- liabilities to unsecured business creditors; and
- personal liability for another practitioner’s professional malpractice

Note that most entities are created by making a filing with the Vermont Secretary of State. These filings are now made online at www.sec.state.vt.us. This website contains helpful information and a list of all fees and forms. You can also search for name availability on the website.

SOLE PROPRIETORSHIPS

A sole proprietorship is a business owned and operated by one person. The business and the individual proprietor are one and the same. Sole proprietors own all the assets of the business
and the profits generated by it. They also assume full responsibility for all of the business liabilities and debts. For this reason, most physicians form an entity through which to operate a medical practice. There are no filings required or contracts required to create a sole proprietorship. However, if the sole proprietor uses a business name (a “d/b/a”), he/she should register the business name as a tradename with the Vermont Secretary of State. A tradename registration costs $50.00 and gives the sole proprietor the right to use the registered business name for five years (and it may be renewed again and again).

**CORPORATIONS**

*What is a corporation?*
A corporation is a separate legal person under the law. A Vermont corporation is created under the laws of the state of Vermont. The owners of a corporation are its shareholders. The shareholders elect a board of directors to oversee major policies and decisions. The corporation does not dissolve when its shareholders change. The greatest advantage to operating a business as a corporation is that the corporation, and not its shareholders, officers or directors, is liable for the debts and obligations of the corporation. While there are exceptions to this rule, generally the owners of the corporation cannot be held personally liable for corporate obligations. This is the fundamental difference between operating as a corporation as compared to a sole proprietorship.

*How is a corporation created?*
A corporation must be created under state law. This is done by filing Articles of Incorporation with the Vermont Secretary of State’s office. This is now an online process. You will need a corporate name, physical and mailing address, registered agent, and number of board members in order to complete the Articles of Incorporation. When the Articles of Incorporation are received and accepted by the Vermont Secretary of State, it will issue a Certificate of Incorporation.


*What is a professional corporation?*
A professional corporation is an entity allowed to be used by certain licensed professionals. The key to a professional corporation is that the professionals get the benefit of the corporate form for the business aspects of the practice, but do not get protection from liability for damages caused by the provision of the professional service. This means that if a physician commits malpractice, that physician can be held personally liable even though he/she operates as a professional corporation.

Vermont’s professional corporation statute was substantially revised in July 2002. (11 V.S.A. §§ 815-881). This professional corporation law does the following:
- broadens the ownership of a professional corporation and allows various different types of professionals to incorporate together;
- permits existing professional corporations to remain under the old statute or elect to be governed by the new law;
- authorizes any licensing authority to restrict the professional corporation’s behavior in the interest of public protection; and
• sets out general procedures for the acquisition and disposition of the shares of stock of a professional corporation shareholder.

Are there ongoing administrative requirements of operating the corporate form?
Yes, corporations have ongoing administrative requirements, including filing documents with the Vermont Secretary of State’s office, the federal Internal Revenue Service, the Vermont Department of Taxes, and the Vermont Department of Unemployment and Training. Every for-profit corporation must file an annual report with the Vermont Secretary of State’s office. Every corporation must hold an annual meeting of stockholders and directors, and must document these meetings in its corporate record book. Every for-profit corporation must issue shares of stock to its shareholders and document the consideration received for the shares of stock.

What is a C corporation?
A “C” corporation is a standard business corporation. A “C” corporation’s earnings are taxed at the corporate level and income to the shareholders is taxed at the personal income tax level.

What is an S corporation?
An “S” corporation is a small business corporation that elects to pass corporate income, losses, deductions and credits through to its shareholders under the Internal Revenue Code. The profit and loss of an S corporation normally passes through to the shareholders in proportion to their shares in the corporation. The shareholders report the profit or loss on their individual tax returns. A professional corporation may elect to be treated as an S corporation or may remain as a C corporation. An S election is made by filing IRS Form 2553 with the IRS. An S corporation has certain limitations including a limit on the number of shareholders (100), a requirement that shareholders be U.S. citizens or resident aliens, and a restriction to only one type of stock. All eligible shareholders must be individuals, estates, certain defined trusts, or certain tax-exempt organizations. Internal Revenue Code section 1361 sets forth complete rules relating to S corporations.

To Do List for Forming a Corporation
Below is a list of tasks to be done when creating a corporation:

Lawyer
1. Draft Articles of Incorporation for review by client
2. File Articles with Vermont Secretary of State
3. Draft Bylaws
4. Prepare Organizational Board Meeting Resolutions
5. Issue Stock Certificates to Shareholders

Accountant
1. Obtain Taxpayer Id Number (EIN) from IRS
2. Discuss with client whether to file S election with IRS
3. Obtain all required business tax account numbers from Vermont Department of Taxes

Medical Practice
1. Open bank accounts
2. Hold a Board of Director organizational meeting and elect officers
3. Obtain insurance
4. If the corporation has employees, obtain workmen’s compensation coverage, unemployment compensation insurance
5. If the corporation has employees, post all labor/employee posters

**PARTNERSHIPS**

A partnership is an association of two or more persons to carry on as co-owners of a business for profit. Vermont’s law provides basic definitions and rules about partnerships some of which can be varied by a partnership agreement. For income tax purposes, a partnership is generally treated as a pass-through entity. The partnership itself does not pay income taxes. The partners report the business profits or losses on their personal income tax returns.

**What is a general partnership and how is it created?**
A general partnership is an association where each owner has unlimited liability. Many partnerships are created without a written agreement. There are many advantages to using a written agreement, including the certainty the partners achieve about their business relationship. If there is no partnership agreement, then the rights and duties of the partners are controlled by Vermont law (See, Title 11, Vermont Statutes Annotated, Chapter 22, 11 V.S.A. § 3201 et seq.).

**What is a limited partnership?**
A limited partnership is a partnership formed by two or more persons having one or more general partners and one or more limited partners. A general partner is personally liable for the debts and obligations of the partnership. A limited partner may not participate in the control of the business. Typically, a limited partner is not liable for the obligations of a limited partnership (unless he or she is also a general partner or unless he/she participates in the control of the business). A limited partnership is formed by filing a Certificate of Limited Partnership in the office of the Vermont Secretary of State. The Certificate must contain the following information: the office and the address of the registered agent; the name and business address of each general partner; the name and residential address of the partners, and amount of cash and description of agreed value of other property contributed by each limited partner; the latest date upon which the limited partnership is to dissolve.

**What is a limited liability partnership?**
A limited liability partnership is a partnership that voluntarily registers with the Vermont Secretary of State as an LLP. Vermont law specifically allows any partnership to register as an LLP. The LLP form shields the individual partners from personal liability for partnership debts. The partner may lose all he/she invested (his partnership interest) but the partner’s personal assets cannot be used to satisfy the debts of the LLP. Thus, a partner in an LLP has similar protection as a stockholder in a corporation. To create an LLP, the partnership must file an LLP registration form with the Vermont Secretary of State. An LLP must end its name with the words “Registered Limited Liability Partnership,” “Limited Liability Partnership,” “RLLP” or “LLP.” See, 11 V.S.A. § 3291 et seq.

**LIMITED LIABILITY COMPANIES**
What is a limited liability company?
A limited liability company (“LLC”) is an entity formed under Vermont law that has elements of both a corporation and a partnership. An LLC is treated like a corporation for purposes of limited liability, and as a partnership (if properly structured) for purposes of income taxation. In order to form an LLC, one must file Articles of Organization with the Vermont Secretary of State. The name of a limited liability company must contain the words “limited liability company” or “limited company” or the abbreviation “LLC” or “LC.” See, 11 V.S.A. § 3001 et seq.

Can one physician form an LLC?
Yes, in Vermont an LLC may be created by one person and that person shall be the sole member (owner). The LLC will protect the sole member from liability for obligations of the LLC.

What is an Operating Agreement?
An Operating Agreement is a private document that sets forth the governance rules for the LLC. The Operating Agreement does not get filed with the Vermont Secretary of State. Many of the provisions of the Vermont LLC statute can be altered or varied by agreement among the LLC members in the Operating Agreement. The Operating Agreement should specify whether the LLC will be managed by its members or by one or more managers. It should also address the following topics:

- how a new member is added to the LLC;
- how the members vote;
- how a member can withdraw or be forced out of the LLC; and
- who makes tax and accounting decisions for the LLC.

Can licensed professionals create an LLC?
Yes, licensed professionals can form a professional limited liability company; however, professionals are also governed by the licensing laws and by Vermont’s professional corporation act. Section 3012(c) of the Limited Liability Company Act (11 V.S.A. § 3012(c)) states that a “limited liability company shall engage in rendering professional services only to the extent that, and subject to the conditions and limitations under which, a professional corporation may engage in rendering professional services.” The name of a professional limited liability company must contain the words “professional limited liability company” or the abbreviation “PLC.” The members of a PLC shall be treated in the same manner as shareholders of a professional corporation. The managers of a PLC shall be treated in the same manner as directors of a professional corporation.

How will an LLC be taxed?
An LLC can elect to be taxed as a corporation or as a pass-through entity. If the LLC has a single member and elects pass-through treatment, it will be treated as a sole proprietorship. If the LLC has more than one member and elects pass-through treatment, it will be treated as a partnership.

CORPORATE AND LLC FORMALITIES
There are certain formalities that a corporation and LLC should comply with in order for it to maintain its status as a separate legal entity, and ensure that the members and shareholders have limited liability to the extent allowed by law, as follows:

1. Maintain a separate bank account and financial records for the entity.
2. Execute the governing documents (Bylaws or Operating Agreement).
3. Obtain all necessary business tax account numbers from the Vermont Department of Taxes.
4. File an annual report with Vermont Secretary of State.
5. Always represent yourself as a duly authorized agent of the entity so that third parties know that they are dealing with an entity and not a sole proprietorship or general partnership.
6. Obtain all-risk insurance, including liability insurance, to provide coverage as appropriate.
7. Do not add a member to the LLC or a shareholder to the corporation without consulting with legal counsel.
8. Work with your accountant to ensure that all ongoing necessary state and federal tax filings get made.

**PARTNER/SHAREHOLDER/MEMBER AGREEMENTS**

Regardless of the type of entity used, if the entity is owned by more than one physician, the physicians should enter into an agreement governing their relationship. If the physicians are doing business as a corporation, they should enter a shareholder agreement. If the physicians are doing business as a partnership, they should enter a partnership agreement. If the physicians are doing business as a limited liability company, they should enter an operating agreement. These agreements should document the following: how decisions will be made; how profits will be shared; how disputes will be resolved; how future owners will be admitted to the entity; how owners will be bought out; and, what happens upon the death or disability of an owner.

**OPENING A MEDICAL PRACTICE**

*What issues should be considered in opening a practice?*

The first issue is the choice of entity issue discussed above. Once the type of entity is selected, the physicians involved should agree on the terms of the appropriate governing agreements. The physicians should contact the local hospital for information about social supports such as the local community mental health agencies and the local agencies on aging. The physicians should determine what medical services the practice will offer, such as lab or x-ray.

Before opening a practice consider the following:

- Create a business plan and pro forma financial statements
- Review choice of entity
- Review licensing requirements
- Look for office space. You may need a lawyer to review a commercial lease for you. Or you may choose to purchase property. In either case you will need a medical office design.
- Billing
- Medical malpractice insurance
- Obtaining provider numbers for Medicaid, Medicare and health insurance companies
- Technology and information system needs
- Policies and procedures manual

These topics are discussed below.

Licensing
The physician must ensure that he/she has all required professional licenses. Licensing in Vermont is obtained from the Vermont Department of Health, Vermont Department of Medical Practice. The website (http://healthvermont.gov/systems/medical-practice-board) has an e-licensing function. Health care professionals who are licensed or certified by the Board can apply for and renew a license or certification, check the status of an application, or update profile information.

Office Space/Equipment/Other Issues
The physician will need to obtain office space for the practice, which involves leasing or purchasing property. The type of service being offered by the physician will often determine the type of space and equipment the physician will need. The physicians need to determine what information and communication systems will be used by the practice. The physician practice will need to obtain insurance to protect against various risks, including all risk insurance protecting against damage to the premises and equipment, general liability insurance, and professional liability insurance. The physician should consider advertising the practice. The physician should determine whether the practice will need to obtain the services of a billing company.

Employees
If the physician hires staff for the office, the physician practice will need to comply with state and federal laws regarding employees and will need to adopt certain employment policies. (See Chapter on Employment Law).

If the physician has previously worked for another practice or medical center, the physician should review his/her employment contract and/or personnel policies to determine whether there are restrictive covenants that prevent the physician from practicing in certain geographical areas. The physician may need to obtain medical records of patients from the patients’ former providers. The physician should prepare a consent form that patients can sign.

What resources are available to provide information on opening a practice?
The physician should check various websites for tips on opening a practice and sample forms and contracts, including the websites for the American Medical Association and the American Academy of Family Physicians.

CLOSING A MEDICAL PRACTICE

The physician must plan ahead before closing a medical practice. The physician must address the following issues in closing the practice: notification to patients; notification to employees;
notification to malpractice and general liability insurers; cancellation of contracts with practice management companies or billing companies; termination of equipment leases; disposal or storage of patient records; and disposal of drug samples. Additionally, if the medical practice is organized as an entity such as a corporation or LLC, the physician must wind up and dissolve the business entity after the closure of the practice.

If the physician intends to sell the medical practice, instead of closing it, then the physician will need to have the practice valued and marketed. The physician will need to hire an accountant and attorney for help in structuring the sale and in negotiating the terms of the sale.

Further guidance on closing a practice can be found from your state or national medical societies; also see the Maine Medical Association Physician’s Guide to Closing a Medical Practice (noting that this reflects some Maine state-specific laws).

**How should a physician notify patients?**

A physician is generally under an ethical and legal obligation to provide services to a patient as long as the patient needs them. In order to avoid a claim of abandonment, the physician should take several steps to terminate the physician-patient relationship. (Abandonment is defined as the termination of the physician patient relationship at an unreasonable time and without giving the patient the chance to find an appropriate replacement.)

The physician should initially notify all patients by a letter which informs the patient of the date the physician will stop practicing and the method by which patients can obtain their medical records or have them transferred to another physician. Ideally, the letter would be sent by certified mail, return receipt requested, but the cost may be prohibitive. The patient notification should include: a brief explanation of the reason for terminating the patient relationship; agreement to continue to provide medical treatment for a reasonable period such as 30 days to allow the patient to find another physician; and, a consent form which the patient can sign in order to authorize the transfer of his or her medical records to the new physician.

The physician should make a list of patients to whom the notice should be sent. Identifying patients could be done by running a list of new patients and patients whom visited the practice during the prior two or three years. The physician should not send a notice to patients of other physicians that he or she shared call coverage with, or patients whom were seen through a hospital consult.

The Vermont Board of Medical Practice website contains an Advisory (from 1999) about the requirement of notifying patients of the termination of the physician-patient relationship. See http://healthvermont.gov/sites/default/files/documents/2016/12/BMP_Policies_Termination%20of%20Relationship.pdf. For more information, see the Guide Sections on Initiation and Termination of the Physician-Patient Relationship and Risk Management.

**Should the physician refer patients to another physician?**

If at all possible, the physician closing a medical practice should avoid referring patients to another named physician. Such a referral puts the referring physician at risk of a claim for negligent referral. The physician should instead refer their patients to general resources such as
the Board of Medical Practice, their local hospital or insurance carrier. Additionally, patients who are under the physician’s ongoing care may require additional arrangements.

Are there employee notice requirements upon the closure of a physician practice?
In Vermont, there are no statutory requirements for notifying employees in a particular manner. The physician should review any employment contracts between the physician practice and its employees. The physician should also review any existing personnel policies or employee handbooks to determine the rights of employees. The physician practice must provide written notice of the termination date of benefits. The physician practice may have certain obligations under COBRA.

Is there a risk of notifying employees about a planned practice closure too early?
Yes, when a physician notifies his/her employees that the practice will be closing, there is a risk that the employees may immediately search for a new job and leave the physician’s practice prior to the closure date. To avoid this, and provide an incentive for employees to remain at the practice, the physician may want to offer a bonus to any employee who works at the practice until the closing date.

How should the physician handle patient records upon the closure of the practice?
Upon the request of the patient, the physician would send a copy of the patient’s records to a new physician or to the patient. In either case, the physician should retain the original (or retain access to the original). The physician should store the original records in a safe place, such as a commercial storage firm. The physician should ask the storage company to enter into a confidentiality agreement so that the physician meets his/her legal obligation to keep patient records confidential. For more information on the length of time to retain records, see the section on Medical Records.

How should the physician dispose of drug samples?
Hopefully your practice already monitors drug samples, organizes them, checks expiration dates, maintains distribution papers in case there is a recall, and keeps the medicines physically secured.

When a practice closes, it may still have possession of unused drugs or drug samples. The physician practice should return any unopened bottles to the distributor or manufacturer to obtain a refund if possible. The physician practice may also return drug samples to the drug company representatives who visit the practice. The physician may also check with the local hospital or local pharmacist for suggestions on how to dispose of the drug samples. There are also comprehensive medical waste disposal companies that can be hired to provide technical advice and disposal services.

ABOUT THE AUTHOR

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CONSENT, PRIVACY AND MEDICAL RECORDS

Topics Covered in this Chapter:
Consent for Treatment
Minor Consent
Privacy and Medical Information
Medical Records
About the Authors

By: Anne Cramer, Shireen Hart and Lauren Layman
Primmer Piper Eggleston & Cramer PC

CONSENT FOR TREATMENT

What are the legal principles supporting a patient’s right to informed consent?
Our common law in the United States has long recognized that every human being of adult years and sound mind has the right to determine what shall be done with his or her body. Initially, this right was defined in terms of an individual’s right to be free of unwanted bodily invasion—essentially that a person cannot be “touched” without his or her explicit permission. Being “touched” without permission from the individual constitutes a battery. In a health care context, a person who has not given permission, or consent, to a “touching” by a health care provider for medical treatment is considered to have a cause of action against the health care provider for a battery.

What is a battery?
A battery is an intentional act that results in harmful contact with another. In a health care context, a health care provider commits a battery if the provider performs a procedure for which the patient has not given consent. A defense to a claim of battery is that the patient consented to the contact.

What is the current law in Vermont regarding the principle of informed consent?
The duty of a health care provider to obtain consent from a patient for treatment has been defined by statute in most states, including Vermont. The Vermont statutes define this duty by limiting the scope of a medical malpractice action based on lack of informed consent, and by outlining a hospital patient’s rights to information and consent in the Hospital Patient Bill of Rights. 12 V.S.A. §1909, 18 V.S.A. §1852(3) and (4).

Medical Malpractice – A Lack of Informed Consent
The Vermont medical malpractice statute defines “informed consent” in the negative (i.e., the consequences of a lack of informed consent). Under the statute, a lack of informed consent is defined as the following:

1. The failure of the person providing professional treatment or diagnosis to disclose to the patient such alternatives thereto and the reasonably foreseeable risks and benefits
involved as a reasonable medical practitioner under similar circumstances would have disclosed in a manner permitting the patient to make a knowledgeable evaluation; or
2. The failure to provide a reasonable answer to any specific question about foreseeable risks and benefits. Additionally, a medical practitioner has a duty to not withhold any information requested by the patient.

12 VSA § 1909(a) and (d).

Patient Bill of Rights – Defining Informed Consent

Under Vermont law, hospital patients have a right to be provided with the necessary information to provide informed consent. The Vermont Hospital Patient Bill of Rights provides the patient the affirmative “right to obtain from the physician coordinating his or her care, complete and current information concerning diagnosis, treatment and any known prognosis in terms the patient can reasonably be expected to understand.” The patient has the right, except in emergencies, to receive from the physician information necessary to give informed consent prior to the start of any procedure or treatment. Such information should include the medically significant risks involved with this procedure or treatment, the probable duration of incapacity and any medically significant alternatives. 18 V.S.A. §1852(3) and (4).

Note that the rights are also set forth in the Medicare Hospital Conditions for Participation. 42 CFR §482.13.

What is the difference between an allegation of battery where the patient has provided no consent to the treatment or procedure, and an allegation of medical malpractice where the patient argues the provider failed to meet the standards of informed consent?

A battery entails a lack of consent. Where a medical professional performs a treatment or procedure for which there is no consent, the patient has a cause of action for battery – an intentional contact or invasion which causes harm.

An action for medical malpractice requires a lack of informed consent. Where the patient has provided consent for the treatment or procedure but receives inadequate disclosure of the alternatives and foreseeable risks and benefits of the alternatives, the cause of action and liability is based on a lack of informed consent and the patient may have a claim of medical malpractice against the health care provider for failing to provide the necessary disclosures. Christman v. Davis, 179 Vt. 99, 101-102 (2005).

Must a medical professional obtain informed consent in an emergency?

No, where treatment has been provided to address an emergency, Vermont law specifies that there can be no cause of action for medical malpractice based on a lack of informed consent. 12 V.S.A. §1909(b); see 18 V.S.A. §1852(4).

Under what other conditions is it permissible for a health care provider to have treated a patient without first having obtained informed consent? What are the exceptions to the principle of informed consent?

If any of the following are present, the health care provider can treat the patient without first obtaining informed consent:
1. the risk not disclosed is too commonly known to require disclosure and that the risk is not substantial,
2. the patient has assured the health care provider he would undergo the treatment procedure or procedures regardless of the risk involved,
3. the patient indicated to the medical practitioner that he/she did not want to be informed of the matters to which he/she would be entitled to be informed;
4. consent either by the patient or on behalf of the patient was not reasonably possible (e.g., as a result of the patient’s incapacity and the unavailability of any patient representative),
5. a reasonably prudent person in the patient’s position would have undergone the treatment or procedure if he had not been fully informed.

12 V.S.A. §1909(c). Further, in the case of an emergency, no informed consent is necessary.

Any discussion about informed consent, or exceptions to informed consent should be documented in the medical record including the reasons for any exception. An executed patient consent form should be maintained in the record or a reason noted as to why a form was not signed.

Must a patient provide consent for a blood draw for a law enforcement purpose?
A patient’s broad consent for medical treatment is distinct from consent for a blood draw for non-medical, law enforcement purposes. O’Brien v. Synnott, 193 Vt. 546, 553 (2013). A health care provider may be liable for failure to obtain informed consent if he or she draws blood for law enforcement purposes from a conscious patient who has refused requests from law enforcement to submit to a blood alcohol test. Id. This is the case even if the health care provider is acting in reliance on an order from law enforcement. Id. The health care provider would not be subject to liability for battery if law enforcement provided a warrant for the blood draw. Id.

Must a patient be advised of the foreseeable risks and benefits of a treatment or procedure in the event that the health care provider believes that this information would adversely affect the patient’s condition?
Yes, Vermont no longer recognizes a “limited” therapeutic privilege that would permit a health care provider to withhold information from a patient if the provider believes that the information would adversely affect the patient’s condition. Prior statutory references to such a privilege have been deleted from Vermont’s statutes.

What type of evidence must be produced by a patient at trial to support a charge of medical malpractice as a result of lack of informed consent?
Vermont law requires that a patient establish the standard of care for the treatment or procedure that was not met or for which insufficient information was provided to give informed consent. 12 V.S.A. § 1908. The law places an affirmative burden on the patient to establish a claim of medical malpractice as a result of lack of informed consent using expert testimony. Christman v. Davis, 179 Vt. 99, 102 (2005), see also 12 V.S.A. § 1909. Courts have interpreted this law to mean that generally the standard of care can only be proved by expert testimony. Lockwood v. Lord, 163 Vt. 210, 213 (1994), see also Larson v. Candlish, 144 Vt. 499 (1984). A limited exception to this rule has been carved out where no expert testimony is necessary to establish the standard of care if the alleged violation of the standard of care is so apparent that it can be

**If a patient gives consent to a treatment or procedure after being fully informed of the risks, benefits and alternatives, and the health care provider performs a treatment or procedure with similar risks, benefits and alternatives, but not the same treatment or procedure, does the patient have a cause of action for battery? Does the patient have a cause of action for malpractice due to a lack of informed consent?**

A patient’s consent is effective for a particular procedure or treatment, or for “substantially the same” proposed conduct. Whether consent is effective for a different treatment or procedure with similar risks and benefits depends upon whether the different conduct or procedure that was performed was “within the bounds” of the conduct for which consent was obtained. *Christman v. Davis*, 179 Vt. 99, 105 (2005). In *Christman v. Davis*, the Vermont Supreme Court held that a “less-extensive operation than discussed with the patient” (i.e., a flap procedure was conducted when the patient had consented to a tissue graft to address a dental root issue) was within the bounds of the patient’s original consent and, thus, there could be no cause of action for battery. *Id.*

**Must a patient’s consent be provided in writing to qualify as informed consent?**

There is no specific requirement that informed consent be given in writing or acknowledged in writing. However, given the time delays attendant to any challenge to a medical procedure for lack of informed consent, obtaining written acknowledgement is prudent and is considered the accepted practice. Additionally, for professionals treating patients in a hospital or ambulatory care setting or performing surgery, properly executed informed consent forms must be included in a patient’s record to meet the Medicare Conditions of Participation. See 42 C.F.R. §§ 482.13, 482.24, and 482.51. Additionally, for Joint Commission Accreditation, hospitals must be able to demonstrate that there are established policies and processes for obtaining informed consent and provide evidence that such consents are being obtained. See Joint Commission Standard PFR.5.1.-5.4.

**Who is responsible for securing informed consent?**

The health care provider who will be providing treatment has the duty to secure the patient’s informed consent. Although the health care provider need not perform this task personally, he or she is the one who faces liability if a patient is not properly informed or does not provide consent. Regardless of which health care provider secures the consent, a patient has a right under the Vermont Bill of Rights for hospital patients to obtain information regarding his or her diagnosis, treatment and prognosis from the physician who is coordinating the patient’s care. 18 V.S.A. § 1852(a)(3).

**What information must the health care professional disclose to the patient before informed consent can be fairly given?**

The professional must give the patient information regarding the specific procedure and treatment, the probable duration of incapacitation, the name(s) of the person responsible for performing the procedures or treatment, and the reasonably foreseeable risks and benefits of the proposed treatment or procedure. 18 V.S.A. § 1852(a)(4). Where medically significant
alternatives for care or treatment exist, or when the patient requests information concerning medical alternatives, the patient has the right to such information. *Id.*

**What risks, benefits and alternatives must be disclosed for a patient to provide informed consent?**

The treating professional must disclose any “reasonably foreseeable risks and benefits that a reasonable medical practitioner under similar circumstances would have disclosed.” 12 V.S.A. § 1909(a)(1). Further, a patient is entitled to a reasonable answer to any specific questions about foreseeable risks and benefits, and a medical practitioner shall not withhold any requested information. 12 V.S.A. § 1909(d). Essentially, the medical professional must provide the patient with sufficient information to permit the patient to make a knowledgeable evaluation of the treatment or procedure. Additionally, the treating professional must disclose medically significant alternatives for care or treatment and provide a patient with information regarding medical alternatives if the patient requests such information. 18 V.S.A. § 1852(a)(4).

**How must the information be disclosed to a patient?**

A face-to-face explanation is advisable although, not necessarily required. Providing the patient with written, digital or visual information about the risks, benefits and alternatives may also satisfy the standard in many instances. However, any communication must be meaningful such that the patient needs to be able to read or understand the language and vocabulary utilized. Additionally, although informed consent is a process rather than a form, documenting the process on a form is important to demonstrate the information provided by the professional to the patient, the questions asked by the patient, and evidence of the patient’s signature at the time such information is given.

**For what treatments or procedures is informed consent required?**

Vermont law does not generally designate the types of treatment, procedure or surgeries which require the professional to obtain a patient’s informed consent before acting. Rather, the law provides for a cause of action for failing to obtain informed consent. See 12 VSA § 1909(a) and (d). However, a defense to a claim of medical malpractice for lack of informed consent may be based on allegations that the risk that was not disclosed is so commonly known and insubstantial that informed consent was not necessary. 12 V.S.A. § 1909(c)(1). Thus, informed consent is not necessary in such circumstances.

Vermont law also requires informed consent for the prescription of opioid medications in certain circumstances. VT Rules Governing the Prescribing of Opioids for Pain, *effective July, 2017.*

**What are the requirements for informed consent for prescribing opiate medications?**

Prior to writing a prescription for an opioid Schedule II, II or IV for the first time to any patient, health care providers must do the following:

- Have an in-person discussion with the patient (or a parent, guardian or legal representative if the patient is a minor or lacks legal competence) regarding the potential side effects, risks of dependence and overdose, alternative treatments, appropriate tapering, and safe storage and disposal of opioid medications;
- Provide the patient with a Vermont Department of Health patient education sheet which will be available on the Department’s website; and
• Receive, and include in the patient's medical record, a signed informed consent from the patient (or from the patient's parent, guardian or legal representative if the patient is a minor or lacks the capacity to provide informed consent) that includes information regarding the drug's potential for misuse, abuse, diversion, and addiction; potential side effects; tolerance; the risks associated with the drug for life-threatening respiratory depression; potentially fatal overdose as a result of accidental exposure, especially in children; neonatal opioid withdrawal syndrome; and potentially fatal overdose when combining with alcohol and/or other psychoactive medication including but not limited to benzodiazepines and barbiturates.

VT Rules Governing the Prescribing of Opioids for Pain, Section 4.3, effective July, 2017.

Chronic Pain
In addition to the requirements described above, before providers prescribe opioids for the treatment of chronic pain (i.e., pain lasting longer than 90 days), providers must also receive and include in the patient's medical record a signed Controlled Substance Treatment Agreement from the patient, or, if the patient lacks the capacity to provide informed consent, from the patient's legal representative. This agreement must include functional goals for treatment, dispensing pharmacy choice, and safe storage and disposal of medication. The agreement must also include other requirements as determined by the provider, such as directly observed urine drug testing and pill counts to reasonably and timely inform the provider if the patient is misusing the prescribed substance. The Controlled Substance Treatment Agreement must be reviewed by the provider and the patient no less than every 365 days to reevaluate the patient.

The provider must also ask, and document asking, the patient (or the parent, guardian or legal representative as necessary) if the patient currently, or has recently, been dispensed methadone or buprenorphine or been prescribed or taken any other controlled substance. The provider must explain to the patient, and document the explanation, that this information is important for the patient’s safety and that the patient is required by law to disclose this information. VT Rules Governing the Prescribing of Opioids for Chronic Pain, Section 5.1.4, through June, 2017; VT Rules Governing the Prescribing of Opioids for Pain, Section 6.2.1.4, effective July, 2017; and 18 V.S.A. § 4223.

Prior to prescribing a Morphine Milligram Equivalent (“MME”) Daily Dose of 90 (a calculator for MME can be found on the Department of Health’s website) for chronic pain, the provider must have an in-person discussion with the patient (or a parent, guardian or legal representative as necessary) regarding the increased risk of fatal and non-fatal overdose and any precautions the patient should take. The informed consent and the Controlled Substance Treatment Agreement must be revised to reflect any changes in the prescription (e.g., the pill count changes, directly observed urine drug testing is required).

These requirements regarding the prescription of opioids for the treatment of chronic pain do not apply to the treatment of patients with chronic pain associated with cancer or cancer treatment, palliative care, end-of-life and hospice care, or patients in skilled and intermediate care nursing facilities.

VT Rules Governing the Prescribing of Opioids for Chronic Pain, Section 5.0, through June, 2017
VT Rules Governing the Prescribing of Opioids for Pain, Section 6.0, effective July, 2017.

Extended Release Hydrocodones and Oxycodones without Abuse-Deterrent Opioid Formulations
In addition to the other requirements for the prescription of opioids, including those pertaining to prescriptions for chronic pain, prior to prescribing extended release Hydrocodones and Oxycodones that are not abuse-deterrent opioids, a provider must include in the Controlled Substance Treatment Agreement a requirement that the patient (or a parent, guardian or legal representative as necessary) agrees to urine testing no less frequently than annually with the actual frequency determined by the provider.

Providers prescribing these medications must have follow-up visits with and evaluations of the patient no less frequently than every 90 days and, during these follow-up appointments, must document that the provider has explained and received acknowledgement from the patient (or a parent, guardian or legal representative as necessary) that a violation of the Controlled Substance Treatment Agreement will result in a re-assessment of the patient’s treatment plan and alteration or institution of controls over medication prescribing and dispensing, which may include tapering or discontinuing the prescription.

VT Rules Governing the Prescribing of Opioids for Chronic Pain, Section 8.0, through June, 2017
VT Rules Governing the Prescribing of Opioids for Pain, Section 8.0, effective July, 2017

When is a health care professional required to discuss the impact a medical condition or medication will have on a patient’s ability to safely operate a vehicle?
In accordance with National Transportation Safety Board recommendations directed at medical and pharmacy licensing boards, health care providers should routinely discuss with patients the effect that medical condition and medication use may have on the ability to safely operate a vehicle in any mode of transportation. NTSB Safety Recommendations I-14-1 and I-14-2.

MINOR CONSENT

What is the legal age of consent in Vermont?
Eighteen (18) is the legal age of consent in Vermont. Individuals under the age of eighteen are minors under Vermont law. In most circumstances, minors are not capable of giving informed consent to their own medical care.

Who can give informed consent to health care for a minor?
Generally, only the following individuals may give informed consent to health care for a minor:

- A guardian or representative who has been appointed by a judge to make health care decisions for the child;
- A parent (adopted or biological).

Can a minor ever give informed consent for his or her own health care?
Yes, minors who are married or have ever been married and minors on active U.S. military duty may give informed consent to their own health care. 12 V.S.A. § 7151. Minors emancipated by a court order may also give informed consent to their own health care. Id. If a minor states that he
or she has been emancipated by court order and is thus authorized to provide informed consent for a medical procedure, the health care provider should obtain a copy of the court order regarding emancipation and retain the court order in the patient’s medical record.

To become emancipated, a probate court must determine that a minor: (1) is sixteen (16) years of age; (2) has lived separate and apart from his or her parents or legal guardians for at least three (3) months; (3) is managing his or her own financial affairs; (4) has demonstrated ability to be self-sufficient in financial and personal affairs; (5) holds a high school diploma or its equivalent or is earning passing grades in an educational program approved by the court and directed toward the earning of a high school diploma or its equivalent; (6) is not under a legal guardianship or in the custody of the Commissioner for Children and Families; and (7) is not under the supervision or in the custody of the Commissioner of Corrections. Id.

Are there any special situations where an unemancipated minor may give informed consent to their own health care?

Yes, minors 12 years or older may give informed consent to treatment for sexually transmitted diseases (including HIV and AIDS), drug dependence, and alcoholism. 18 V.S.A. § 4226. But if a minor requires immediate hospitalization for treatment of any of these conditions, the parents must be notified of the hospitalization. Id.

Minors 14 years or older may also voluntarily admit themselves to a hospital for mental health related treatment if they give informed consent in writing. Minors under 14 may admit themselves to a hospital for mental health related treatment by providing their own written informed consent and a written application from a parent or guardian. 18 V.S.A. § 7503.

Minors of any age may give informed consent to medical treatment associated with rape, incest, or sexual abuse. Health care providers are required to report such incidents to the Department of Children and Families (“DCF”) within 24 hours. 33 V.S.A. § 4911 et seq.

Who can give informed consent for HPV vaccinations of an unemancipated minor?

Informed consent from a parent or guardian should be obtained before administering an HPV vaccine to a minor. Although HPV is a sexually transmitted disease (“STD”), it is unclear whether the vaccine for the HPV is considered a “treatment” for an STD. Thus, it is unlikely that the HPV vaccine falls within the Vermont statutory exception permitting minors 12 years or older to give informed consent to treatment for STDs. See “Are there any special situations where an unemancipated minor may give informed consent to their own health care?”, above, and 18 V.S.A. § 4226. As a result, as with other vaccines, health care providers should obtain informed consent from a parent or guardian before administering the HPV vaccine to minors.

Further, in addition to obtaining informed consent from a parent or legal guardian for the vaccination, health care providers must give the parents or legal representatives of a child a separate Vaccine Information Sheet (“VIS”) prior to administering every dose of the HPV vaccine. 42 U.S.C. § 300aa-26(d) (Emphasis added) see also Centers for Disease Control and Prevention, Vaccine Information Statements: Frequently Asked Questions, https://www.cdc.gov/vaccines/hcp/vis/about/vis-faqs.html (last visited January 9, 2017).
Per the federal National Childhood Vaccine Injury Act (“NCVIA”), if the parent/legal representative is present at the time of the HPV vaccination, the VIS must be provided to the parent/legal representative. If the parent/legal representative is not present at the time of the vaccination (e.g., the minor has driven him or herself to the appointment), the provider must provide the legal representative with a VIS prior to vaccination (i.e., before the minor arrives to receive the vaccination) and the VIS must be coupled with a method to acknowledge receipt/review of the VIS (e.g., adding a written statement that the parent/legal representative received and reviewed the current edition of the VIS, with the edition date specified, on the medical consent form authorizing vaccination).

**Who can give informed consent to health care when the minor’s parents are divorced?**

In general, both parents can make health care decisions for the child. If the parents are divorced, the informed consent of either parent may be assumed to be sufficient. That said, every divorce decree is different and there are situations where a judge has entered an order in which one or both of the parents is no longer authorized to make health care decisions for their child. In those situations, health care professionals are obligated to follow the judge’s order. If one parent claims that he or she has exclusive control of medical decision-making, he or she should be asked to present relevant family court documents.

**Can a parent or guardian delegate authority for giving informed consent to medical treatment for a minor?**

Vermont law is silent on this question. It is reasonable to assume that where the parent(s) or guardian(s) will not be reasonably available to give informed consent to medical treatment (e.g., vacations, illness, etc.), they may delegate this authority to a selected adult. Because there are no clear guidelines for determining the legality of an apparent delegation of this parental authority, providers should make reasonable efforts to obtain parental informed consent and should use their professional judgment and exercise greater caution in providing services involving any increased risk. If the office has a written consent on file that has been signed by the parent authorizing the health care provider and his/her staff to provide medical care to the child, there is little or no risk in providing the child with routine medical care. In situations where the treatment is non-routine or poses some degree of risk to the child, it is always advisable to consult with the parent before proceeding. Asking parents ahead of time to document authorization for medical treatment in their absence is recommended.

**When faced with a situation where the minor has been brought to the office by a grandparent or adult sibling, should a provider refuse treatment until informed consent from the parent/guardian has been obtained?**

No. As long as the health care provider has exercised due care and made a good faith determination that the grandparent or adult sibling has been authorized by the child’s parent or guardian to act as the agent in obtaining medical care for the minor, liability is unlikely if the parent later claims that the grandparent or sibling lacked authorization. That said, if the medical care will involve anything more than routine, low-risk procedures, the health care provider should make every effort to obtain the parent’s specific informed consent before providing this type of care. Health care providers are expected to use good judgment in determining which procedures require specific parental informed consent.
Is parental informed consent required before a minor can terminate a pregnancy or receive contraceptive devices/medications?
U.S. Supreme Court rulings and Vermont state law permit unemancipated minors to give informed consent to abortions and medical treatment necessary to obtain contraceptive devices and medications. In determining whether the minor’s informed consent is sufficient, health care providers should carefully assess whether the minor understands the nature and risks of the proposed treatment and is capable of making an informed, rational choice. The following factors should be considered when making such an assessment: age of the minor, ability, experience, education, maturity level, conduct and demeanor. If the provider determines that notifying the parent or guardian is in the best interests of the child, this should be discussed with the minor. It is always important to document decisions and the basis for the decision.

Is parental informed consent required before minors are provided pregnancy, prenatal and delivery care?
Vermont law is silent as to whether pregnant, unemancipated minors can give informed consent to reproductive services. Because cases involving other reproductive services, such as abortion and contraception, have allowed minors considerable freedom, health care providers should evaluate whether the minor’s informed consent is sufficient considering the following factors: age of the minor, ability, experience, education, maturity level, conduct and demeanor.

Are there any situations where providers need not obtain informed consent from anyone before providing the minor with health care?
Yes. When dealing with an emergency – a situation in which immediate treatment is needed to save the patient’s life or health – where informed consent cannot be obtained, treatment may be provided without obtaining informed consent. It is good practice to try and obtain informed consent as soon as possible even in an emergency situation. 12 V.S.A. § 1909(b).

If adolescents drive themselves to their appointments, should providers obtain parental informed consent before treating the minor?
In such a situation, the health care provider is expected to exercise sound judgment as to whether the parent must be contacted before treating an unemancipated minor. If the office has a written consent on file that has been signed by the parent authorizing the health care provider and his/her staff to provide medical care to the child, there is little or no risk in providing the child with routine medical care. In situations where the treatment is non-routine or poses some degree of risk to the child, it is always advisable to consult with the parent before proceeding.

In situations where it is not required to obtain the parent or guardian’s informed consent (e.g., treatment for venereal disease or substance abuse), is the parent or guardian responsible for the costs of this medical care?
Yes. Generally, parents or guardians are responsible for support of their unemancipated minors if the treatment is medically necessary. Note, if a minor patient is seeking alcohol or drug abuse treatment, federal law prohibits disclosure of patient identifying information to parents for the purposes of seeking reimbursement. 42 C.F.R. § 2.14. Thus, providers must obtain a minor’s consent before seeking reimbursement for these services from the minor’s parent or guardian.
In situations where it is not required to obtain the parent or guardian’s informed consent, may the health care provider inform the minor’s parents of the medical treatment?

In the case of substance abuse, federal regulations prohibit a health care provider from disclosing this treatment to the parent or guardian unless the minor lacks the capacity to make rational decisions, the situation poses a substantial threat to the child’s life or well-being, and the health care provider determines that this threat may be reduced by communicating the treatment to the parents. 42 C.F.R. § 2.14(d). In other circumstances, ethical and/or medical rather than legal concerns control the decision about whether providers should inform parents when there is no requirement to obtain the parent’s or guardian’s informed consent. There should be little legal risk of informing parents or guardians if the disclosure is in the best interests of the child. Note that parents may become aware of the visit if their insurance is used to pay for care.

In addition to obtaining informed consent from the parent or guardian, must providers also obtain informed consent from the minor?

No. If the parent or guardian is authorized to provide informed consent, health care providers are not also required to obtain the informed consent of the minor. That said, depending on the age and maturity level of the child, it is good practice to explain the procedure to the child and attempt to obtain his or her informed consent and cooperation.

In situations where an unemancipated minor has a child of her own, who is authorized to give informed consent for care of the minor’s child?

The child’s parent, regardless of age, is authorized to make decisions for her own minor child. This may lead to the anomalous situation where the parent is not legally capable of giving informed consent to her own health care, but can make decisions on behalf of her infant.

When minors are in the state’s custody, who is authorized to provide informed consent for their health care?

When a minor is in the custody of the State, the Commissioner of the Department of Children and Families (“DCF”) has authority to provide informed consent for a minor’s health care just as a parent would as outlined above. The Commissioner has additional authority to delegate this responsibility to other members of DCF. Social Services Policy Manual: Working with Families No. 74, 10/27/99.

Are minors in DCF custody able to receive contraceptive services?

According to the policies of the DCF, caseworkers will ensure that appropriate supportive counseling and contraceptive services are available to teens in custody. Social Services Policy Manual: Working with Families No. 74, 10/27/99.

May minors in DCF custody receive pregnancy-related services without informing their parents?

DCF staff may or may not inform parents of teens in custody about their pregnancy-related care, depending upon the DCF’s determination of what is in the best interests of the minor. Social Services Policy Manual: Working with Families No. 74, 10/27/99.
Who is authorized to consent to treatment when a child is in custody of the Department of Corrections?
The State of Vermont, through the Commissioner of the Department of Corrections (“DOC”), has exclusive authority to consent to medical treatment for children in the custody of the DOC. 28 V.S.A. § 1104. The DOC may try to involve the child's parents to obtain their input and background information.

When a minor in DOC custody receives medical care from a provider outside the prison, can the corrections officer accompanying the minor receive medical records of the visit?
Yes, a health care provider may give the records to the DOC officer transporting the patient in a sealed envelope marked: "To the attention of the responsible Department of Corrections health authority." Records may also be mailed to the "designated health authority" at the facility where the child is residing.

When minors are in custody of the Commissioner of Corrections who has access to their medical records or health care information?
The following individuals have access to health care information about a minor in custody of the Department of Corrections:
- The Commissioner, Deputy Commissioner, Director of Correctional Services, clinical director, medical director and their designees - on an as needed basis;
- Health care providers designated by the Department of Corrections;
- Non-health staff employed by the Department of Corrections have access as determined by health services staff.
Persons in custody are entitled to reasonable opportunities to discuss their medical care with health care providers. Guardians, including parents, have the same type of access to discuss care that the person in custody has so long as the person in custody has signed a written release approved by the DOC health care provider or a court has approved the guardian to act on behalf of the minor in custody. Department of Corrections Directive 254.02, Access to Health Care Records.

PRIVACY AND MEDICAL INFORMATION

What federal and state rules govern the confidentiality of medical records in Vermont?
Laws pertaining to general medical records are described below. Where pharmacy, mental health treatment, or substance abuse treatment records are at issue, there are additional federal and state laws to consider as discussed elsewhere in this Guide.

Federal Law
Health care providers must consider the requirements of the Health Insurance Portability and Accountability Act of 1996 (HIPAA), as modified by modified by the American Recovery and Reinvestment Act of 2009 (the “Health Information Technology for Economic and Clinical Health” (“HITECH”) Act), and its implementing regulations governing the electronic transfer and security of data and the privacy of medical information (collectively, “HIPAA privacy and security rules”). See 45 CFR Parts 160 and 164. These laws and regulations allow a provider to use and disclose health information for (1) providing treatment to the individual patient, (2) seeking payment for services, and (3) for the provider’s “health care operations.”
The HIPAA privacy and security rules provide a “floor” for protecting the privacy of identifiable health information, but they do not preempt state laws which are more protective of privacy or provide greater patient rights to access information.

**Vermont Laws**

Vermont law adopts the HIPAA privacy and security rule requirements for the disclosure of protected health information. 18 V.S.A. § 1881. There is also a patient privilege statute, which provides greater protection than the HIPAA privacy and security rules, and requires that doctors, chiropractors, dentists, nurses and mental health professionals must not disclose any information acquired in attending a patient unless the patient waives the confidentiality or it is waived by an express provision of law. 12 V.S.A. § 1612. A recent Vermont Supreme Court decision has limited the application of this privilege to judicial proceedings. *Kuligoski v. Brattleboro Retreat, et al*, 2016 Vt. 54A.

Hospital patients and nursing home residents are further protected by the confidentiality provisions contained in the Vermont Bill of Rights for hospital and nursing home patients, which require that the patient/resident must authorize the release or use of their records outside of the treatment team or the facility. 18 V.S.A. §1852(7) and the Nursing Home Bill of Rights, 33 V.S.A. §7301(2)(H). A nursing home resident’s record may be released without authorization when the resident is transferred to another health care institution or when required by a third-party payment contract. *Id.*

Health care providers are advised to have patients execute a general consent authorizing the provider to disclose protected health information for treatment, payment, healthcare operations and coordination of care purposes. This consent should be retained in the patient’s medical record.

*Are prescription records given the same confidentiality protection as other medical records?*

Under the federal HIPAA privacy and security rules, prescription records are treated the same as other health records and are subject to the same confidentiality provisions. Vermont law allows inspection of prescriptions, orders and prescription records only by federal or state officers or their specifically authorized agent whose duty it is to enforce the federal and state drug laws, or to authorized agents of professional licensing boards. 18 V.S.A. §§ 4211 and 4218.

The law also specifies that no privilege of confidentiality shall apply to information communicated to a physician in an effort to unlawfully procure a regulated drug or the administration of any such drug. 18 V.S.A. §4223(b). Such unlawful means include fraud, deceit, misrepresentation, subterfuge, forgery, concealment, or other unlawful or deceitful means. 18 V.S.A. §4223(a). Examples of such unlawful behavior may include failing to disclose that the patient is receiving regulated drugs from another prescriber, pretending to be an established patient of another physician to a covering physician, altering a prescription for 10 pills to be a prescription for 100 by adding a zero, or a patient saying they are taking a drug when a screen shows they are not.
When are pharmacists and providers required to report prescription records regarding dispensing of Schedule II, III and IV controlled substances?

Pharmacies and prescribers (i.e., health care professionals licensed to prescribe Schedule II, III or IV controlled substances) that dispense Schedules II, III or IV controlled substances (“Controlled Substances”) to their patients are required to submit a report to the Vermont Prescription Monitoring System (“VPMS”) no less frequently than every calendar week detailing the Controlled Substances dispensed by the prescriber or pharmacy during the preceding seven (7) days. Vt. Admin. Code Rule 12-5-21:4.0. Effective July, 2017, reports will be required within 24 hours of dispensing the prescription. See Vermont Prescription Monitoring System Rule, effective July, 2017.

Reports to the VPMS must include information related to the patient, prescription, dispenser and prescriber.

Dispensing prescribers and pharmacists must submit a “zero controlled substances report” during any week that no Controlled Substances are dispensed. Vt. Admin. Code Rule 12-5-21:4.0.

Each pharmacy shall provide to every customer to whom a controlled substance is dispensed an advisory notice informing the customer that all prescriptions for Controlled Substances are entered into a statewide VPMS database in order to protect patients and the public. The notice, available on the Department of Health’s website, must either be posted by the pharmacy in a prominent manner readily accessible to customers or duplicated in its entirety on a written insert for delivery to the patient. Vt. Admin. Code Rule 12-5-21:4.0.

Reporting to VPMS is not required when a drug is administered directly to a patient or dispensed by a health care provider at a facility licensed by the Department of Health, provided that the quantity dispensed at the facility is limited to an amount adequate to treat the patient for a maximum of 48 hours. A pharmacy that does not stock or dispense Controlled Substances may request an exemption from reporting from the VPMS program office. The exemption shall terminate when the pharmacy dispenses any controlled substance. Vt. Admin. Code Rule 12-5-21:4.0.

When are pharmacists and providers required to access prescription records of other providers regarding dispensing of Schedule II, III and IV controlled substances?

Prescribers and pharmacists registered with the VPMS are permitted to access the prescription records in the VPMS when the requested information is for the purpose of providing medical or pharmaceutical treatment to a bona fide current patient. 18 V.S.A. § 4284(b)(1).

Prior to prescribing Controlled Substances, Prescribers are required to query the VPMS in the following circumstances:

- The first time the provider prescribes an opioid Schedule II, III, or IV controlled substance written to treat pain when such a prescription exceeds 10 pills or the equivalent.; Rule Governing the Prescription Monitoring System, Section 6.2, effective July 1, 2017; see also 18 V.S.A. § 4289 (c)(3).
- When starting a patient on a Schedule II, III, or IV controlled substance for nonpalliative long-term pain therapy of 90 days or more; see also 18 V.S.A. § 4289 (c)(2).
• Prior to writing a replacement prescription for a Schedule II, III, or IV controlled substance; *see also* 18 V.S.A. § 4289 (c)(4).
• At least annually for patients who are receiving ongoing treatment with an opioid Schedule II, III, or IV controlled substance; *see also* 18 V.S.A. § 4289 (c)(1).
• The first time a provider prescribes a benzodiazepine. Rule Governing the Prescription Monitoring System, Section 6.2, *effective July 1, 2017 effective July, 2017*.
• When prescribing Schedule II, III or IV controlled substances to treat acute pain for a duration longer than 21 days; *(Sunsetting June 30, 2017)*
• In an Emergency Department or Urgent Care setting:
  o When a patient requests an opioid prescription for chronic pain from an Emergency Department or Urgent Care prescriber if the prescriber intends to write a prescription for an opioid.
  o When a patient requests an extension of a current opioid prescription for acute pain from an Emergency Department or Urgent Care prescriber if the prescriber intends to write a prescription for an opioid. *(This provision regarding extensions of current opioid prescriptions in the ED or urgent care will sunset on June 30, 2017)*
  o Before prescribing an opioid for longer than 10 days. *(The provisions described in these last two sub-bullets on June 30, 2017)*
• Prior to prescribing buprenorphine or a drug containing buprenorphine that exceeds the dosage threshold approved by the Vermont Medicaid Drug Utilization Review Board and published in its Preferred Drug List, prescribers must receive prior approval from the Chief Medical Officer or Medical Director of the Department of Vermont Health Access or designee;
• Prior to first prescribing an extended release hydrocodone or oxycodone that is not an abuse-deterrent opioid, and a query no less frequently than once every 120 days thereafter for any patient prescribed 40 mg or greater of hydrocodone or 30mg or greater of oxycodone per day of an extended release hydrocodone or oxycodone that is not an abuse-deterrent opioid;
  o During the initial query, the Prescriber must conduct a review of the other Controlled Substances prescribed to the patient prior to the first prescription of an extended release hydrocodone or oxycodone that is not an abuse-deterrent opioid.


As of July 1, 2017, all dispensers (with the exception of hospital-based dispensers) dispensing a quantity of a Schedule II, III, or IV opioid controlled substance that is sufficient to treat a patient for fewer than 48 hours will also be required to query the VPMS in the following circumstances:
• Prior to dispensing a prescription for a Schedule II, III, or IV opioid controlled substance to a patient who is new to the pharmacy;
• When an individual pays cash for a prescription for a Schedule II, III, or IV opioid controlled substance and the individual has prescription drug coverage on file;
• When a patient requests a refill of a prescription for a Schedule II, III, or IV opioid controlled substance substantially in advance of when a refill would ordinarily be due; and
• When the dispenser is aware that the patient is being prescribed Schedule II, III, or IV opioid controlled substances by more than one prescriber.

**Rule Governing the Vermont Prescription Monitoring System**, Section 5.2, **effective July 1, 2017**

What additional protection is given to mental health treatment records in Vermont?

Information and records pertaining to the treatment of mental illness and developmental disability or to involuntary hospitalization must be kept confidential, including information which directly or indirectly identifies the patient, unless one of a very limited list of exceptions applies (e.g., the mental health provider has a duty to warn or is required to file a report with the National Instant Criminal Background Check System), the patient or legal guardian consents in writing to the disclosure, or a court orders the disclosure. 18 V.S.A. §7103(a).

Under the HIPAA privacy and security rules, psychotherapy notes require specific authorization from an individual in order to be disclosed except when disclosure is required by law, or when disclosure is to a person reasonably able to prevent or lessen a serious and imminent threat to the health or safety of a person or the public. 45 CFR § 164.512(j)(1)(i). Psychotherapy notes refer to notes recorded by a mental health professional and documenting or analyzing a conversation during a private, group, joint, or family counseling session, are kept separate from the rest of an individual health record. 45 CFR §164.501 and §164.508(a)(2). Psychotherapy notes do not include medication prescription and monitoring, counseling session start and stop times, the modalities and frequencies of treatment furnished, results of clinical tests, and any summary of the following items: diagnosis, functional status, the treatment plan, symptoms, prognosis, and progress to date. *Id.*

When must a mental health provider disclose confidential patient information to warn about a patient who may present a serious risk of harm?

There are two situations which require a mental health provider to disclose confidential patient information to warn of a serious risk of harm:

**Duty to Warn An Identifiable Individual**

When a mental health provider has good reason to believe that a patient poses a serious risk of danger to an identifiable individual, the provider has a duty to exercise reasonable care to protect the identifiable victim from that danger even if it requires the disclosure of confidential patient information. This “duty to warn”, which has evolved in case law in Vermont and other states, is an exception to the patient privilege of confidentiality.

The duty to warn exception originated in California with the case of *Tarasoff v. Regents of the University of California*, 551 P.2d 334 (1976). In *Tarasoff*, a university hospital psychologist was told by his patient that the patient intended to kill a woman, Tatiana Tarasoff. Two months later, he did so. The Supreme Court of California ruled that the “public policy favoring
protection of patient-psychotherapist communications must yield to the extent to which disclosure is essential to avert danger to others.” 551 P.2d at 347.

In Peck v. Counseling Service of Addison County, the Vermont Supreme Court adopted the Tarasoff ruling in a situation where a patient receiving mental health services threatened to burn his father’s barn down. 146 Vt. 61 (1985). The Court held that if a mental health provider knows or should know that his patient poses a serious risk of danger to an identified person, the provider has a duty to take whatever steps are reasonably necessary to protect the identifiable victim from that danger, which could include reporting confidential patient information. The health care provider has a duty to exercise due care in determining what steps may be necessary to protect the identifiable victim of a patient’s threat of harm and what confidential information must be disclosed. Id. at 67. The health care provider must report confidential information discreetly, and in a fashion that would preserve the privacy of the patient to the fullest extent compatible with the prevention of the threatened danger, insuring that only that information which is necessary to protect the potential victim is revealed. Peck, 146 Vt. at 67-68, citing Tarasoff, 7 Cal.3d at 441.

Although to date the Vermont courts have not had occasion to extend the waiver of physician-patient confidentiality beyond mental health services, other states have done so when a patient is incapacitated as a result of medical treatment or disease and the person poses an obvious risk of serious harm to others. In these situations, a health care provider may have a duty to warn reasonably identifiable potential victims or a duty to take action to avoid the harm.

Duty to Warn of Potential Acts of Violence
The Vermont Supreme Court has recognized that there are additional circumstances where a mental health provider has a duty to warn (and thus disclose confidential patient information) even when there is no identifiable individual to whom the patient poses a risk. In Kuligoski v. Brattleboro Retreat, et al., the Vermont Supreme Court held that mental health providers have a duty to warn individuals within a “zone of danger” that a patient with dangerous propensities presents a serious risk of harm. 2016 WL 5793088 (September16, 2016).

The Court did not define the “zone of danger” precisely, but it includes those most likely to be victims if a patient engages in violence. While the Court declined to hold that the duty to warn extends to the general public, it did find that the “zone of danger” includes the identifiable “caretakers” of a potentially violent patient and thus these “caretakers” must be warned.

The Vermont Supreme Court held that the duty to warn applies when: “a caregiver is actively engaged with the patient’s provider in connection with the patient’s care or the patient’s treatment plan (or, in this case, discharge plan), the provider substantially relies on that caregiver’s ongoing participation, and the caregiver is himself or herself within the zone of danger of the patient’s violent propensities.” A “caretaker” does not need to be formally charged with legal responsibility for the patient, as in guardianship.

The Court also held that the “duty to warn” encompasses a “duty to inform,” which the Court described as a duty “to provide sufficient information to [caretakers] so they could fully assume their caretaker responsibilities to assist [the patient] and protect against any harmful conduct in
which he might engage.” Mental health providers must not merely warn of potential risks, according to the decision, they must also provide sufficient information to assist the “caretakers” in controlling the patient’s conduct and protecting third parties from harm.

*How do I determine if the patient poses a risk of danger to unidentified individuals requiring a warning?*

If a patient is considered to have violent propensities, Vermont law places the burden on mental health professionals to determine who may be a caregiver or within the “zone of danger” and what information must be disclosed in order to assist the “caretakers” in controlling the patient’s conduct and protecting third parties from harm.

In almost all instances, consultation with a colleague is advised to determine the most appropriate course of action given the conflicting duties imposed on the mental health practitioner.

*How does the duty to warn relate to HIPAA privacy and security rule requirements?*

The HIPAA privacy and security rules do not create a duty to warn or mandate disclosure. Rather, under the HIPAA privacy and security rules, a covered entity is permitted to use and disclose protected health information – including psychotherapy notes – if the covered entity believes, in good faith, that the use or disclosure is necessary to prevent or lessen a serious and imminent threat to the health or safety of a person or the public, and the disclosure is to a person or persons reasonably able to prevent or lessen the threat, including the target of the threat. 45 C.F.R. § 164.512(j). A covered entity may also disclose protected health if the covered entity believes, in good faith, that the use or disclosure is necessary for law enforcement to identify or apprehend an individual because of a statement by an individual admitting participation in a violent crime that the covered entity reasonably believes may have caused serious physical harm to the victim, or where it appears from all the circumstances that the individual has escaped from a correctional institution or from lawful custody. *Id.*

In the above situations, medical records and psychotherapy notes may be disclosed without authorization or consent of the patient. The disclosure must be made to a person or persons reasonably able to prevent or lessen the threat or to the target of the threat. The disclosure must be limited to the minimum amount of information necessary to prevent the harm from occurring.

What rules apply to the confidentiality of substance abuse treatment records?

Federal law affords separate provisions and protection to the confidentiality of alcohol and substance abuse treatment program records. 42 CFR Part 2. These provisions prohibit disclosure of the identity of anyone receiving alcohol and substance abuse treatment, and require a specific consent or authorization form (distinct from the form required by the HIPAA privacy and security rules) be executed by the patient before the disclosure of any treatment related information. The patient’s written consent must include the following information:

- The specific name or general designation of the program or person permitted to make the disclosure.
- The name or title of the individual or the name of the organization to which disclosure is to be made.
- The name of the patient.
• The purpose of the disclosure.
• How much and what kind of information is to be disclosed.
• The signature of the patient and, when required for a patient who is a minor, the signature of a person authorized to give consent; or, when required for a patient who is incompetent or deceased, the signature of a person authorized to sign in lieu of the patient.
• The date on which the consent is signed.
• A statement that the consent is subject to revocation at any time except to the extent that the program or person which is to make the disclosure has already acted in reliance on it.
• The date, event, or condition upon which the consent will expire if not revoked before. This date, event, or condition must insure that the consent will last no longer than reasonably necessary to serve the purpose for which it is given.
• The following written statement:
  o “This information has been disclosed to you from records protected by Federal confidentiality rules (42 CFR part 2). The Federal rules prohibit you from making any further disclosure of this information unless further disclosure is expressly permitted by the written consent of the person to whom it pertains or as otherwise permitted by 42 CFR part 2. A general authorization for the release of medical or other information is NOT sufficient for this purpose. The Federal rules restrict any use of the information to criminally investigate or prosecute any alcohol or drug abuse patient.”

45 C.F.R. § 2.32-2.33.

The written consent from the patient only permits the disclosure of the alcohol or substance abuse treatment records to the individual or organization and for the purposes specified on the form. The records may not be further disclosed or released by the receiving individual or organization absent the patient executing another consent. 42 C.F.R. § 2.32.

The federal rules related to substance abuse treatment records are being revised. This guide will be updated as additional information and guidance becomes available.

**Under what circumstances may a Vermont provider disclose health information or medical records without a patient’s consent or authorization?**

Under Vermont law, a health care provider may disclose health information or medical records without a patient’s consent or authorization only where there are “express provisions of law” or a court order. Disclosures required by “express provisions of law” include the following:

• **Child Abuse.** Health care providers are obligated to contact the Department of Children and Families (“DCF”) when they have reasonable cause to believe that any child has been abused or neglected. The health care provider must file a report with DCF within 24-hours of the time information regarding the suspected abuse or neglect was first received or observed. 33 V.S.A. §§ 4911-4920.

• **Crime Victim Under Age of Sixteen.** Health care providers are required to disclose information indicating that a patient under the age of sixteen has been a victim of a crime. 12 V.S.A. § 1612(b).

• **Abuse, Neglect or Exploitation of Vulnerable Adults.** Health care providers must report to the Department of Aging and Independent Living when they have reasonable cause to believe that a disabled adult or an adult suffering from infirmities of age or an adult
receiving personal care services at home or at a licensed facility has been abused, neglected or exploited. 33 V.S.A. §§6901-6914. Note, however, that under the HIPAA privacy and security rules, the victim must be notified of such a report or disclosure unless it is believed that such notification might place the individual at risk of serious harm. If the health care provider would be informing a personal representative of the victim, and the covered entity reasonably believes the personal representative is responsible for the abuse, neglect, or other injury, and that informing such person would not be in the best interests of the individual, the health care provider may use professional judgment in determining whether to inform the personal representative. 45 C.F.R. § 164.512(c)(2).

- **Firearm-Related Injuries.** Health care providers treating bullet wounds, gunshot wounds, powder burns or other injuries caused by the discharge of a gun, pistol, or other firearm must report such cases to local law enforcement officials or to the state police. 13 V.S.A. §4012.

- **Suspicious Deaths.** When a person dies from violence, or suddenly when in apparent good health or when unattended by a physician or a recognized practitioner of a well-established church, or by casualty, or by suicide or as a result of injury or when in jail or prison, or any psychiatric hospital, or in any unusual, unnatural, or suspicious manner, or in circumstances involving a hazard to public health, welfare, or safety, a physician notified of the death must report it to a medical examiner. 18 V.S.A. § 5205(a).

- **Requests by Chief Medical Examiner.** Information regarding the mental or physical condition of a deceased patient must be released by a physician, dentist, chiropractor, mental health provider, or nurse upon request from the chief medical examiner. 12 V.S.A. § 1612(c)(2).

- **Communicable Diseases.** If a health care provider has reason to believe that a person is sick or has died of a diagnosed or suspected communicable disease, identified by the Department of Health as a reportable disease and dangerous to the public health, he or she shall transmit a report thereof within 24 hours and identify the name and address of the patient and the name of the patient's physician to the Commissioner of Health. A list of reportable communicable diseases, including tuberculosis and venereal diseases, can be found at [http://healthvermont.gov/sites/default/files/documents/2016/11/hs_orid_vt_reportable_disease.pdf](http://healthvermont.gov/sites/default/files/documents/2016/11/hs_orid_vt_reportable_disease.pdf). 18 V.S.A. §§ 1001, 1004, 1007, 1041-1048, 1091-1106.

- **Fetal Deaths.** Health care providers are obligated by statute to report fetal deaths of certain gestational age or size and all therapeutic or induced abortions to the Department of Health within seven days after delivery. A physician who is treating a woman as a result of miscarriage and does not know if a report of fetal death has been made to the State shall file such a report. If there is evidence of violence or other unusual or suspicious circumstances surrounding the fetal death, the physician must report the death to the medical examiner immediately. 18 V.S.A. § 5222.

- **Cancer.** Providers are required to report each case of cancer to the Department of Health within 180 days of diagnosis, unless the patient has been previously diagnosed or admitted for cancer treatment at a hospital facility in Vermont. All health care facilities and health care providers who provide diagnostic or treatment services to patients with cancer shall report to the Department of Health any further demographic, diagnostic, or treatment information requested by the Commissioner concerning any person now or
 formerly receiving services, diagnosed as having or having had a malignant tumor. Additionally, the Commissioner or his or her authorized representative shall have physical access to all records that would identify cases of cancer or would establish characteristics of the cancer, treatment of the cancer, or medical status of any identified patient with cancer. 18 V.S.A. §§151-156.

- **Lead Poisoning.** All health care providers who analyze blood samples for lead levels or who use laboratories outside Vermont to analyze blood samples for lead levels shall report all information required by the Department of Health to the Department immediately by telephone if the result of any analysis is 45 micrograms or more of lead per deciliter of blood, or by electronic means within 14 days of analysis if the result of the analysis is less than 45 micrograms of lead per deciliter of blood. Any laboratory in Vermont that analyzes blood samples of Vermont residents for lead levels shall make reports as required by the Department. 18 V.S.A. §1755(d).

- **Blood Alcohol Level Reporting.** If a health care provider who is providing health services to a person in the emergency room of a health care facility as a result of a motor vehicle accident becomes aware as a result of any blood test performed in the health care facility that the person's blood alcohol level meets or exceeds the level prohibited by law, the health care provider shall report that fact, as soon as is reasonably possible, to a law enforcement agency having jurisdiction over the location where the accident occurred. 23 V.S.A. §1203b.

- **Duty to Warn.** In 2016, the Vermont Supreme Court held in *Kuligoski v. Brattleboro Retreat* that mental health providers have a duty to warn individuals within a “zone of danger”, including caretakers, that a patient presents a serious risk of harm. 2016 WL 5793088 (September 16, 2016). The Court further found that mental health providers must not merely warn those in the “zone of danger” of potential risks, they must also provide sufficient information to assist the “caretakers” in controlling the patient’s conduct and protecting third parties from harm. *Id.* For further discussion, see “When does a mental health provider have a duty to warn about a patient that may present a serious risk of harm?”

- **Bioterrorism.** Health care providers must report all cases of patients who exhibit any illness, disease, injury or death identified by the Department of Health as likely to be caused by a weapon of mass destruction, including illnesses, diseases, injuries or deaths which can result from bioterrorism, epidemic or pandemic disease, or novel and highly fatal infectious agents or biological toxins, and that might pose a risk of a significant number of human fatalities or incidents of permanent or long-term disability; or which can be caused by biological agents identified under federal law. A pharmacist must report any unusual or increased prescription requests, unusual types of prescriptions, or unusual trends in pharmacy visits that may result from bioterrorist acts, epidemic or pandemic disease, or novel and highly fatal infectious agents or biological toxins, and might pose a substantial risk of a significant number of human fatalities or incidents of permanent or long-term disability. Prescription-related events that require a report include an unusual increase in the number of prescriptions to treat fever, respiratory or gastrointestinal complaints, an unusual increase in the number of prescriptions for antibiotics, an unusual increase in the number of requests for information on over-the-counter pharmaceuticals to treat fever, respiratory or gastrointestinal complaints, and any
prescription that treats a disease that is relatively uncommon and may be the result of bioterrorism. 13 V.S.A. § 3504.

Can a hospital or health care provider inform visitors or callers about a patient’s location in the facility and general condition?
Yes, the HIPAA privacy and security rules permit a hospital or other health care provider to maintain a directory of certain information about patients and to use this facility directory to inform visitors or callers about a patient’s location in the facility and general condition. 45 C.F.R. § 164.510(a). A patient must be informed about the information to be included in the directory, and to whom the information may be released, and must have the opportunity to restrict the information or to whom it is disclosed, or opt-out of being included in the directory. The patient may be informed, and make his or her preferences known, orally or in writing.

The directory can include the following information: patient name, location in the facility, health condition expressed in general terms that does not communicate specific medical information about the individual and religious affiliation. The facility or health care provider may provide the appropriate directory information – except for religious affiliation – to anyone who asks for the patient by name unless the patient has restricted the information or opted-out of the directory. Religious affiliation may be disclosed to members of the clergy so long as the patient has been informed of this use and disclosure, and does not object.

When, due to emergency treatment circumstances or incapacity, the patient has not had an opportunity to express his or her preference about how, or if, the information may be disclosed, directory information about the patient may still be made available if doing so is in the individual’s best interest as determined in the professional judgment of the provider, and would not be inconsistent with any known preference previously expressed by the individual. In these cases, as soon as practicable, the health care provider must inform the patient about the directory and provide the patient an opportunity to express his or her preference about how, or if, the information may be disclosed.

What obligation does a provider have to notify a patient of an unauthorized use or disclosure of medical information of a Vermont patient?
Although Vermont law does not require any notification to a patient of a wrongful disclosure or use of medical information, the federal HIPAA privacy and security rules do. Under the HIPAA privacy and security rules, “breach” means the acquisition, access, use, or disclosure of protected health information in a manner not permitted by the HIPAA privacy and security rules, which compromises the security or privacy of the protected health information. 45 C.F.R. § 164.402. A breach does not include limited disclosures to and access by workforce members or other individuals authorized to access protected health information under the authority of the provider or through a business associate agreement.

Providers are required to notify a patient whose protected health information has been or is reasonably believed to have been breached no later than 60 days after the discovery of the breach. 45 C.F.R. § 164.404. The content and methodology of notification of a breach are set forth in federal regulations. Id. If the breach of unsecured protected health information involves more than 500 patients, a provider must notify media outlets as well as the individual patients.
45 C.F.R. § 164.406. Providers are required to keep a log of any breaches involving fewer than 500 patients and provide the log to the U.S. Department of Health and Human Services not later than 60 days after the end of each calendar year. 45 C.F.R. § 164.408.

If medical information that has been subject to a breach includes personal information, such as a social security, driver’s license, credit card, or financial account numbers (or passwords), notice of the breach must be made to the Vermont Attorney General within 14 business days of its discovery and to the individual as soon as possible and no later than 45 days after its discovery. 9 V.S.A. §§ 2430 and 2435.

**What liability or penalties could be imposed on a provider who wrongfully discloses the medical information of a Vermont patient?**

Under the HIPAA privacy and security rules, a civil penalty may be imposed by the U.S. Department of Health and Human Services Office of Civil Rights (OCR) ranging from $100 to a maximum of $1,500,000. 45 C.F.R. § 160.404. The Vermont Attorney General’s office, along with all states attorney generals, is authorized to enforce the HIPAA privacy and security rules through injunction or civil penalties ranging from $100 up to a maximum of $25,000 per calendar year. 42 U.S.C. § 1320d-5(d). Criminal penalties for certain egregious wrongful disclosures may be pursued by the U.S. Department of Justice, as well. 42 U.S.C. § 1320d-6.

There is no Vermont-specific penalty or cause of action for the wrongful release of confidential medical information. However, Vermont law does prohibit the disclosure of protected health information in violation of the HIPAA privacy and security rules. 18 V.S.A. § 1881. Additionally, there is a $2000 fine or up to one year of imprisonment or both for the wrongful disclosure of information related to hospitalization for mental illness. 18 V.S.A. § 7103(c).

Although in other states a person harmed by a wrongful disclosure of medical information may have a cause of action for invasion of privacy, defamation or breach of contract or fiduciary trust, Vermont has no statutory law or reported case law granting an individual a cause of action.

**May a health care or mental health provider in Vermont release medical records or disclose medical information in response to a subpoena?**

No, unless the patient has authorized or consented to the release, or, in the rare instance, that the subpoena has been issued by a Vermont or federal court rather than an attorney. A civil subpoena, commonly issued by attorneys, does not override the patient privilege of confidentiality. The Vermont patient privilege statute and the mental health information statute permit disclosures without patient consent only if required by law or ordered by a court of law. 12 V.S.A. §1612 and 18 V.S.A. §7103. In turn, health care providers should not release medical records or disclose medical information in response to a civil subpoena absent an authorization from the patient (or the guardian of a minor), a provision of law requiring the disclosure, or an order from a court.

Vermont law is to be distinguished as much stricter than the HIPAA privacy and security rules, which permit a health care provider to disclose health information in response to a subpoena if certain efforts have been made to notify the individual. 45 CFR §164.512(e). Because it...
provides for more stringent privacy protections, in the event of a subpoena, Vermont law should be adhered to rather than HIPAA privacy and security rules.

**What should a provider do if he is served a subpoena for medical records or to testify about medical information?**

The following steps are recommended:

- Determine whether there is a proper legal basis for releasing the patient’s records. For instance, a patient authorization may be attached to the subpoena. If so, the provider is authorized to release patient information in response to the subpoena.
- If no proper legal basis appears for releasing the records, contact the party issuing the subpoena to seek a voluntary resolution. Explain that the subpoenaed documents or information cannot be released without the patient’s authorization, and that none appears. Often, a subpoena will be withdrawn on this basis or authorization will be obtained. If the former occurs, document the withdrawal of the subpoena in writing by a follow-up letter.
- Under appropriate circumstances, it may be desirable to contact a patient in response to a subpoena to determine whether the patient will consent to the release of the requested documents or records.
- If the subpoena is not voluntarily withdrawn, and there remains no evident legal basis for releasing the records or information sought, determine whether the subpoena has been validly issued. Note that a subpoena issued by another state court is not valid in Vermont and requires no response. There may be other technical defects in the subpoena as well. (Check with your attorney.)
- If the subpoena is valid, a response must be made before the return date. Counsel should be contacted to make an appropriate response, either by written objection, a motion to quash or by a motion for protective order.

Some special comments are in order regarding responding to investigative or criminal subpoenas issued by prosecutors and law enforcement agencies. Subpoenas in this context may be seeking information about a criminal defendant or suspect, or about a victim (as in a rape case). Resistance from practitioners to release information may sometimes be interpreted by law enforcement agencies as lack of cooperation in criminal enforcement. These problems must be handled delicately and diplomatically, and may require meetings with local law enforcement agencies to explain the practitioner’s obligations to protect patient records and communications. At a minimum, law enforcement officials could be asked to obtain a district court order in support of a criminal subpoena. Such an order would ensure that due consideration is accorded the confidentiality rights of the patient whose records are being sought.

**What should a practitioner do if an officer has a search warrant for medical records?**

The practitioner must comply with the warrant as a search warrant is a court order issued after prior judicial approval and a showing of probable cause. The practitioner should call legal counsel as soon as possible.

Search warrants have a distinctly different purpose than a subpoena. Search warrants are limited to criminal proceedings to obtain evidence against a defendant suspected of a criminal violation. The defendant may be a health care provider or facility or a patient. Because a search warrant is only issued after a showing of probable cause, they are executed immediately and there is no
time between issuance and execution for the person subject to the warrant to challenge its legal validity. Therefore, challenges to search warrants are always made after the warrant has been issued and executed and prior to the introduction of the documents or objects in evidence.

A copy of the search warrant will be served on the person from whom or from whose premises the property was taken. In addition, the officer executing the warrant must provide a receipt for the property taken. (The basic requirements for state search warrants are set forth in V.R.C.P. 41.)

**What rights do the media or press have to patient information?**
The media’s rights to access information are no different than anyone else’s. They must either have patient authorization or a court order.

**MEDICAL RECORD**

**Who owns the medical record?**
Although the medical record contains patient information, the record belongs to the licensed organization or health care provider group who creates or maintains the record in order to provide medical services to the patient. As discussed below, federal and state law provide patients with rights to access medical information about themselves.

**When a provider leaves a group practice, what happens to the medical records?**
Medical records belong to the group practice, unless a provider’s employment or practice agreement provides otherwise. Patients, however, may always request that a copy of their medical record information be forwarded to another physician.

**What rules govern a patient’s access to his or her medical record?**
Under Vermont law, practitioners are required to provide patients prompt access to their records upon written request. A failure to do so constitutes unprofessional conduct. 3 V.S.A. §129a(a)(8) and 26 V.S.A. §1354(a)(10). Vermont law also requires that copies of medical records be provided to a patient’s representative or succeeding health care practitioner upon the patient’s written request, and that providers notify patients about how to obtain their records when a practice closes. *Id.*

The HIPAA privacy and security rules also require that medical records (except for psychotherapy notes) or copies thereof be made available to patients upon request. 45 C.F.R. § 164.524. Providers may require that the request for records be in writing but the provider must inform the patient of this requirement. *Id.* The HIPAA privacy and security rules require a health care provider to respond to a patient’s request to inspect or obtain a copy of their medical record within thirty (30) days. *Id.* This response time may be extended for a second thirty-day period. *Id.*
May access to a patient’s medical record ever be denied to the patient or his or her representative?

Vermont professional conduct laws require the prompt disclosure of a patient’s medical information to that patient, his/her representative or succeeding health care professionals. 3 V.S.A. §129a(a)(8) and 26 V.S.A. §1354(a)(10).

Unlike the Vermont statutes pertaining to unprofessional conduct, the HIPAA privacy and security rules specify the following limited circumstances under which a practitioner may deny access to an individual’s health information:

- When the portion of the medical record requested contains
  - Psychotherapy notes; or
  - Information compiled in reasonable anticipation of, or for use in, a civil, criminal, or administrative action or proceeding;
- When a health care provider acting under the direction of the Department of Corrections determines that permitting an inmate to obtain a copy of his/her medical record would jeopardize the health, safety, security, custody, or rehabilitation of the individual or of other inmates, or the safety of any officer, employee, or other person at the correctional institution or responsible for the transporting of the inmate;
- Under limited circumstances, when the protected health information is created or obtained by the health care provider in the course of research that includes treatment, an individual's access to the protected health information may be temporarily suspended for as long as the research is in progress;
- When an individual’s protected health information is contained in records that are subject to the Privacy Act, an act granting privacy to government-held records, if the denial of access under the Privacy Act would meet the requirements of that law;
- When the protected health information was obtained from someone other than a health care provider under a promise of confidentiality and the access requested would be reasonably likely to reveal the source of the information;
- When a provider has determined, in the exercise of professional judgment, that the access requested is reasonably likely to endanger the life or physical safety of the individual or another person (sometimes referred to as the “therapeutic privilege”);
- When the protected health information makes reference to another person (unless such other person is a health care provider) and a provider has determined, in the exercise of professional judgment, that the access requested is reasonably likely to cause substantial harm to such other person; or
- When the request for access is made by the individual's personal representative and a provider has determined, in the exercise of professional judgment, that the provision of access to such personal representative is reasonably likely to cause substantial harm to the individual or another person.

45 C.F.R. § 164.524(a).
The provider must provide the patient with a written denial of access detailing the basis for the denial of medical records. If the provider is relying on one of the last three bullets as the basis for denying access to the medical record, the patient has a right to seek review of the decision to deny access. The provider must, to the extent possible, give the patient access to any other protected health information requested, after excluding the protected health information as to which the provider has a ground to deny access.

As noted above, Vermont professional licensing law contains no similar exceptions permitting providers to deny access to medical information. Because Vermont law would be considered to be more beneficial to a patient, it is likely to govern and to preempt any contrary federal provision allowing for providers to deny access to records. Thus, before denying a patient access to his or her medical records based on the HIPAA privacy and security rules set forth above, a provider should consult counsel.

If a health care provider receives records from another provider, must the receiving provider produce the received records to a patient where a patient has authorized the release of all information?
Yes. The patient has a right to all of their medical information in the possession or under the control of a health care provider regardless of who generated the original documentation. 3 V.S.A. § 129a(a)(8) and 26 V.S.A. § 1354(a)(10).

Must health care providers release to a patient a record that has been labeled “Do Not Rerelease”?
Yes. Under Vermont law, a patient has access to all medical records in the possession or under the control of a health care provider. Generally, records labeled “Do Not Rerelease” are related to alcohol or drug abuse treatment. Records of such treatment are subject to federal regulations that require the provider to obtain specific consent from the patient before the provider further releases the record of alcohol or drug abuse treatment. 42 C.F.R. Part 2.

Who may authorize the disclosure of the medical record of a deceased individual?
For disclosures outside of a judicial proceeding, the HIPAA privacy and security rule disclosure requirements should be followed. (As noted elsewhere, Vermont law has specifically adopted them for disclosures of protected health information. 18 V.S.A. § 1881.) The relevant HIPAA provision allows that if an executor, administrator or other person has the authority to act on behalf of the deceased individual under applicable law, then that person, as the decedent’s personal representative, must be treated as having the same rights as the deceased individual. 45 CFR Section 164.502(g)(1) & (4). See also https://www.hhs.gov/hipaa/forprofessionals/privacy/guidance/health-information-of-deceased-individuals/.

In judicial proceedings, the Vermont patient privilege statute requires health care providers to disclose information regarding the mental or physical condition of a deceased patient (except for information that “would tend to disgrace the memory of the decedent”), if the privilege of confidentiality is waived:
- by the decedent’s personal representative, the surviving spouse or the next of kin of the decedent
• by any party in interest if, in any litigation, the interests of the personal representative are deemed by a trial judge to be adverse to those of the estate of the decedent or
• by the executor named in the will, or the surviving spouse or any heir-at-law or any of the next of kin or any other party in interest if the validity of the will of the decedent is in question.
12 V.S.A. § 1612(c).

No further Vermont law guidance has been provided by either legislation or court precedent to define the scope of this provision despite multiple areas of confusion (e.g., the definition of “next of kin” and what information “tends to disgrace the memory of the decedent”). Prior to the Vermont Supreme Court decision in Kuligoski vs. Brattleboro Retreat, et. al., 2016 Vt. 54A, the patient privilege statute referenced above was broadly applied in all disclosure situations, not just judicial proceedings.

What are health care providers allowed to charge patients or others, such as lawyers and insurance companies, for providing copies of a patient’s health care record?
A provider is allowed to charge a reasonable, cost-based fee, not to exceed a $5.00 flat fee or $.50 per page, whichever is greater. 18 V.S.A. § 9419 and 45 C.F.R. § 164.524(c)(4). The fees may only include costs for the following:
• Labor for copying the protected health information requested by the individual, whether in paper or electronic form;
• Supplies for creating the paper copy or electronic media if the individual requests that the electronic copy be provided on portable media;
• Postage, when the individual has requested the copy, or the summary or explanation, be mailed; and
• Preparing an explanation or summary of the protected health information requested (if the patient agrees to receive the summary in lieu of providing access to the full record and to pay for the costs of preparing the summary).
45 C.F.R. § 164.524(c)(4).

Practitioners and health care facilities are required to provide an itemized bill to the recipient of the records copied. Vermont law prohibits any charge for copies of records needed to support a claim or an appeal for public benefits such as welfare, Social Security, Medicare or Medicaid. 18 V.S.A. § 9419. The HIPAA privacy and security rules do not override this provision.

What is the difference between the federal regulation and state law on medical record copy charges?
Both Vermont law and the HIPAA privacy and security rules address the amount that may be charged for providing copies of a patient’s health care record. The two provisions provide as follows:
• Vermont law permits a health care provider or custodian of health records to charge a fee that is no more than a flat $5.00 fee or $0.50 per page, whichever is greater. No charges may be imposed when the records are requested to support a claim or an appeal for public benefits, such as welfare, Social Security, Medicare or Medicaid. 18 V.S.A. § 9419.
• The federal HIPAA privacy and security rules permit health care providers to charge a reasonable, cost-based fee for copying a patient’s medical records, based only on the cost
of labor, supplies, postage and, where applicable, the drafting of an explanation or summary of the protected health information. 45 C.F.R. § 164.524(c)(4).

A health care provider must follow whichever provision above results in the lower copying fee. For example, if your office’s reasonable cost-based fee of copying the records is less than the Vermont allowance of $.50 per page or $5.00 fee, you must charge the reasonable cost-based fee for the copies, unless it is to support a public benefits claim or appeal in which case no copy charge may be imposed. On the other hand, if the actual cost of providing copies exceeds the amount permitted by Vermont law, you are capped by the Vermont statutory allowance and can charge no more than $.50 per page or a $5.00 fee.

A provider is permitted to charge for the preparation of an explanation or summary of a record, in lieu of the full record, if the recipient has agreed in advance to receive an explanation or summary and if they have agreed in advance to the fees. 45 C.F.R. § 164.524(c)(2)(iii). A provider may also charge a fee for mailing, if the patient agrees in advance. 45 C.F.R. § 164.524(c)(4). A provider should document in the patient’s record whether they have requested that the record be mailed to them or have agreed to have the record summarized.

**What am I allowed to charge patients for providing copies of images?**

Vermont law allows health care provider offices to charge a cost-based fee for providing copies of x-rays, films, models, disks, tapes or health information maintained in other formats. 18 V.S.A. § 9419. This provision is consistent with the HIPAA privacy and security rules.

**How long must a provider retain medical records?**

Hospitals are required to retain medical records for a minimum of ten years as part of their state licensure obligations. 18 V.S.A. § 1905(8). Vermont law does not specify how long other providers should retain medical records. However, based on the statutes of limitation for certain causes of action under Vermont and federal law, all health care providers are advised to retain medical records for at least ten years after the patient was last treated by the provider.

The Vermont statute of limitations governing medical malpractice actions allows a cause of action to be brought within three years of the date of the incident, or two years of the discovery of the injury, but not later than seven years from the date of the incident. 12 V.S.A. § 521. Where fraudulent concealment has prevented the patient from discovering the injury, there is no time limit on when the patient can commence an action against the provider. Where the action is based on the discovery of a foreign object in the patient’s body, discovered later than seven years after the incident, the patient may commence an action within two years of the discovery of the object. Additionally, actions against health care providers brought under the federal or state False Claims Acts have a six-year statute of limitation, or up to ten years if the facts could not be reasonably discovered. 31 U.S.C. §§ 3731; 32 V.S.A. § 639.

Children’s records should be retained until at least three years following their eighteenth birthday because the statute of limitations for a minor to sue for a cause of action does not begin until the minor reaches the age of majority. 12 V.S.A. § 551(a).
Hospitals and other health care providers should establish medical record retention and disposal policies and procedures and manage all medical records accordingly.

What period of time is recommended for retaining the records of a deceased patient?
As a general rule, it is recommended that a provider retain records of deceased patients for no less than three years after the patient’s death or ten years after care was provided, whichever is longer. The wrongful death statute requires court actions to be commenced two years from the discovery of the death unless the death was deemed murder, in which case the cause of action must be commenced within seven years of the discovery of the death. 14 V.S.A. § 1492. The survival of actions law, however, permits court actions to be commenced two years after the date of issuance of letters testamentary or administration (i.e., if a person, by or against whom an action may be brought, dies before such action may be commenced) by the probate court. 12 V.S.A. § 557(a). Letters testamentary or administration can be issued some time after the death. Physicians should check with the court with respect to the timing of the issuance of letters testamentary.

What procedures should be followed if a health care provider seeks to amend a medical record previously created?
If it is necessary to correct an entry in a medical record, health care providers must make the correction without erasing, obliterating or deleting the original medical record entry. Alterations to or deletions of original records may raise the suspicion of an attempt to conceal the truth. Whether the correction is made in an electronic record or a paper record, the original entry should not be changed. The amendment should include the date and time of the amendment, a notation that the entry is an amendment (to distinguish it from the original record), and the reason for the amendment. Health care facilities and professionals should develop policies on correcting medical records and delegate authority to make corrections to specific identified individuals.

What rights does a patient have to amend his or her medical record?
Patients have the right under HIPAA privacy and security rules to seek to amend or supplement their own medical records for as long as the covered entity maintains the information. 45 C.F.R. § 164.526(b)(1). The provider may require patients to make requests for amendment in writing and to provide a reason to support a requested amendment, provided patients are informed in advance of such requirements. Id. The provider must act on a patient’s request for amendment no later than 60 days after it receives the request. 45 C.F.R. § 164.526(b)(2). The deadline may be extended up to 30 days. Id.

If a request to amend is accepted, the covered entity must make the appropriate amendment and inform the patient that the amendment is being made. The provider must also obtain consent from the patient for the provider to share the amended protected health information with other relevant persons, including the following individuals:

- Persons identified by the patient as having received protected health information about the patient and needing the amendment; and
- Persons, including business associates, that the provider knows
  - Have the protected health information subject to the amendment, and
May have relied, or could rely, on such information to the detriment of the patient. 45 C.F.R. § 164.526(c).

A request to amend may be denied if the health care professional or facility determines that the information or record:

- Was not created by the covered entity, unless the individual provides a reasonable basis to believe that the originator of the protected health information is no longer available to make the amendment;
- Is not part of the designated record;
- Would not be available for inspection under the patient’s right of access; or
- Is accurate and complete. 45 C.F.R. § 164.526(a)(2).

When a provider denies the patient’s request for an amendment, the provider must give the patient written notice of its decision that describes:

- The basis for the denial;
- The patient’s right to submit a written statement disagreeing with the denial and how to do so;
- A statement that the patient can request the health care professional or facility to include the patient’s request and the denial with any future disclosures of the information (if the patient does not file a statement of disagreement); and
- How the individual can file a complaint with the covered entity or the secretary of HHS. 45 C.F.R. § 164.526(d).

A health care provider that is informed by another provider of an amendment to a patient’s protected health information must amend the protected health information in his or her own records for the patient. 45 C.F.R. 164.526(e).

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EMPLOYMENT LAW

Topics Covered in this Chapter:
The Employment Relationship
Hiring
Employee Benefits
Documents and Record-Keeping
On the Job
Problems and End of Employment
About the Author

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This Employment Law Chapter is for informational purposes only and is not for the purpose of providing legal advice. Employers facing a specific issue or problem should seek the assistance of an attorney. Please note that the information in this Chapter assumes a non-unionized workforce.

THE EMPLOYMENT RELATIONSHIP

After an employer determines that a staff person is needed but before advertising or hiring, there are several decisions to be made. These include whether the worker will be classified as an employee or independent contractor, whether the person will have a contract or not, what wages can or must be paid, and what duties will be performed.

When can/should a health care professional or facility hire an independent contractor rather than an employee?

Some organizations may prefer to retain an independent contractor to save money on employment taxes and to avoid the cost of benefits, workers’ compensation, and unemployment. Some workers would rather be treated as independent contractors to avoid withholding from their paychecks and other reasons. However, the decision is not up to the individuals involved; different federal and state laws dictate whether a worker can be classified as an independent contractor rather than an employee. To make it even more complicated, the test is not the same under all laws applicable to the employment relationship. For example, a worker may be an independent contractor under the IRS test for purposes of federal tax withholdings, but be considered an employee for Vermont workers’ compensation and unemployment purposes, although some of the tests are similar. The following is a summary of these three tests.

(1) IRS Independent Contractor Test

For purposes of distinguishing between employees and independent contractors for federal tax withholding purposes, the IRS cares primarily about the degree of control exercised by the employer and the degree of independence experienced by the worker. More detail on the IRS’s independent contractor test can be found in IRS Publication 15-A, Employer’s Supplemental Tax Guide. The IRS test is grouped into three categories—behavioral control, financial control, and
type of relationship. For example, within the issue of behavioral control, the IRS looks at the
degree to which the person is generally subject to the organization’s instructions about when,
where, and how to work. If the organization directs when and where to do the work, provides
tools and equipment to the worker, purchases or tells the worker where to purchase supplies,
informs the worker what order or sequence to follow, and similarly has significant control over
how the work is performed, the worker will generally be considered an employee for federal tax
withholding purposes.

Within the issue of financial control, the IRS expects that an independent contractor will often
perform the same work for other organizations, receive a flat fee rather than an hourly wage,
have an opportunity for profit or loss, have unreimbursed expenses, or otherwise have a
significant investment in the work. Finally, the type of relationship helps determine the
classification. An employee is less likely to have a written contract, is more likely to receive
employee benefits, usually is not hired for a specific project or period, and performs work that is
a key aspect of the organization’s business. For example, generally the IRS would assume that a
physician or nurse hired by a physician’s office would be an employee, not an independent
contractor, as either professional would be performing work essential to the operations of the
organization. In contrast, the person who mows the lawn or cleans the office performs peripheral
duties and is more likely to be performing work under their own business, have other customers,
bring their own supplies, etc.

(2) Vermont Unemployment Insurance “ABC” Test

The tests for distinguishing between employees and independent contractors for Vermont
unemployment insurance and workers’ compensation coverage are narrower. For purposes of
Vermont unemployment insurance, all persons who perform services for wages are presumed to
be employees, entitled to unemployment benefits. To rebut this presumption, an employer must
show that a worker meets all three elements of the “ABC” test:

(A) such individual has been and will continue to be free from control or direction over
the performance of such services, both under his or her contract of service and in fact;
(B) such service is either outside the usual course of the business for which such service
is performed, or that such service is performed outside of all the places of business of the
enterprise for which such service is performed; and
(C) such individual is customarily engaged in an independently established trade,
occupation, profession or business.

21 V.S.A. § 341(a).

Unless all three elements of the ABC test are met, the worker must be classified as an employee
for purposes of unemployment compensation, even if the worker is classified as an independent
contractor for federal tax withholding purposes. The Vermont Department of Labor’s website
summarizes the test for unemployment insurance at: http://labor.vermont.gov/unemployment-
insurance/employers/who-is-an-employee-vs-independent-contractor/.

(3) Vermont Workers’ Compensation “Nature of Business” Test
To determine whether an employee-employer relationship exists for purposes of Vermont workers’ compensation law, a “nature of the business test” is used, i.e., whether the work that the owner contracted for is part of, or process in, the trade, business or occupation of the owner. Under this test, an individual is an employee for Vermont workers’ compensation purposes if he/she is hired to carry out some phase of the hiring entity’s business. The Vermont Department of Labor’s provides more detail on this test at: http://labor.vermont.gov/are-you-an-employee-or-contractor/.

The state tests for unemployment insurance and workers’ compensation are more inclusive than the IRS tests, so it is possible that a worker could be classified as an employee for certain purposes and an independent contractor for others. For example, a temporary replacement physician is not performing services outside the usual business of a physician’s office and therefore will be considered an employee for unemployment insurance and workers’ compensation, even though it may be possible to classify the physician as an independent contractor under the IRS test, depending on the nature and extent of his/her services.

Proper classification is important because misclassifying a worker as an independent contractor could lead to substantial liability for back taxes, overtime pay, benefits, and other liabilities.

**What is employment at-will?**

An employer must next decide whether to hire an employee on an at-will basis, or some other defined basis. Vermont law presumes that an employee who is hired for an indefinite period is an employee “at-will.” This means that the employee remains employed only as long as both the employer and employee agree. An at-will employee may leave employment at any time, and there is no law requiring that the employee give any notice. Similarly, an at-will employee can be terminated from employment with or without cause and with or without notice, as long as the employee is not terminated for an unlawful reason. Unlawful reasons for employment termination are discussed below in other sections.

An employer can modify the at-will relationship by hiring an employee for a definite period, making promises about job security, entering into a contract with the employee, or taking other steps that a court would deem sufficient to change the presumption. Creating personnel policies or practices that assure employees that they will have continued employment or that certain procedures will be followed before the employee is terminated is a common way the at-will relationship can be modified (sometimes inadvertently). If an employer wishes to preserve at-will status, all employment-related documents, including handbooks, policies, and offer letters, should reaffirm employees’ at-will status.

**How is an employment contract formed?**

An employment contract is formed similarly to other contracts — an offer is made, it is accepted, and consideration is given for it. It does not have to be in writing to be binding on the parties. Generally, if an offer of employment (which usually states or at least implies a promise of payment) is made and then is accepted by the employee, an employment contract is formed. Thus, virtually every employee is working under some form of contract; the important issue is what the terms of the contract are. Remember, if the contract did not specify a length of time or a term, it is generally considered an at-will contract and may be terminated at any time by either
party. If the contract is at all ambiguous, it is generally construed against the drafter. Therefore, if an employer sends an offer letter or prepares personnel policies or makes oral promises to an employee or applicant and the contents are at all unclear, they will be interpreted to favor the employee. Thus, any communications with employees should be made clearly and carefully. The organization should determine before hiring whether it intends to hire its employees on an at-will or other contractual basis and craft its offer letter, personnel policies, and practices accordingly.

What laws govern how an employee is paid?
The federal Fair Labor Standards Act (FLSA) and Vermont wage and hour laws govern certain aspects of how an employee must be paid. The federal law does not apply to all employers, so an organization must first determine whether it is covered by the FLSA or only by state law. The FLSA applies to enterprises with annual gross volume of sales made or business done of $500,000 or more; the FLSA applies to all hospitals, businesses providing medical or nursing care for residents, schools and public agencies are covered regardless of the dollar amount of business; in addition, the FLSA applies to any employee who engages in interstate commerce (this includes activities such as ordering supplies from another state over the telephone or Internet).

Under both federal and Vermont law, an employer must pay employees at least the applicable minimum wage and must pay time-and-a-half for any time worked over 40 hours in a workweek, unless the employee is specifically exempted from such overtime provisions. Both the FLSA and Vermont law require employers to keep accurate records of the hours worked by non-exempt employees and the wages paid to all employees. 21 V.S.A. § 393.

In Vermont, the minimum wage for employees is $9.60 an hour, as of January 1, 2016. Vermont’s minimum wage will increase to $10.00 per hour, starting January 1, 2017, and $10.50 per hour, starting January 1, 2018. Vermont law provides that its minimum wage will continue to increase each year thereafter. 21 V.S.A. § 384(a). Non-exempt employees of hospitals, public health centers, nursing homes, therapeutic community residences, maternity homes, and residential care homes must be paid overtime under Vermont law for any time worked over 40 hours in a workweek, unless the employer files an election and instead pays them time and a half for any hours more than eight in any day and 80 in a two week pay period. See 21 V.S.A. §384(b)(4); 29 C.F.R. § 778.601.

Vermont law requires that employers pay employees weekly unless they notify the employees in writing that they will be payed bi-weekly or semi-monthly, and payment must be made no more than six days after the end of the pay period. 21 V.S.A. §342(a). An employee who is terminated must be paid within 72 hours of discharge. 21 V.S.A. §342(b)(2). An employee who voluntarily leaves his/her employment must be paid on the next regular payday. 21 V.S.A. §342(b)(1).

How does overtime work?
If an employer is covered by the FLSA, every employee is presumed to be entitled to overtime pay for any time worked over 40 hours in a workweek (not 80 hours in a pay period). When an employee is eligible for overtime and minimum wage, it is referred to as “non-exempt”. Some positions are “exempt” from the requirements of the law, but the exemptions are intended to be
quite narrow and construed against the employer. An employee must specifically meet an exemption or she/he is entitled to overtime pay.

The three primary exemptions applicable to physician’s offices are the executive, administrative, and professional exemptions. The first requirement for all three is that the employee be paid on a salary basis of at least $455* for each week in which the employee performs any work (with a few exceptions). (*An Overtime Rule was set to increase the minimum weekly salary to $913 effective Dec. 1, 2016, but as of the date of this Guide, the implementation of the Rule had been halted by a nationwide preliminary injunction.) In addition to this “salary basis” test, the employee must also satisfy a “job duties” test to be exempt. The executive employee’s primary duty must be to manage a department, direct the work of two or more other full time employees, and have the authority to hire and terminate (or at least have his/her recommendations carry particular weight). The administrative employee must perform office work directly related to the organization’s general business operations, and the employee’s primary duty must include the exercise of discretion and independent judgment with respect to matters of significance. The professional employee must have advanced, specialized knowledge in a recognized field of science or learning (such as an MD or an RN) customarily acquired from a prolonged course of specialized intellectual instruction. Generally, a Licensed Practical Nurse does not fit the “professional” exemption. Unless an employee meets the test to be exempt under the FLSA, the employee must be classified as non-exempt. Determining whether or not an exemption applies to a particular position or employee can be complicated and may require consultation with an attorney or other human resource professional. It is prudent to seek legal guidance with any questions about employee classification.

An employer must pay a non-exempt employee one and a half times the employee’s regular rate of pay for any time worked over 40 hours in any workweek, even if the employer has not authorized or expressly permitted the overtime. The employee may not waive the overtime pay, and an employer may not offer compensatory time in a different workweek in lieu of the overtime pay. If an employee works unauthorized overtime, the employer may take disciplinary action, including termination, against the employee, although the employee must be paid for the time worked.

Why should an employer have job descriptions?
Written job descriptions help define the expectations an employer has for its employees, reduce the risk for disagreements about the scope of job requirements, and provide a basis for evaluation of employee performance and appropriate compensation. In addition, job descriptions identify the essential functions of a job for purposes of engaging in the interactive process and providing reasonable accommodations for qualified individuals with disabilities. Ideally, a job description should identify those functions the employer deems essential before the employee is even hired.

HIRING

Like other employers, physicians may wish to use a range of measures to choose employees, including advertisements, background checks, applications, interviews, and references.
What should be in a job advertisement?
The content of a job advertisement varies by the nature of the job and extent of the search. It is unlawful for an employer to publish a job advertisement that shows a preference for or discourages someone from applying based on a category protected by law (e.g., their age, race, color, religion, sex, gender identity, sexual orientation, pregnancy, national origin, disability, genetic information or any other protected category). The advertisement must be truthful and not misleading, although it need not include all of the particulars of the job. Many advertisements include an equal employment opportunity statement.

What should be included in an EEO policy?
An equal employment opportunity (“EEO”) policy in Vermont should include the statement that an employer provides equal employment opportunities to all employees and applicants and that it will not discriminate on the basis of race, color, religion, ancestry, national origin, place of birth, sex, gender, sexual orientation, gender identity, age, pregnancy, HIV-positive status, veteran status, military service or obligation, genetic information, or against a qualified individual with a disability, citizenship, immigration status, or any other category protected by law. In addition, an employer must ensure that there is no retaliation against an individual who has opposed any alleged discrimination, has lodged a complaint of discrimination, has cooperated with a state investigation into a discrimination complaint, or who the employer believes may be about to lodge such a complaint. Some employers voluntarily choose to add protection in their policies for categories that may not be protected by law, but are particular to the employer’s clientele or mission.

Though there are different thresholds for coverage under each of these laws, the following is a non-inclusive list of laws that prohibit discrimination in employment based on protected categories: Title VII of the Civil Rights Act of 1964, Age Discrimination in Employment Act, Americans with Disabilities Act, Americans with Disabilities Act, Rehabilitation Act, Pregnancy Discrimination Act, Equal Pay Act, Family and Medical Leave Act, Immigration Reform and Control Act of 1986, Title II of the Genetic Information Nondiscrimination Act, Vermont’s Fair Employment Practices Act, Vermont’s Parental and Family Leave law, Vermont’s Healthcare Whistleblower’s Protection Act, and Vermont’s Sexual Harassment law.

What questions are allowed or prohibited in an application or interview?
An employer should not ask questions that elicit information about protected characteristics such as age, sex, race, religion, national origin, sexual orientation, gender identity, disability, or any other category protected by law as the question alone may be evidence supportive of a discrimination claim. Employers should also not ask an applicant about his/her workers’ compensation history, birthplace, dates of attendance at school, arrest record, or marital status. Instead, a job application or interview should focus on assessing the applicant’s skill set, ability, and possession of qualifications necessary to perform the job – licensing, education, employment history, performance of expected duties. If the applicant will be driving a car, inquiries about a driver’s license may be appropriate.

The law requires that employers provide reasonable accommodations to both employees and job applicants with a disability, unless doing so would cause undue hardship (significant difficulty or expense for the employer).
There are also strict limits on when employers can ask applicants or employees about medical issues, disabilities or require a medical examination. Under the Americans with Disabilities Act (“ADA”), a disability is defined in one of three ways: (i) a physical or mental impairment that substantially limits one or more major life activities; (ii) a record or past history of impairment; or (iii) being regarding as having an impairment. 29 C.F.R. § 1630.2(g)(1). Under the law, the definition of disability must be construed in favor of broad coverage of individuals. The ADA does not contain a list of all conditions which constitute disabilities, but impairments such as epilepsy, diabetes, cancer, HIV infection, and bipolar disorder nearly always meet the definition of a disability.

An employer cannot ask a job applicant if they have a disability (or about the nature of an obvious disability), to answer medical questions, or take a medical exam before extending a job offer. At the application and interview stage, employers may only ask job applicants whether they can perform the job and how they would perform the job (but an employer cannot ask the applicant if she/he needs a reasonable accommodation to do the job). Thus, an applicant who is blind or visually impaired may be asked specifics of how she/he would perform the duties of the job but cannot be asked whether she/he has any sight or how she/he became blind or whether his/her vision is expected to improve in the future. After a conditional job offer has been made, the law allows the employer to ask disability-related questions or require a medical examination, but only if they are required of all entering employees within a job classification. To withdraw a job offer following such inquiries, the employer must demonstrate the reason was job-related and consistent with business necessity and that no reasonable accommodation would enable the applicant to perform the essential functions of the job.

Once an employee has started working, an employer generally can only ask medical questions or require a medical exam if the employer needs medical documentation to support an employee’s request for an accommodation or if, based on current, objective evidence, the employer believes that an employee is not able to perform a job successfully or safely because of a medical condition, i.e., when it is job related and consistent with business necessity. Employers must treat any medical information obtained from a compliant inquiry or disclosed by the employee as a confidential medical record and maintain it separately from the employee’s personnel file.

**Can I ask about criminal histories on job applications?**

Effective July 1, 2017, most Vermont employers will be prohibited from asking about an individual’s criminal history record on initial employment applications. 21 VSA § 495j. Once the law takes effect, employers can still ask about a prospective employee’s criminal history record during a job interview or once the applicant has been deemed otherwise qualified for the position. Under the law, if an applicant divulges criminal history in response to an employer inquiry, the employer is required to offer that applicant the opportunity to explain the circumstances, including any post-conviction rehabilitation, if the criminal history record does not disqualify the applicant under federal or state law. An employer will only be able to ask about criminal offenses on an initial application if it is subject to an obligation imposed by federal or state law or regulation that prohibits it from employing an individual who has been convicted of certain criminal offenses. However, the questions on the application must be limited only to the types of criminal offenses that create the legal disqualification.
What kind of background checking may or should an employer do before hiring an employee?
The extent of background checking to be done before hiring an employee depends on the nature of the position and duties to be performed and must be performed in compliance with the law. Some employers have a legal requirement to check background checks.

When can an employer do a criminal background check on an applicant?
Effective July 1, 2017, an employer must not perform a background check on an applicant until after a job interview or the applicant has been deemed otherwise qualified for the position. 21 VSA § 495j.

Under Vermont law, any employer may obtain criminal records directly from the Vermont Criminal Information Center (VCIC) if the requirements of 20 V.S.A. § 2056c are met.

When an employer uses a third party (i.e., not the employer’s own employee) to conduct a background check or to obtain reports from outside agencies, such reports are subject to the federal Fair Credit Reporting Act (“FCRA”). Background information is either classified as a “consumer report” or an “investigative consumer report.” The third party company performing the background check is called a “consumer reporting agency.” A “consumer report” is defined as any communication that contains information about an individual’s credit worthiness, credit standing, credit capacity, character, general reputation, personal characteristics or mode of living. A “consumer report” includes reports on criminal history, driving records, verification of education, etc., not just credit reports. Generally, the following notices are required when an employer is obtaining a “consumer report” that is covered by the FCRA:

1. Written disclosure to the applicant/employee;
2. Written authorization from the applicant/employee;
3. Certification to the reporting agency that it will comply with the FCRA;
4. Before taking adverse action based on the consumer report, provide the person with a copy of the report, and a statement of his/her rights from the FCRA; and
5. Providing notice of the adverse action. This must include (a) a statement that adverse action was taken based on the consumer report; (b) contact information for the consumer reporting agency; (c) a statement that the consumer reporting agency did not make the adverse decision and cannot explain why that decision was made; (d) an explanation of the applicant's or employee's rights under the FCRA to obtain a free copy of the consumer report from the reporting agency and to dispute with the agency the accuracy or completeness of the report.

An “investigative consumer report” is when the third party consumer reporting agency obtains information about character, etc. from personal interviews. Employers must ensure they are following the FCRA’s additional disclosure requirements if they decide to have a background check company or private investigator obtain an “investigative consumer report.”

The FCRA does not cover an employer’s own background investigation conducted by the employer’s own staff; the FCRA applies to background reports from private investigators or other consumer reporting firms.
The Equal Employment Opportunity Commission emphasizes that in all cases with background checks, employers treat everyone equally. It is unlawful, for example to only check backgrounds for individuals based on their membership in a protected category (e.g., race, national origin, age, etc.) or to use a background check in a manner that discriminates disproportionately against individuals in protected categories. Employers should have a business necessity to justify using a conviction record (and should generally never ask about arrest records) taking into account the nature and gravity of the offense(s), the time that has passed since the conviction, and the nature of the job.

When can an employer do a credit check?
In addition to limitations under the FCRA set forth above, Vermont law (21 V.S.A. § 495i) prohibits employers from inquiring into an applicant or employee's credit report or credit history except if certain conditions are met. Under Vermont law, a credit history (which includes a credit report) may only be required if the position involves access to confidential financial information, a financial fiduciary duty to the employer, access to payroll information, or the employer can demonstrate the information is a “valid and reliable predictor of employee performance in the specific position of employment.” If an employer will obtain a credit history or act on a credit history, it must obtain written consent each time, disclose in writing the reasons for accessing the credit history, disclose in writing the reason for adverse action, if any, not charge the employee for the cost associated with obtaining a credit history, keep the information confidential, and if the applicant is not hired, either destroy the credit history in a secure manner or give it to the applicant.

When can an employer conduct drug or alcohol testing?
Vermont has a strict drug testing laws. An employer may only require an applicant to submit to a drug test if:

- the applicant has already been offered the job conditioned on a negative test result;
- the applicant receives written notice of the employer’s policy, the procedure, and list of drugs to be tested;
- the employer follows the specific procedures required by the law, which includes a strict chain of custody, use of designated state labs, an appointed medical review officers and collector, etc.; and
- the employee is given an opportunity to retest any positive result.

No random or company-wide drug tests are permitted unless federal law requires it (i.e., for certain transportation employees). To test an employee, the employer must:

- have probable cause to believe the employee is using or under the influence of a drug on the job;
- have an employee assistance/rehabilitation program (EAP);
- not terminate the employee if the employee agrees to complete the EAP; and
- follow the specific testing process in the act.

More information is available at Vermont’s Drug Testing Act, 21 V.S.A. §§511-520.
When can/must an employer check an applicant’s licensing or professional status?

An employer can check an applicant’s (or employee’s) licensing or professional status at any time. The National Practitioner Data Bank (NPDB) and Healthcare Integrity and Protection Data Bank (HIPDB) allow certain entities, including a “health care entity,” to obtain information about medical malpractice or similar claims. 45 C.F.R. Part 60. A “health care entity” is a hospital or an organization that provides health care services and follows a formal peer review process for the purpose of furthering quality health care, or a committee of that entity, including a professional society, HMO, or medical or dental group practices following a formal peer review process. A health care entity can check a health care practitioner’s licensing or professional status when the health care entity is employing, affiliated with or granting privileges to the health care practitioner. The health care entity may check the health care practitioner’s status at any time. Only an authorized entity may obtain information from the databanks, and there are requirements about registration, fees, and other criteria available from the NPDB website. Authorized entities include state licensing and professional agencies, law enforcement, other state and federal agencies, utilization and quality peer review organizations, health plans and, in limited circumstances, attorneys or individuals pursuing medical malpractice claims. For more information, see the National Practitioner Databank section in the Professional Liability chapter.

The Vermont Board of Medical Practice maintains a database of licensed physicians. (http://healthvermont.gov/health-professionals-systems/board-medical-practice/look-license; see also 26 VSA § 1368). The Vermont Secretary of State’s Office of Professional Regulation licenses most other health care professionals, including nurses, pharmacists, radiologic technologists, occupational therapists, physical therapists, and many others. Its website includes a searchable database of professionals and their licenses. (https://www.sec.state.vt.us/professional-regulation.aspx). Licenses and specific information regarding the discipline of a licensee are public records, although ongoing investigations or prior complaints that did not result in discipline are not. 3 VSA § 131. Given the easy availability of the information, an employer would be hard-pressed to explain why it did not check to ensure that an employee was properly licensed.

May an employer require an employee to sign a covenant not to compete before beginning work?

Yes. As of the date of this Guide, Vermont law does not prohibit an employer from requiring that an employee sign a restrictive covenant agreement not to compete with the employer, as long as the restraint is narrowly tailored in geographical, temporal, and subject matter restrictions to protect the employer’s legitimate interests.

EMPLOYEE BENEFITS

Earned Sick Leave

Vermont passed earned sick leave legislation in 2016. 21 V.S.A. §§ 481-486. All employers with more than five employees who work on average 30 or more hours per week must start providing earned sick time on January 1, 2017. Employers with five or fewer employees who are employed for an average of 30 or more hours per week must start providing earned sick time on January 1, 2018.
Generally, all employees who work on average at least 18 hours per week must receive earned sick time. The law contains several classifications of employees who may be excluded from eligibility for earned sick time, including: individuals under 18 years of age, seasonal employees (employees who work 20 or fewer weeks in a 12-month period in a job that is scheduled to last 20 weeks or fewer), and per diem employees of health care facilities. Employees can use earned sick time for any of the reasons listed in 21 V.S.A. § 483(a):

1. The employee is ill or injured.
2. The employee obtains professional diagnostic, preventive, routine, or therapeutic health care.
3. The employee cares for a sick or injured parent, grandparent, spouse, child, brother, sister, parent-in-law, grandchild, or foster child, including helping that individual obtain diagnostic, preventive, routine, or therapeutic health treatment, or accompanying the employee's parent, grandparent, spouse, or parent-in-law to an appointment related to his or her long-term care.
4. The employee is arranging for social or legal services or obtaining medical care or counseling for the employee or for the employee's parent, grandparent, spouse, child, brother, sister, parent-in-law, grandchild, or foster child, who is a victim of domestic violence, sexual assault, or stalking or who is relocating as the result of domestic violence, sexual assault, or stalking. As used in this section, "domestic violence," "sexual assault," and "stalking" shall have the same meanings as in 15 V.S.A. § 1151.
5. The employee cares for a parent, grandparent, spouse, child, brother, sister, parent-in-law, grandchild, or foster child, because the school or business where that individual is normally located during the employee's workday is closed for public health or safety reasons.

Covered employees must accrue one hour of earned sick time for every 52 hours worked. All hours actually worked by non-exempt employees will count towards the accrual of earned sick time. However, an employer can limit the number of hours in a workweek that will count towards the accrual of earned sick time for exempt employees to 40 hours. Regardless of the number of hours worked by an exempt or non-exempt employee, employers can limit the total amount of earned sick time an employee can accrue to 24 hours in a 12-month period during 2017 and 2018. In 2019 and thereafter, employers can limit the total amount of earned sick time an employee can accrue to 40 hours in a 12-month period.

Employers can impose a waiting period of up to one year during which an employee must accrue earned sick time, but can be prohibited from using the earned sick time until after he or she has completed the waiting period.

All earned sick time that remains unused at the end of every annual period must generally carry over to the next annual period. However, carry-over is not required if an employer offers a paid time off policy and provides the employee with access to their full accrual of earned sick time at the beginning of each annual period (i.e, the time is “front loaded”). Also, an employer, at its discretion, can pay out accrued but unused earned sick time at the end of an annual period, and then the amount of earned time for which the employee was compensated would not carry over.
Employers do not have to pay out earned sick time at separation from employment.

An employer’s existing paid time off policy will comply with Vermont law if the paid time off accrues at a rate that is greater than equal than the rate in Vermont’s law and may be used for the same purposes and with at least the same rights as they would be able to use earned sick time under Vermont’s law. Employers can offer paid time off benefits that are more generous than the provisions under Vermont law, if they choose to. Employers who already offer paid time off should review and revise existing policies for compliance with the provisions of Vermont’s earned sick time law.

All employers must post a poster on earn sick leave issued by the Vermont Department of Labor. Employers are also required to notify employees about earned sick leave benefits at the time of hire.

For more information, including a link to the poster, frequently asked questions and the Department of Labor’s final proposed rules on earned sick time, see the Department of Labor’s website [here](#).

**Paid Vacation or Holidays**

Employers are not required to provide employees with paid vacation or paid holidays. However, if an employer and employee have an oral or written agreement providing specific benefits to the employee and the employer fails to provide the benefits, the employer is liable for actual damages caused by failure to pay the benefits. 21 V.S.A. § 345(b).

**Breaks**

Vermont law does not require an employer to provide employees with any specific break or lunch periods but requires that employers provide “reasonable opportunities during work periods to eat and to use toilet facilities in order to protect the health and hygiene of the employee.” 21 V.S.A. §304.

**Nursing Mothers**

Vermont law requires employers to provide nursing mothers, for three years after the birth of a child, with reasonable time (paid or unpaid) throughout the day to express breast milk and to provide accommodations, including appropriate private space to express milk that is not a bathroom stall. 21 V.S.A. §305.

The federal Fair Labor Standards Act also requires employers to provide reasonable break time for an employee to express breast milk for her nursing child for one year after the child’s birth, each time such employee has the need to express milk. 29 U.S.C. § 207(r). Under the FLSA, employers are required to provide a place for the employee to express breast milk, other than a bathroom, that is shielded from view and free from intrusion from coworkers and the public.

Employers are not required under state or federal law to compensate nursing mothers for breaks taken for the purpose of expressing milk. However, where employers already provide compensated breaks, an employee who uses that break time to express milk must be compensated in the same way that other employees are compensated for break time. In addition,
employers must follow the FLSA’s general requirement that the employee must be completely relieved from duty or else the time must be compensated as work time applies. Employers should treat breaks taken by nursing mothers to express milk consistently with the way other break time is compensated.

Flexible Working Arrangements
All Vermont employers are required by law to consider employee requests for flexible working arrangements. 21 V.S.A. § 309. “Flexible working arrangements” mean intermediate or long-term changes in the employee’s regular working arrangements, including changes in the number of days or hours worked, changes in the time the employee arrives at or departs from work, work from home, or job-sharing. Under the law, employers must engage in good faith discussions concerning employee requests for flexible working arrangements. Employers are obligated to respond to employee requests at least twice per calendar year. An employer has discretion to deny a request when it is inconsistent with business operations. Vermont law lists eight possible reasons denial may be appropriate, including situations where a request: imposes additional costs, has a detrimental effect on aggregate employee morale or the ability to meet consumer demand, or there is an inability to reorganize work among existing staff. It is important to note that if an employee requests a flexible work arrangement in writing and the employer denies any part of the request, the employer must provide a written denial. Retaliation against employees who exercise their rights under this law is also prohibited. Employers should implement internal procedures to document employee requests and efforts taken to evaluate and respond to requests.

Group Health or Retirement Benefits
If an employer does choose to offer group health or retirement benefits, the Employee Retirement Income Security Act of 1974 (“ERISA”), the Internal Revenue Code, and the Affordable Care Act (group health only) are the principal laws that affect such plans. Employers should consult with an employee benefits advisor, a human resources professional or an attorney concerning an employer’s obligations in connection with offering group health or retirement benefits.

Workers’ Compensation and Unemployment Insurance
Employers are required to provide workers’ compensation and unemployment compensation coverage for all employees. Under Vermont's workers’ compensation law, an employee is entitled to compensation if the worker “receives a personal injury by accident arising out of and in the course of his employment.” Generally, any employer who employs at least ten employees for more than 15 hours per week must reinstate to the first available, suitable job, any employee if the employee recovers within two years of the onset of the work related injury. Recovery is determined if the employee can “reasonably be expected to perform safely the duties of his or her prior position or an alternative suitable position.” 21 V.S.A. § 643b(a)(2).

The employer need not reinstate the employee to the same job, but must reinstate him/her to “the first available position suitable for the worker given the position the worker held at the time of the injury.” 21 V.S.A. § 643b(b). The employee also regains seniority and unused leave time that had been earned up to the date of the injury. Reinstatement is not required if the employee had prior notice or had already given notice before the injury that his/her employment would end, if the employment would have ended on its own, or if the employee fails to stay in touch
with the employer about his/her interest in reinstatement, status of recovery, and current mailing address. 21 V.S.A. § 643b(d). Under Vermont’s workers’ compensation law, an employer need not hold a job open for an employee who is out on workers’ compensation leave, unless the employee’s leave also qualifies for job-protected leave, such as leave under the FMLA or VPFLA. While an employee is on workers’ compensation leave and the employee’s prior job or a similar job is open when the employee recovers (and less than two years has passed since the employee was injured), the employer must reinstate the employee. Vermont employers are not required to establish light duty positions, but may do so.

For more information on Unemployment Insurance, see the section below, What is Unemployment Compensation.

Family and Medical Leave
In addition to Earned Sick Leave requirements under Vermont law, many employers are required to provide unpaid, job-protected leave for qualifying family, medical and parental reasons. Depending on their size, employers may be covered by the federal Family and Medical Leave Act (“FMLA”) and/or Vermont’s Parental and Family Leave Act (“VPFLA”).

Who must comply with the FMLA and VPFLA?
The FMLA allows eligible employees to take reasonable unpaid leave for medical reasons, for the birth or adoption of or bonding with a child, to care for a child, spouse, or parent who has a serious health condition, or for qualifying exigencies related to the foreign deployment of a military member. It applies to all private employers who employ 50 or more employees. Vermont has enacted a family leave law as well, which requires any employer that employs fifteen or more eligible employees to provide medical leave and any employer with at least ten eligible employees to provide parental and short-term leave. If an employee is covered by both federal and state leave law, the employee must receive the more favorable benefits so that the leave of absence will run concurrently under the FMLA and VPFLA.

Which employees are covered?
To be eligible under the FMLA, an employee must have been employed for at least twelve months (not needing to be consecutive) with at least 1,250 hours of service during the previous twelve months and work at a worksite that has at least 50 employees within 75 miles. The VPFLA applies to any employee who has been continuously employed by the same employer for at least one year, who has worked an average of at least 30 hours per week during the previous year.

May an employer require advance notice and medical certification?
An employer may require an employee to give at least 30 days’ notice of his/her intent to take leave when the need for leave is foreseeable. When the need for leave is unforeseeable, notice must be provided as soon as practicable. If the leave is for a serious illness, the employer may require certification from a health care provider of the condition and the need for the leave. The federal Department of Labor has created specific FMLA forms for employers to use, including certifications for health conditions which include only the information to which an employer is entitled. The FMLA requires that an employer notify the employee if medical certification will be required and allow the employee 15 days to return the form; it also limits the information the
employer can ask. For example, the employer generally cannot call the employee’s physician and ask questions, except where it is within limited circumstances permitted by FMLA regulations.

A covered employer must notify the employee requesting leave whether or not she/he is eligible under the FMLA. If the employee is eligible for FMLA leave, and once the employer has enough information to determine that leave is being taken for a FMLA-qualifying reason, the employer must notify the employee that the leave is designated and will be counted as FMLA leave, and the amount of leave counted against the employee’s leave entitlement. The employer must also notify the employee if the leave is not FMLA-protected. The federal Department of Labor has also created FMLA forms for Eligibility and Designation Notices.

How much leave time must be given and for what reasons?
The FMLA and VPFLA allow covered employees to take up to twelve weeks of unpaid leave in a twelve-month period for their own serious health condition, the serious health condition of an immediate family member, the birth and care of the employee’s newborn child, or for the adoption or foster care placement of a child. The VPFLA law uses the term serious illness and defines it as any condition that poses an imminent danger of death, requires inpatient care in a hospital, or requires continuing in-home care under direction of a physician. The VPFLA also limits parental leave to the initial placement of a child 16 years of age or younger for adoption. Vermont law defines immediate family members as the employee’s child, stepchild or ward who lives with the employee, foster child, parent, spouse (including civil union partner), or parent of the employee’s spouse or civil union partner. The employee may use any earned, accrued paid time during the 12 weeks leave, but she/he cannot use the paid leave to extend the leave beyond 12 weeks. The employee does not have to use any paid leave and may elect to take the entire 12 weeks unpaid. An employer can cap the total amount of paid time off an employee can use during VPFLA leave to 6 weeks.

Employees do not need to use their leave entitlement in one block. If medically necessary, FMLA/VPFLA leave for a serious health condition may be taken intermittently (in separate blocks of time due to a serious health condition) or on a reduced leave schedule (reducing the usual number of hours you work per workweek or workday). FMLA leave may also be taken intermittently or on a reduced leave schedule for a qualifying exigency relating to covered military service.

The FMLA also includes a special leave entitlement that permits eligible employees to take up to 26 weeks of leave to care for a covered service member during a single 12-month period.

What happens when the employee returns from leave?
At the conclusion of FMLA or VPFLA leave, an employee must generally be reinstated to the same or an equivalent position with equivalent benefits, pay, and other terms and conditions of employment. In other words, an employee generally cannot be permanently replaced while on VPFLA or FMLA leave. Also, during the leave, the employer must continue the employee’s group health insurance as if the employee remained continuously employed, and leave cannot result in the loss of an employment benefit that accrued prior to the start of the employee’s leave.

What is short-term family leave?
Vermont law also entitles eligible employees to an additional 24 hours of leave in any 12-month period, but limited to four hours in any 30-day period, for short-term family leave which can be used to participate in academic activities at their child’s school, attend routine medical appointments with their child or parent, or similar matters that might not qualify for FMLA. The employee is expected to make a reasonable attempt to schedule the appointments outside of work time and to give the earliest possible notice of the need for leave.

**DOCUMENTS AND RECORD-KEEPING**

*What documents must a new employee complete?*

After an employee is hired (but not before), the employee must complete a form I-9 ([https://www.uscis.gov/i-9](https://www.uscis.gov/i-9)), and the employer must examine evidence of the employee’s identity and employment eligibility, as specified on the form I-9, on the employee’s first day of employment. The employee must also fill out Form W-4 ([https://www.irs.gov/pub/irs-pdf/fw4.pdf](https://www.irs.gov/pub/irs-pdf/fw4.pdf)) for federal income tax withholding. Any insurance forms should also be completed, as well as emergency notification information. In Vermont, new hire reporting is mandatory, and more information is available at the Vermont Department of Labor’s website. ([http://labor.vermont.gov/unemployment-insurance/employers/new-hire-reporting/](http://labor.vermont.gov/unemployment-insurance/employers/new-hire-reporting/))

In addition to documents that must be completed by an employee, all employers have workplace posting requirements. Many state and federal laws require employers to notify employees of their rights under such laws through the conspicuous display of posters in the worksite.

*What records should be in an employee’s personnel file?*

The following types of documents are generally maintained in an employee’s personnel file: employee contact and emergency contact information, employment application, resume, offer or hiring letter, salary or other compensation information, including information confirming pay changes, records related to promotion, demotion, transfer, or reporting structure, performance reviews, documentation of disciplinary action, signed acknowledgment(s) of receipt of: the employee handbook or other company policies or procedures, job descriptions, any other signed agreements with the employee, including non-competes, non-disclosure (confidentiality), non-solicitation agreements, if any. Some employers also choose to keep other documents with sensitive employee information, such as the employee’s social security number, separate from the employee’s main personnel file.

Any information concerning an employee’s health or medical conditions must be kept in confidential files, separate from other personnel documents. Otherwise, Vermont has no law governing what documents must be or may not be in an employee’s personnel file. Additionally, there is currently no Vermont law requiring a private employer to provide an employee with access to his/her personnel file.

*What records must be kept and for how long?*

Numerous state and federal laws require retention of specific employment records, but the length of retention varies. Most financial records, particularly those to support federal income tax filings, should be kept for at least seven years after the relevant return is filed. Employee
benefits documents under ERISA must be kept for at least six years after the plan ends. Vermont’s unemployment law requires payroll and work hour records to be kept for four years, and various employment laws, including the FLSA, require retention for at least three years after an employee leaves (or as long as any legal proceeding involving them continues). OSHA requires that documents related to certain toxic exposures be kept for the duration of employment plus thirty years. Applications, even of unsuccessful candidates, must be kept for at least one year after the hiring decision is made.

Generally, employers should consider keeping all employee personnel records (except records relating to exposure to toxic substances) for seven years after termination of employment. At the end of whatever time is chosen, records should be destroyed pursuant to an established document-destruction plan.

**Must an employer have an employment handbook?**

Vermont law requires that every Vermont employer have one policy: a sexual harassment policy. A sexual harassment policy must include at least all information required under Vermont law. 21 V.S.A. § 495h. These policies do not have to be in a handbook per se, but they do have to be written and disseminated (and the sexual harassment policy must be posted). For more information, see the section in this chapter on Unlawful Harassment.

There are pros and cons to having a handbook with more than these policies. However, every employer has policies, even if they are not written down. Those policies should be applied consistently and fairly, and putting them in writing may make it easier to do so. If an employer has a handbook, it should be clear and simple. It should say what the employer means, and the employer should really mean everything it says. Which policies to include and the extent of detail in them should reflect the employer’s values, style, and culture. The contents of a handbook must be compliant with state and federal law and may be used to argue a contractual obligation has been created on the part of the employer, so any handbook should be reviewed by the organization’s attorney.

Vermont employers must prohibit smoking in the workplace.

**What policies should an employer consider including in a handbook?**

Policies should address the employer’s expectations for employees - behavior, attendance, meal and rest breaks, personal use of equipment, drugs and alcohol, anti-discrimination/harassment/retaliation, etc. They should also provide information employees need to know about administrative matters - benefits information, payroll practices, filing harassment complaints, overtime rules, expense reimbursement policies, etc. Consequences for an employee’s breach of policy or failure to perform their job are often addressed. Leave time, particularly vacation, Earned Sick Leave, and unpaid family leave issues, are important topics, as there are certain choices an employer can make on such policies. Although a confidentiality provision is often included in a personnel handbook, most operational issues (how to do proper billing, what to put in a client’s chart, how to answer the phone, etc.) belong elsewhere.

**ON THE JOB**
What should an employer do to supervise and evaluate employees?

Each workplace has its own culture about how supervision and evaluation is carried out, ranging from formal to informal. When a problem arises, however, it is critical that the employer have documentation of all corrective action it has taken or attempts it has made to improve the employee’s performance. Best practice suggests that employers provide informal supervision on an ongoing basis but that formal supervision or evaluations occur at least annually. Evaluations should be as objective as possible and based on the employee’s job description. If an employee has worked for an employer for several years, there is no documentation that any problems were ever discussed, and the employee is then terminated, the employer may have difficulty defending the performance-based reasons for its decision if the employee asserts the firing was for an unlawful reason. Documentation can be as simple as handwritten notes in a file, or as formal as a typed evaluation form. Above all, evaluations and supervision should be honest and not ignore or minimize problems or concerns.

What must an employer do to protect an employee’s privacy?

As an employer, a physician practice may be subject to the privacy requirements of HIPAA, if it provides any health services to the employee (e.g., free screenings to employees), if it is self-insured (even partially) for health care, if it offers a medical reimbursement plan, or if it receives an employee’s protected health information (PHI) for any insurance-related purpose. An employer is not subject to HIPAA merely because it has information about an employee’s health for employment purposes, such as determining sick leave, family leave, workers’ compensation, or disability accommodations. Therefore, each employer must determine if it is subject to the HIPAA privacy regulations related to records created or maintained for employment-related purposes. If it is, the employer must have a HIPAA policy (for employee records), must enact appropriate procedures to ensure that none of the PHI is used for employment purposes, must appoint a privacy officer, and take other steps to comply with the regulations.

Employees also have a common law right to expect privacy in certain areas, such as on their person, in bathrooms, and in changing areas. The extent of the privacy right depends on the situation and circumstances, and employers can take steps to limit the expectation. For example, a private employer may notify employees that it will monitor email or Internet use, monitor telephone calls, or search employees’ workstations or handbags, as long as the notification is clear and the practice is limited to business-related matters.

When and how can an employer monitor an employee’s electronic communications?

The Federal Wiretapping Act, as amended by the Electronic Communications Privacy Act, generally makes it criminal for anyone to intercept anyone else’s telephone or electronic communications. The Act contains an exception that allows employers to intercept and monitor the communications of employees on its premises for work-related purposes. An employer may monitor an employee’s personal call only so long as needed to determine the call is personal, or if the employee consents. Additionally, anyone may record or intercept any communication if one party to the communication consents (i.e., a person may record a conversation she/he is involved in, even without notice to the other parties). Some states prohibit this kind of interception without consent (so interstate calls may raise problems), but Vermont does not. A written policy clearly stating what communications will be monitored and obtaining employee
acknowledgment or consent is advisable before an employee’s electronic communications are monitored.

The Stored Communications Act covers stored communications, such as employee websites or e-mails stored on a server. (18 U.S.C. §§ 2701-12). The Stored Communications Act does not prevent an employer from reviewing communications stored on employer-provided wire or electronic communications services, if the review is authorized by the employer’s policies.

There are certain circumstances where employers are prohibited from surveillance. For example, under the federal National Labor Relations Act, which applies to all employers, employers must not conduct surveillance of employees engaging in union organizing activities. Employers also must not target employees for surveillance in an unlawfully discriminatory manner.

What is an employer’s obligation to employee health and safety?
Both federal and state laws require that employers maintain a safe and healthful working environment. To that end, VOSHA provides technical assistance or has the right to inspect a workplace to ensure that employers are following state and federal regulations.

When is an employer liable for an employee’s acts?
Generally, an employer can be held liable by a third party for negligent acts committed by its employees within the scope of their employment. Sometimes employers are even liable for the intentional or reckless acts of employees, but usually only if the employer was aware of the potential that the employee would act as she/he did and the scope of the employee’s employment allowed the employee to engage in that act. Employers generally are not liable for the acts of their employees outside the scope of their employment.

What must an employer do to accommodate an employee with a disability?
The federal Americans with Disabilities Act of 1990 (ADA), as amended by the ADA Amendments Act of 2008 and the Vermont FEPA both require an employer both refrain from discriminating against qualified individuals because of a disability and provide qualified individuals with a disability a reasonable accommodation, unless doing so would cause an undue hardship. All employers are covered by Vermont’s FEPA and a private employer is covered under the ADA if it has 15 or more employees on its payroll for 20 or more calendar workweeks (which do not need to be consecutive) in either the current or preceding calendar year. 29 C.F.R. § 1630.2(e).

A reasonable accommodation is a change in the work, workplace, or application process that helps make it possible for an individual with a disability to perform the duties of or apply for a job, or enjoy the benefits and privileges of employment. An employer does not have to provide the exact accommodation the employee or applicant wants, if more than one accommodation would be effective.

Under the ADA, the term “qualified,” with respect to an individual with a disability, means that the individual satisfies the requisite skill, experience, education and other job-related requirements of the employment position such individual holds or desires and, with or without
reasonable accommodation, can perform the essential functions of such position. 29 C.F.R. § 1630.2(m).

An undue hardship means the accommodation would cause significant difficulty or expense for an employer, in light of the employer’s size, financial resources, and the needs of the business. It is not an undue hardship just because an accommodation request involves some cost to the employer.

When does an employee have a disability?
According to the Americans with Disabilities Act and the Vermont Fair Employment Practices Act, a disability is a physical or mental impairment that substantially limits one or more major life activities (such as walking, talking, seeing, hearing, learning, or working). 42 U.S.C. § 12102(1); 29 C.F.R. § 1630.2(g); 21 V.S.A. § 495d(5). The ADA and VT FEPA also prohibit discrimination against individuals who have a history or record of a disability (e.g., cancer that is in remission) or who are regarded as having a disability (e.g., the employer believes the employee is disabled). The ADA further prohibits discrimination against individuals who have a relationship with or are associated with a person with a disability.

A physical impairment is further defined as:
- any physiological disorder, or condition, cosmetic disfigurement or anatomical loss affecting one or more of the following body systems: neurological, musculoskeletal, special sense organs, respiratory (including speech organs), cardiovascular, reproductive, digestive, genitourinary, hemic, lymphatic, skin, endocrine, immune, or circulatory. 29 C.F.R. § 1630.2(h)(1).

A mental impairment is defined as “any mental or psychological disorder, such as intellectual disability, organic brain syndrome, emotional or mental illness, and specific learning disabilities.” 29 C.F.R. § 1630.2(h)(2).

Under the ADA, the definition of disability is to be interpreted in favor of broad coverage. Nonetheless, not every impairment will constitute a disability within the meaning of the ADA. However, the term “substantially limits” shall be construed broadly in favor of expansive coverage, to the maximum extent permitted by the terms of the ADA. “Substantially limits” is not meant to be a demanding standard. 29 C.F.R. § 1630.2(j)(1)(i). An impairment is a disability within the meaning of the ADA if it substantially limits the ability of an individual to perform a major life activity as compared to most people in the general population. An impairment does not need to prevent, or significantly or severely restrict, an individual from performing a major life activity in order to be considered substantially limiting. 29 C.F.R. § 1630.2(j)(1)(ii).

Under ADA regulations, the following (non-exhaustive) list includes impairments that should be “easily” be found to be disabilities: deafness, blindness, an intellectual disability, partially or completely missing limbs or mobility impairments requiring the use of a wheelchair, autism, cancer, cerebral palsy, diabetes, epilepsy, human immunodeficiency virus (HIV), multiple sclerosis, muscular dystrophy, major depressive disorder, bipolar disorder, post-traumatic stress disorder, obsessive compulsive disorder, and schizophrenia. 29 C.F.R. § 1630.2(j)(3)(iii).
Mitigating or corrective measures (such as hearing aids or medications) must not be considered in the analysis of whether an individual has a disability.

Alcoholism may meet the definition of a disability, but the ADA specifically provides that employers may prohibit the use of alcohol in the workplace and may require an employee who is an alcoholic or who engages in the illegal use of drugs to meet the same performance and behavior standards of other employees. 42 U.S.C. § 12114 (c)(4). The ADA does not protect an individual who currently engages in the illegal use of drugs, but it may protect a recovered drug addict who is no longer engaging in the illegal use of drugs, who is qualified and can meet the definition of disability under the ADA. 42 U.S.C. § 12210.

What documentation can an employer require to determine if an employee has a disability?
After a request for a reasonable accommodation has been made, and whether the need for accommodation is not obvious, an employer may request that the employee provide reasonable documentation from a health care provider confirming the existence of a disability, the employee’s job-related functional limitations, and the need for a reasonable accommodation. An employer must be careful to avoid seeking more information than is necessary to make those determinations. For example, an employer can only seek documentation pertaining to the disability that requires a reasonable accommodation.

What must an employer do if an employee requests an accommodation?
Generally, a request for an accommodation comes from the disabled individual. Alternatively, a request may come from a third party asking for an accommodation on behalf of the covered individual. Requests can be made verbally or in writing, and the individual does not specifically have to say the word “accommodation.” An employer cannot obligate that an employee’s request take a certain form before considering it. EEOC guidance also suggest that employers should be offered or provided without request in situations where the employer knows the employee has a disability and is experiencing workplace problems because of the disability.

When an individual requests a reasonable accommodation for a disability, the employer must promptly engage in an “interactive process” in good faith to determine what limitation are created by the disability and to explore potential, appropriate and reasonable accommodation(s).
When engaging in the interactive process, an employer should do the following:

- Document the individual’s accommodation request;
- Determine (if not known or obvious) that the individual has a disability;
- Ask the individual to provide relevant information, if necessary;
- Meet with the individual to discuss job-related limitations, the accommodation request, and possible alternatives; and
- Document the discussion about the accommodation and the determination on the request.

The ADA and Vermont law require an employer to provide a reasonable accommodation if the employee (or applicant) is otherwise qualified for the job and if the employee needs the accommodation to perform the essential functions of his/her job. A reasonable accommodation means any modification or adjustment to the job or work environment. The essential functions of a job are key to any reasonable accommodation analysis under the ADA. If an employee cannot perform one or more of the essential functions of the job she/he was hired to do, and there
is no reasonable accommodation that can enable him/her to perform that function, the employee is not qualified to do the job. In such cases, an employer must examine whether the employee can be reassigned to a vacant position as a reasonable accommodation.

An employee may request that marginal or non-essential duties of a position (for example, a function that can be performed by another employee or a function that is only occasional) be reallocated or redistributed as a reasonable accommodation.

The laws do not define, in relation to any particular job, what is reasonable and what is not. That determination is made on a case-by-case basis, although employers do not have to change the essential functions or nature of the job to accommodate an employee. Also, the duty to provide an accommodation is ongoing - an individual can make more than one accommodation request and may modify a request if circumstances change.

What are examples of reasonable accommodations?
Examples of reasonable accommodation include: making existing facilities used by employees readily accessible to and usable by an individual with a disability; restructuring a job; modifying work schedules; acquiring or modifying equipment; providing qualified readers or interpreters; or appropriately modifying examinations, training, or other programs; providing a leave of absence; allowing the use of reserved parking spaces, etc. Reassigning an employee to a vacant position may be a reasonable accommodation if it is requested or there are no accommodations that would keep the employee in his/her current position.

An employer is not required to provide personal use items such as glasses, wheelchairs, or hearing aids unless the employee only needs them to perform a specific function at work and in no other situation. It is not a reasonable accommodation to eliminate or reassign an essential function of the job.

What can an employer do if an employee presents a health or safety risk?
An employer does not have an obligation to accommodate an employee who poses a direct threat to the safety of him/herself or others. A “direct threat” means “a significant risk of substantial harm” that cannot be reduced or eliminated by any reasonable accommodation. The risk of harm must not be speculative but must be reasonable and based on current, objective medical knowledge and/or evidence.

**PERFORMANCE MANAGEMENT AND END OF EMPLOYMENT**

*When can an employee be disciplined or terminated?*
In Vermont, an at-will employee may be disciplined or terminated at any time, with or without notice, and for any reason or for no reason at all, as long as it is not an unlawful reason. Having said that, there are several specific laws that define unlawful reasons for termination or that govern an employer’s behavior towards an employee, many of which have been addressed in this Guide. For example, common law may provide an employee with a civil remedy if an employer denies an employee any protection or benefit the employer has promised and on which the employee has detrimentally relied, defames an employee or former employee, intentionally
inflicts emotional distress on him/her, or intentionally interferes with the employee’s contractual relations with others.

What are unlawful reasons to discipline or terminate an employee?
There are numerous unlawful reasons to terminate an employee, but some of the most common follow (this list is not all-inclusive):

- because of the employee’s race, color, sex, gender, gender identity, age, religion, national origin, ancestry, place of birth, ethnicity, pregnancy, sexual orientation, disability, physical or mental condition, genetic information, HIV-positive status, military service or obligation, veteran status, citizenship, or immigration status;
- because the employee has disclosed the amount of his/her wages or inquired about or discussed the wages of other employees;
- because the employee has complained about discrimination, filed a charge of discrimination, or cooperated in an employment discrimination investigation or lawsuit or because the employer believes that the employee may lodge a complaint or cooperate with officials in an employee discrimination investigation;
- because the employee filed a workers’ compensation or VOSHA claim;
- because the employee has exercised or attempted to exercise his/her rights to medical, parental or family leave under the law, if applicable;
- because the employee missed work to serve on a jury;
- because the employee reported a health or safety violation;
- because the employee attempted to unionize other employees or engaged in other protected concerted activities for employees’ mutual aid or protection;
- because the employee refuses to take a polygraph test (except where permitted by law); or
- because the employee engaged in activities protected by public policy.

Employees of hospitals and nursing homes also have whistleblower protection. 21 V.S.A. §507.

What is discrimination?
Discrimination is an action that adversely affects an employee in any aspect of the terms, conditions, or privileges of employment, such as hiring, firing, compensation, transfer, promotion, testing, recruiting, or receipt of benefits. Treating one employee or one class of employees differently from others because of one or more legally protected characteristics is unlawful discrimination.

What kind of discrimination is prohibited?
The Vermont Fair Employment Practices Act (21 V.S.A. § 495) applies to all Vermont employers, regardless of size, and prohibits discrimination on the basis of race, color, religion, national origin, sex, sexual orientation, gender identity, ancestry, place of birth, age (18 and up), physical or mental condition, or HIV-positive status.

There are many state and federal laws that prohibit discrimination in employment based on various additional protected characteristics, each of which has a different threshold level of employees for coverage (some of these characteristics are included under the unlawful reasons to
terminate employee, set forth above).

What is unlawful harassment?
Harassment is a form of unlawful employment discrimination. Harassment is unwelcome conduct based on a legally protected characteristic, where 1) enduring the offensive conduct becomes a condition of continued employment; or 2) the conduct is severe or pervasive enough to create a work environment that a reasonable person would consider intimidating, hostile, or abusive. Employers should take appropriate steps to correct and prevent unlawful harassment of its employees. Any employee who is affected by the offensive conduct can be the victim of harassment, even if he or she is not the person being harassed.

Vermont law specifically defines sexual harassment as unwelcome sexual advances, requests for sexual favors, and other verbal or physical conduct of a sexual nature when:
- submission to that conduct is made either explicitly or implicitly a term or condition of employment; or
- submission to or rejection of such conduct by an individual is used as a component of the basis for employment decisions affecting that individual; or
- the conduct has the purpose or effect of substantially interfering with an individual's work performance or creating an intimidating, hostile or offensive work environment. 21 V.S.A. § 495d(13).

Vermont law requires every employer to have a sexual harassment policy, which shall include:
- a statement that sexual harassment in the workplace is unlawful;
- a statement that it is unlawful to retaliate against an employee for filing a complaint of sexual harassment or for cooperating in an investigation of sexual harassment;
- a description and examples of sexual harassment;
- a statement of the range of consequences for employees who commit sexual harassment;
- if the employer has more than five employees, a description of the process for filing internal complaints about sexual harassment and the names, addresses, and telephone numbers of the person or persons to whom complaints should be made; and
- the complaint process of the appropriate state and federal employment discrimination enforcement agencies, and directions as to how to contact such agencies. 21 V.S.A. § 495h(b).

A poster containing the policy must be in every workplace, and each employee must receive an individual written copy.

In addition, both federal and Vermont law require an employer to take prompt, remedial action reasonably calculated to end the harassment whenever the employer receives a report of unlawful harassment. The first step in any response is to investigate and document the allegations. What other action to take will depend on the nature of the conduct at issue and the relationship of the perpetrator to the victim. An employer will be held liable for a supervisor’s sexual harassment, where that supervisor has taken a tangible employment action against the victim. Conduct by a co-worker may involve liability to the employer if the employer has knowledge of the harassment and fails to take appropriate and prompt remedial action.
What can an employer do to reduce exposure to claims for harassment or discrimination?

First, all changes in an employee’s terms and conditions of employment should be based on legitimate performance reasons or be consistent with the business needs and mission of the organization. Second, employee performance problems should be addressed promptly and honestly, rather than being tolerated, downplayed, or ignored. Third, if an employer learns that an employee feels she/he is being discriminated against or harassed, the employer should document the concerns, immediately investigate them, and take prompt action to try to end the behavior. An employer must promptly investigate allegations of unlawful discrimination or harassment, even if the employee or witness refuses to submit a complaint in writing or the employee asks that the matter be kept confidential. While an employer should keep employee complaints as confidential as possible, complete confidentiality should never be promised because some information must be disclosed when investigating a complaint in order to gather relevant information. Fourth, the employer should conduct periodic training of employees to ensure that they are aware of the laws affecting the workplace and that their conduct conforms with them. Fifth, employment policies should be clear and consistently followed.

What laws govern reductions in force or mass layoffs of employees?

The federal law specifically governing layoffs is the Worker Adjustment and Retraining Act (the “WARN Act”), which applies to organizations with 100 or more employees. The WARN Act requires notification 60 days before a facility or operating unit of a business is closed or a mass layoff is held that involves employment loss for 50 or more employees.

Vermont also has the Notice of Potential Layoffs Act (“NPLA”) 21 V.S.A. §§ 411-418, which applies to employers with 50 or more employees. The NPLA generally requires employers to provide 45 days’ notice to the State government and 30 days’ notice to employees and local government in advance of covered mass layoffs. Notice obligations are triggered when there is to be a mass layoff or business closing that will result in the permanent employment loss of at least 50 employees at one or more worksites in Vermont during any 90-day period. An employer who fails to provide required notice may be liable for administrative penalties as well as severance pay and health benefit continuation for affected employees.

To ensure compliance with both the state and federal layoff notification laws, potentially covered employers should consult with counsel in advance of implementing any layoffs, reductions in force or plant closings.

What is unemployment compensation?

Under Vermont and federal laws, employees who lose their jobs, generally through no fault of their own, are entitled to unemployment compensation during periods between jobs. The unemployment compensation fund is funded entirely by employer taxes, and the amount of tax is based on the employer’s experience rating - that is, the number of claims attributable to it within the relevant period. Non-profit 501(c)(3) organizations may elect not to contribute (and pay any claims directly themselves).

A former employee will generally be eligible for unemployment benefits if the separation is through no fault of their own and he/she is willing and able to work. There are certain circumstances where a claimant may be partially or totally disqualified from unemployment
benefits in circumstances of misconduct or gross misconduct, as these terms are narrowly defined by Vermont unemployment compensation law. The Vermont Department of Labor’s unemployment office particularly looks to find out if the employee has received notice and warning about the consequences of his/her behavior, as well as its severity, before denying benefits. A claimant may also be denied benefits if the employee quits (leaves voluntarily without good cause attributable to the employer). For more information, see the Vermont Department of Labor’s unemployment insurance webpage.

What can/should an employer say in giving a reference?
Employers must be careful in giving references for former employees, as defamation, retaliation and tortious interference with contract claims have been made against employers for making false statements that damage a person’s reputation or ability to find another job. On the other hand, employers have been sued by a person’s new employer because they have given a falsely positive reference that fraudulently misrepresented the former employee in an overly positive light. Many of the legal cases have involved extreme situations, such as accusing an employee of stealing when the employer had no evidence the employee stole (and particularly if the employee is later able to prove that she/he did not) or giving a glowing reference for an employee who was terminated for sexual assault. To minimize exposure to claims, many employers refuse to give references for anyone (there is no legal requirement to provide a reference) or choose to give only limited information (e.g., to confirm solely dates of employment, positions held). Still others give a reference only after receiving a written release from the former employee to do so. Whatever the approach, employers should consistently follow the same practice when providing references.

What are the requirements for continuation of health benefits for employees who leave employment?
The Consolidated Omnibus Budget Reconciliation Act (COBRA) requires group health plans offered by employers with 20 or more employees to offer continuation coverage for employees or their dependents who experience a “qualifying event” and requires employers to give notice of COBRA rights. A qualifying event may be termination of employment, a reduction in hours, divorce, or similar acts that result in a loss of group health benefit coverage. The continuation of coverage may require that the employee pay the cost of the coverage and may continue for up to 18 months (36 months for certain qualifying events) from the date the health plan would have ended. If the beneficiary does not pay the premium, the employer ceases to offer any group health plan, or the employee becomes eligible for another health insurance plan or Medicare, the continuation coverage ends. When the employee leaves employment, the employer or insurer must send the employee a notice to elect continuation coverage.

Vermont law requires all group health insurance plans, even those for employers with fewer than 20 employees, to provide 18 months of continuation coverage. 8 V.S.A. § 4090a, et. seq. The standards are similar to COBRA, and an employer must send notice of the right to elect continuation of coverage to the employee (or former employee) within 30 days following the occurrence of any qualifying event. 8 V.S.A. § 4090a(e).

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FRAUD AND ABUSE COMPLIANCE

Topics Covered in this Chapter:
Introduction
The Federal False Claims Act
The Anti-Kickback Statute
The Physician Self-Referral Ban (The Stark law)
Other Federal Laws
Vermont Laws
Preventative Action Compliance Programs
Conclusion
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INTRODUCTION

The regulatory enforcement atmosphere for health care providers has been steadily intensifying since the early 1990s, roughly paralleling the growth in the nation’s government-funded health care programs. Indeed, passage of major health care reform initiatives in 2009 and 2010 brought with it expanded liabilities and penalties for fraud, enhanced screening of providers and suppliers, and additional funding for regulatory enforcement.

Under the best circumstances, it is sometimes difficult for providers (and even regulatory professionals!) to predict what types of arrangements and practices might constitute fraud or unlawful conduct under the various enforcement statutes. And penalties for unlawful conduct can be substantial, including criminal fines (or even imprisonment), civil monetary penalties, and exclusion from Medicare, Medicaid, and other federal programs.

All this means that providers are finding it absolutely essential to become familiar with the basic tenants of fraud and abuse laws to understand the potential pitfalls associated with their business activities. This section briefly outlines the primary enforcement tools used to root out and combat fraud in the health care system, including the federal False Claims Act, the federal health care program “Anti-Kickback law,” the physician self-referral or “Stark law”, the Vermont False Claims Act and the state Medicaid fraud law. In addition to these primary enforcement mechanisms, this section also discusses a number of other statutes that may be implicated by conduct triggering the primary fraud and abuse laws.

THE FEDERAL FALSE CLAIMS ACT

The federal False Claims Act (FCA), 31 U.S.C. §§ 3729-3733, was first enacted in 1863 to combat defense procurement fraud during the Civil War. Since then, the FCA has been amended a number of times, and in 1986, as the federal government continued to take on more responsibility for health care, Congress made changes that shifted the primary focus of the law from the defense industry to health care. In 2009, the statute was substantially amended by the
Fraud Enforcement and Recovery Act (FERA) as a part of broader health care reform efforts. Today, the federal government recovers billions of dollars each year from companies and individuals in the health care industry through FCA enforcement actions and settlements.

In the most general terms, the FCA creates stiff penalties for any person who “knowingly presents, or causes to be presented, a false or fraudulent claim for payment or approval” to the federal government. In a provision commonly known as the “reverse” false claims section, the law also prohibits actions to improperly avoid paying money owed to the government.

In the health care industry, FCA liability arises most commonly out of the miscoding of or misrepresentation in ICD codes, improper billing for tests not performed, or misrepresentations involving places or providers of services. And importantly, certain conduct that violates the anti-kickback and Stark laws (see below for more on these laws) may also violate the FCA.

**Key Elements of the FCA**

Under the FCA as amended by the FERA, a “claim” is “any request or demand, whether under a contract or otherwise, for money or property” made to the U.S. government or certain government contractors (including contractors implementing Medicare Part C and D plans). A claim must be “false” to be covered by the FCA, and falsity may take the form of an express claim that is false, or an express or implied certification that is false. False claims must also be “material,” which means the claim must have had a “natural tendency to influence, or [must have been] capable of influencing, the payment or receipt of money or property.”

The FCA primarily prohibits a person from “knowingly” presenting (or causing someone else to present) a false or fraudulent “claim” for payment to the U.S. government. The statute itself defines “knowingly” as (1) having actual knowledge of the falsity of the claim; (2) deliberate ignorance of the truth or falsity of the claim; or (3) reckless disregard of the truth or falsity of the claim. That means that a person need not specifically intend to defraud the government in order to be held liable; rather the FCA prohibits all claims that a person knows or should know are false.

In determining whether a claim is submitted “knowingly,” courts and the U.S. Department of Justice (which is responsible for enforcing the FCA) may ask the following questions:

- Did the provider have either actual or constructive knowledge of the rules or policies governing the submission of the particular claim? Were there Medicare or Medicaid bulletins or advisories on point?
- How clear is the policy that was violated? Is there room for reasonable mistake?
- How many false claims were submitted? Reckless indifference can be inferred from a large number of claims.
- Does the practitioner or provider have a compliance plan, and is it being followed?
- Does the practitioner have a record of reporting suspected compliance problems and attempting to fix them?
- Did the provider seek guidance on billing issues from CMS or a third-party carrier, and was any guidance given clear?
Penalties and Damages
Penalties and damages in a successful FCA case can be severe. As of August 1, 2016, civil penalties under the FCA dramatically increased from up to $11,000 ($5,500 minimum) per claim to a maximum of $21,563 (and at least $10,781) for every false claim.

The civil penalties apply whether or not the government actually suffered a financial loss. If the government did suffer such a loss, it is also entitled to up to three times the amount of damages it suffered, or twice the amount if the provider self-reported and was cooperative during the enforcement action.

Finally, in the event a successful FCA case is brought by a private whistleblower, the provider may also be responsible for paying the whistleblower’s costs and attorneys’ fees.

Whistleblowers and “Qui Tam” Actions
A qui tam action is a mechanism through which a private party may sue in the name of the government to enforce its rights. In the context of the FCA, this means that a private party – frequently a whistleblower – may sue a provider on behalf of the federal government and receive a portion of any judgment or settlement that results. In such cases, the private party bringing suit is known as the “relator.”

By design, relators are provided with a substantial financial incentive to bring FCA cases. If the government decides not to participate in the suit (in every case, the United States Attorney General must be given notice and an opportunity to take over the case), the relator is entitled to 25-30 percent of any amount recovered. And even if the government does decide to handle the case, the relator is still entitled to 15-25 percent.

In a qui tam action, a health care whistleblower may include a count alleging retaliation by the provider. If successful, the whistleblower may be entitled to reinstatement (if he or she was fired as a result of his or her whistleblowing activity), awarded twice his or her backpay plus interest, compensation for any special damages, attorney’s fees, and the costs of bringing suit.

Health care whistleblowers are also protected by Vermont’s Healthcare Whistleblower’s Protection Act, 21 V.S.A. §§ 507-509, and can also pursue claims under Vermont’s False Claims Act, 32 V.S.A. §§ 630 et seq., both of which are discussed later on in these materials.

Steps to Reduce Risk of FCA Violations
The primary way for providers to reduce FCA risk is to ensure that employees and business colleagues understand the law’s basic requirements and know that the provider takes FCA and fraud and abuse issues seriously. The following are a few additional, specific steps providers can take to help minimize FCA risk:

- Ensure that corporate policies clearly spell out FCA compliance and reporting requirements. Make sure that the consequences of failing to report fraud are clearly communicated and understood. Policies should also protect those who come forward with billing questions, complaints or concerns.
- Conduct periodic trainings on FCA compliance.
• Maintain clear personnel policies and meticulous personnel records. This is important as a means to identify and prevent employees from committing health care fraud, and to help defend against claims of retaliation.
• Have a system of review in place for claims and invoices submitted to the government. Try to spot billing errors or regulatory compliance issues before they are submitted.
• Regularly engage employees in areas of potential FCA problems. Solicit open communication about potential problems, and again, make sure that those who come forward are adequately protected.
• Have a plan in place to deal with allegations of fraud. Ensure prompt investigatory and remedial action.
• Consider self-reporting any problems to potentially avoid larger exposure. Always consult with an attorney first, though.

THE ANTI-KICKBACK STATUTE

The Federal Health Care Program Anti-Kickback Statute (its full name), 42 U.C.S. § 1320a-7b, creates criminal liability for individuals and entities that “knowingly and willfully” solicit, receive, offer or pay anything of value to induce referrals of items or services paid for by a federal health care program (including Medicare and Medicaid). The statute is designed primarily to prevent kickbacks, bribes and rebates, as well as similar arrangements with a potential to improperly increases costs or decrease efficiencies in the federal health care benefit programs.

“Knowing and Willful”

To be held liable under the Anti-Kickback Statute, an individual or entity must “knowingly and willfully” enter into a prohibited arrangement. The precise meaning of “knowingly and willfully” has been the subject of some debate, but in general, it must be established that at least one purpose of the challenged arrangement must have been to unlawfully induce referrals of items or services paid for by a federal program.

In a 2010 amendment to the Anti-Kickback law, Congress clarified that a person or entity need not have actual knowledge of the law or specific intent to violate it – a construction some courts had previously placed on the law that had made it difficult for criminal prosecutions to be successful. Rather, the prosecution must show only that the person or entity knew that its conduct was generally illegal.

Exceptions to the Anti-Kickback law

Congress has expressly recognized that the Anti-Kickback law’s language is quite broad and that the potential for confusion and unfair application of the statute is great. Therefore, the law and its regulations include a set of statutory exceptions and regulatory “safe harbors” for a variety of business arrangements. If a particular arrangement meets the requirements of one of the safe harbors, it will be presumed not to violate the statute and will not be subject to an enforcement action by the agency responsible for administering the law, the U.S. Department of Health & Human Services.
It is difficult to provide a concise description of all the specific requirements for an arrangement to fall within an exception or safe harbor. In general, though, the exceptions and safe harbors encompass the following types of arrangements:

**Statutory Exceptions**
- Payments to bona fide employees
- Discounts
- Payments to purchasing agents
- Some types of risk-sharing arrangements
- Prescription drug discounts for certain beneficiaries in the Medicare coverage gap (or “donut hole”)

**Safe Harbors**
- Investment interests in certain entities
- Certain space, equipment, personal services and management contracts
- Sale of a practice
- Legitimate referral services
- Warranties (so long as certain reporting requirements are met)
- Group purchasing organizations
- Certain Medicare Part A waivers of coinsurance and deductibles
- Beneficiary incentives offered by group health plans and price reductions offered to such plans
- Certain efforts to recruit physicians to underserved areas
- Investment interests in group practices by physicians
- Obstetrical malpractice insurance subsidies
- Investment interests in surgeon-owned ambulatory surgical centers
- Cooperative hospital service organizations
- Referral agreements for specialty services
- Ambulance replenishing arrangements
- Certain arrangements involving electronic prescribing and electronic medical records
- Certain transfers of goods and services in the context of federally qualified health centers

To fit within any of the Anti-Kickback safe harbors, a person must satisfy certain specific criteria that are set forth in the applicable statutory or regulatory provisions. Notably, a business arrangement that does not fall within a safe harbor is not automatically illegal. Rather, such arrangements will be evaluated on a case by case basis by the Department of Health & Human Services Office of Inspector General (OIG).

**Penalties**
Violation of the Anti-Kickback law is a federal felony, punishable by up to five years of imprisonment and/or fines up to $25,000. Moreover, a violation of the law can result in exclusion from participation in the federal health care programs. Discipline imposed under the law may also be a basis for state unprofessional conduct charges and license suspension.
Relationship of Anti-Kickback law with the False Claims Act

As part of health care reform efforts in 2010, Congress made clear that a violation of the Anti-Kickback law is also a violation of the federal False Claims Act. Thus, although the Anti-Kickback law does not itself provide for a private cause of action for its violation, civil liability still could be established through either a direct or qui tam action under the FCA.

Additional Guidance for Anti-Kickback Statute Compliance

Under the Anti-Kickback statute, OIG is authorized to issue advisory opinions assessing proposed business arrangements for compliance with the law. 42 C.F.R. §§ 1008.5(a)(1)-(3). The opinions may address (1) whether the arrangement constitutes a prohibited referral and/or (2) whether the arrangement fits within a safe harbor.

Previous advisory opinions are available to the public on OIG’s website and are helpful for understanding how OIG might view a particular arrangement. However, only parties to a particular advisory opinion can rely on it as evidence in an enforcement action.

Anyone may request an advisory opinion from OIG, but the decision to request an opinion should be made only after careful consideration of the costs, benefits and risks. For example, though the process for obtaining an advisory opinion is far less costly than an enforcement action, it might still be considered somewhat expensive, and it could result in either an adverse decision or a decision based on a set of facts that might no longer exist once the opinion is issued.

OIG also issues more generalized Special Fraud Alerts, Special Advisory Bulletins and other guidance, all available on its website. If it appears that no helpful advisory opinion is available, one of these other resources may include useful information.

THE PHYSICIAN SELF-REFERRAL BAN (THE STARK LAW)

The physician self-referral statute, 42 U.S.C. § 1395nn, was first introduced by Representative Pete Stark, and therefore became known as the Stark law. As originally enacted in 1992, it prevented physicians from referring Medicare beneficiaries to clinical laboratories in which the physician (or a member of his or her immediate family) has a financial interest. In 1995, however, the law was expanded to cover referrals for a much wider range of health services that were considered to be particularly susceptible to over-utilization. The statute as originally passed is commonly referred to as “Stark I,” while the 1995 expansion is referred to as “Stark II” and a 2007 series of regulations is known as “Stark III.”

Broadly speaking, the Stark law now prohibits a physician from (i) making a “referral” of a patient; (ii) for the furnishing of “designated health services” payable by Medicare (or possibly Medicaid – see discussion below); (iii) to an entity with whom the physician (or the physician’s family) has a “financial interest.”

Key Elements of the Stark law

“Referral” by a Physician
The application of the Stark law is triggered by a referral by a physician. The law’s definition of “referral” contains two components. In the case of referrals for which payment is to be made under Medicare Part B, a “referral” encompasses a request by the physician for designated health services, the “ordering of” any such services or the “certifying or recertifying” of the need for such services. For all other referrals the definition also includes the “establishment of a plan of care” by a physician that includes provision of designated health services.

The key point is that this is a broad definition. Indeed, under the Stark law, it is not necessary for a physician to expressly “refer” a patient to a particular service provider – all that is required is an “order” from the physician and use of the services by the patient.

“Designated Health Services”
Furthermore, for the Stark law to apply, the referral must be for “designated health services” (DHS). Those DHS include:
- Clinical laboratory services
- Physical therapy, occupational therapy, and speech language pathology services
- Radiology/imaging services and radiation therapy services
- Durable medical equipment and supplies
- Parenteral and enteral nutrients, equipment and supplies
- Prosthetics, orthotics, and prosthetic devices and supplies
- Home health services
- Outpatient prescription drugs
- Inpatient and outpatient hospital services

“Financial Relationship”
The Stark law prohibits referrals to DHS providers with whom a physician, or an immediate family member of the physician, has a “financial relationship.” A “financial relationship” is a “direct or indirect ownership interest or compensation arrangement.”

An “ownership interest” includes stock, partnership interests, limited liability company memberships, loans, bonds or other financial interests secured by an entity’s property or revenue. The law also includes ownership interests in any third-party entity that has an ownership interest in the DHS provider. Interests in retirement plans, stock options or unsecured loans are not included.

For purposes of the Stark law, “immediate family members” include the physician’s:
- Husband or wife
- Birth or adoptive parent, child or sibling
- Stepparent, stepchild, stepbrother, or stepsister
- Father-in-law, mother-in-law, son-in-law, daughter-in-law, brother-in-law, or sister-in-law
- Grandparent or grandchild and spouse of grandparent or grandchild

Payable by Medicare (or Medicaid?)
As discussed above, the Stark law originally applied only in the context of DHS payable by Medicare, and still, no express language provides for direct application of the Stark law to self-referrals of Medicaid patients. But that, unfortunately, is not the end of the story.

Under Stark II (the 1995 expansion), a provision was added that prohibits the federal government from reimbursing state Medicaid claims that would violate Stark law prohibitions. Recently, based on that provision, courts have held that physicians and DHS providers may be held liable under the False Claims Act if they cause a state to submit a Medicaid reimbursement claim that would have violated the Stark law in the context of Medicare.

In other words, the Stark law does not expressly apply to referrals of Medicaid patients, but it is likely that referrals of Medicaid patients that would violate the Stark law if made in the context of Medicare patients could also serve as a basis for liability under the False Claims Act. Thus, providers should not assume that the Stark law has no application in the context of Medicaid.

 Exceptions

The Stark law contains a number of exceptions permitting referrals in the case of certain specified “financial relationships.” Generally, most of these exceptions require that the financial relationship be established in advance through a written agreement and that any interest or compensation be made at fair market value and not depend on the amount or value of any referral business.

Unlike the safe harbor provisions in the Anti-Kickback law, the exceptions in the Stark law are the only way an otherwise unlawful referral will be protected. If a referral does not satisfy all of the requirements of an exception, the referral will be deemed to be unlawful.

The Stark law exceptions are grouped into three separate categories: those relevant to both ownership interests and compensation arrangements; those relevant only to ownership interests; and those relevant only to compensation arrangements.

Exceptions to Prohibited Ownership Interests and Compensation Arrangements

- Physician services within the same group practice
- In-office ancillary services (in some case, so long as the physician provides a list of alternative suppliers of such services in the area)
- Services furnished by certain organizations and health plans to enrollees in the organization
- Services provided by academic medical centers
- Implants furnished by an ambulatory surgical center
- EPO and other dialysis related drugs
- Preventative screening tests, immunizations and vaccines
- Eyeglasses and contact lenses following cataract surgery
- Intra-family rural referrals

Ownership Interest Exceptions

- Publicly traded securities and mutual funds
- Hospitals in Puerto Rico
- Ownership in hospitals (under certain, very limited conditions)

**Compensation Arrangement Exceptions**
- Rental of office space and equipment
- Bona fide employment relationships
- Fair market value personal service arrangements
- Payments unrelated to the provision of DHS
- Physician recruitment and retention
- Isolated financial transactions
- Certain group practice arrangements with a hospital
- Payments by physicians for items or services
- Non-monetary compensation up to $392 (2016 figure – adjusted each year for inflation)
- Certain fair market value transactions
- Medical staff incidental benefits (parking, meals, free internet, etc.)
- Risk-sharing arrangements
- Compliance training
- Indirect compensation arrangements
- Referral services that meet anti-kickback safe harbor rules
- Obstetrical malpractice insurance subsidies that meet anti-kickback safe harbor rules
- Professional courtesy
- Retention payments in underserved areas
- Community-wide health information systems
- Charitable donations made by physicians
- Electronic prescribing and health record items and services

**Penalties**
Violations of the Stark law are punishable with civil monetary penalties up to $15,000 for each bill or claim presented for a service that a person knew or should have known violates the Stark prohibition, and up to $100,000 for “circumvention schemes” where physicians or entities enter into arrangements that have the principal purpose of assuring referrals in violation of the Stark prohibition.

Claims will not be paid for any DHS provided in violation of the law, and reimbursement is required for any payments already collected. Violations may also result in exclusion from Medicare, Medicaid and any other federal health care program, and as discussed above, may serve as the basis for liability under the False Claims Act.

**Exclusion from Participation in the Federal Health Care Programs**
Physicians and other providers who are excluded from participation in the federal health care programs may no longer submit claims, either directly or through an entity by which they are employed, to those programs. In fact, health care providers face civil monetary penalties for submitting claims for items or services provided, directly or indirectly, by excluded persons or entities.
Health care providers have an affirmative duty to check the Department of Health & Human Services’ Office of Inspector General’s “List of Excluded Individuals” before hiring or contracting with anyone. This list is available on OIG’s website, www.oig.hhs.gov. If a provider contracts with or hires and excluded individual, it faces penalties of up to $10,000 for each item or service furnished by the excluded individual and submitted in a claim for federal program reimbursement, plus three times the amount claimed, and program exclusion.

**OTHER FEDERAL LAWS**

In addition to the federal False Claims Act, Anti-Kickback law and Stark law, there are a number of additional pertinent federal laws that pose a threat of criminal liability relating to fraud and abuse. Although several are specific to health care claims, there are also general federal criminal statutes that punish false or fraudulent conduct that deprives the federal government of money or property.

**HIPAA**

The Health Insurance Portability and Accountability Act of 1996 (HIPAA) contains certain provisions aimed at preventing health care fraud and abuse. The first provision prohibits health care fraud, which consists of knowingly and willfully executing or attempting to execute a scheme:

- to defraud any public or private health care benefit program; or
- to obtain, by means of false or fraudulent pretenses, representations or promises, any money or property owned or controlled by a health care benefit program in connection with the delivery of or the payment for health care benefits, items or services. 18 U.S.C. § 1347.

Any such health care fraud violations are punishable by fines or imprisonment of up to 10 years, or both, provided that a violation resulting in serious bodily injury or death may result in longer prison sentences.

HIPAA also prohibits theft or embezzlement in connection with health care. 18 U.S.C. § 669. This provision prohibits embezzling, stealing or otherwise, without authority, converting to the benefit of any other person, or intentionally misapplying money, funds, securities, premiums, credits, property, or other assets of a health care benefit program.

HIPAA substantially strengthened health care enforcement authority and established a Fraud and Abuse Control Program to coordinate federal, state, and local health care anti-fraud investigation and enforcement, which is jointly administered by the U.S. Attorney General and the U.S. Department of Health and Human Services Office of the Inspector General. It also authorized the Fraud and Abuse Data Collection Program, a nationwide database reporting final adverse actions taken against health care providers, practitioners, and suppliers by federal and state enforcement authorities. Finally, HIPAA prohibits the concealment of a material fact or the knowing and willful making of a material false statement in connection with the delivery of or payment for benefits, items or services pursuant to a health care program. 18 U.S.C. § 1035.
Additional Criminal Provisions under the Social Security Act

The Social Security Act (SSA), 42 U.S.C. § 1320a-7(a), makes it a felony to knowingly and willfully make or cause to be made any false statement of a material fact in any application for payment, or for use in determining rights to payments, under a federal health care program. It also makes it unlawful to conceal or fail to disclose an improper payment with an intent to fraudulently secure such benefit or payment.

Further, the SSA makes it unlawful for a person to knowingly and willfully make or cause to be made any false statement of a material fact about the operation of any institution, facility or entity to qualify for certification or recertification (if certification is required) or when responding to any required disclosure of information. It additionally makes a misdemeanor the accepting of an assignment of benefits or agreeing to be a participating physician or supplier and knowingly, willfully and repeatedly violating the term of such assignment or agreement. Any claim under the provisions of the SSA described in this section also constitutes a false claim under the False Claims Act.

Mail and Wire Fraud

The mail and wire fraud statute, 18 U.S.C. §§ 1341, 1343, prohibits using the mail or a wire to execute a scheme or artifice to defraud or obtain money or property by means of false or fraudulent representation. Mail or wire fraud is a felony punishable by fines and/or up to twenty years imprisonment.

False Statements

The making false statements statute, 18 U.S.C. § 1001, subjects a person to a fine and/or imprisonment of up to five years when the person, in any matter involving a health care program, knowingly and willfully falsifies, conceals or covers up any trick, scheme or device of material fact, or makes any materially false, fictitious, or fraudulent statement, or makes or uses any materially false document or writing, knowing that the document contains a materially false statement or entry.

Conspiracy Statutes

The conspiracy to defraud the government with respect to claims statute, 18 U.S.C. § 286, prohibits conspiring to defraud any federal agency by obtaining or helping someone else to obtain payment from any false, fraudulent or fictitious claim.

The conspiracy to commit offense or to defraud the United States statute, 18 U.S.C. § 371, applies when two or more persons conspire to commit any offense against, or defraud the federal government or any of its agencies, and at least one of the persons acts in furtherance of the conspiracy.

Theft of Government Property

The theft of government property statute, 18 U.S.C. § 641, prohibits a person from embezzling, stealing, or knowingly converting to his/her use or the use of another, or selling, conveying, or disposing of anything of value to the United States or any of its agencies. Anyone who receives, conceals or retains any such thing of value, intending to use it to his or her gain, and knowing it to have been embezzled or stolen, is subject to fines or imprisonment or both.
RICO
The Racketeer Influenced and Corrupt Organizations Act (RICO), 18 U.S.C. §§ 1961 et seq., the application of which is not limited to organized crime, prohibits a person from receiving income, directly or indirectly, from a pattern of racketeering activity. A pattern is two or more occurrences of a “predicate act” over a 10-year period. RICO contains a private right of action for persons injured in their business or property.

Money Laundering
The criminal statute governing money laundering, 18 U.S.C. §§ 1956, 1957, prohibits knowingly engaging or attempting to engage in a “monetary transaction in criminally derived property” valued over $10,000 and derived from “specified unlawful activity.”

Obstruction
The obstruction of criminal investigation statute, 18 U.S.C. § 1510, prohibits willfully bribing someone to obstruct, delay or prevent communication of information relating to violation of any criminal statute to a criminal investigator.

The obstruction of criminal investigations of health care offenses statute, 18 U.S.C. § 1518, prohibits preventing, obstructing, misleading, delaying or attempting to do any of those things with respect to communicating information or records relating to a violation of a federal health care offense to a criminal investigator.

Aiding and Abetting
People who aid or abet in the commission of an offense are punishable as if they were principals. 18 U.S.C. § 2. In addition, people who, knowing that an offense against the United States has been committed, assists the offender in order to hinder his or her apprehension or punishment is an accessory after the fact. 18 U.S.C. § 3.

Vermont Laws
In addition to the federal laws on health care fraud and abuse, health care practitioners should also become familiar with Vermont’s fraud and abuse laws. Of particular importance are (i) the State False Claims Act (the State FCA), (ii) the criminal false representation law, (iii) the Medicaid fraud statute, and (iv) the Healthcare Whistleblower’s Protection Act.

State False Claims Act
In 2015, Vermont became the thirtieth state to adopt a civil false claims act modeled on the federal FCA (see discussion in Part A above on the federal FCA). The State FCA, 32 V.S.A. §§ 630 et seq., authorizes civil enforcement for any false or fraudulent claims made for State funds. Vermont’s definition for the term “claim” under the State FCA tracks the federal definition: a claim is “any request or demand, whether under a contract or otherwise, for money or property” made to the State of Vermont or certain government contractors. The act is broad and prohibits any person from, among other things:
- knowingly submitting, or causing to be submitted, a false claim;
knowingly making or using any false record or statement material to a false claim or to an obligation to pay the State;
knowingly submitting a claim that violates the State’s criminal false claims act (see below) or the federal false claims act, anti-kickback statute or Stark law (see above for a discussion of these federal fraud and abuse laws);
knowingly and improperly avoiding or decreasing an obligation to pay the State;
entering into a written agreement with the State knowing the agreement contains false information;
if the person discovers after submission that a false claim has been made resulting in overpayment, failing to timely disclose the false claim or receipt of overpayment to the State; and
conspiring to submit a false claim.

General Categories of False Claims
The above-listed prohibitions of the State FCA can be summarized as prohibiting five general types of behaviors:

- **Mischarge.** The act prohibits a person from billing the State for goods or services that are not actually provided. This category of prohibited acts includes overcharging for goods and services, such as upcoding for health care services.
- **Fraud-in-the-inducement.** The act prohibits a person from making false statements or taking illegal actions during the formation of an agreement with the State. Examples include making false representations and warranties in a contract with the State or making an increased number of claims due to a kickback arrangement.
- **False certification.** Under the act, a person may not falsely certify that he or she qualifies for a State program, that he or she is eligible for a State benefit, or that his or her claims are complaint with the State program’s requirements.
- **Substandard service or product.** The act prohibits a person from providing a substandard service or product under a State contract. For example, a false claims action could be brought against a physician for failing to meet applicable quality of care standards under a State contract.
- **Reverse false claim.** Finally, the act prohibits so-called “reverse false claims.” A reverse false claim occurs when a person has an obligation to pay the government and tries to avoid that obligation. In other words, it is the government that is owed the payment, as opposed to the traditional false claim that involves the government paying a person. One such reverse false claim arises when a person keeps what the State has given him or her, as the beneficiary of an inadvertent false claim or overpayment, if that person fails to disclose the claim or overpayment within the later of (i) 120 days after discovery of the false claim or overpayment, or (ii) the date of a corresponding cost report. Another type of reverse false claim arises if a person knowingly uses a false record or statement to avoid paying or reducing the amount owed to the State.

Investigation and Enforcement.
Similar to the federal FCA, the State FCA may be enforced either by the State attorney general or by a private citizen who is aware of the false claim(s), called a “relator.” A suit by a relator under the State FCA is filed in the name of the State and under seal, pending review by the State attorney general’s office. When the suit is under seal, it means that the filing of the case is not
public record and is confidential; the named defendant is not notified of the suit until the court issues an unsealing order. The attorney general has an initial 60-day period during which to determine whether to intervene in the suit or to decline the case, though the attorney general may seek an extension of the seal period.

If the attorney general elects to intervene, then it generally has full control over the suit and may move to dismiss or settle the case. If the attorney general declines to intervene, then the relator has the right to conduct the action. Even if the attorney general declines to intervene, the office retains the right to receive all pleadings and transcripts, and may choose to intervene at a later date by showing good cause. And even in declined cases, the State typically will be involved in settlement negotiations, either by intervening or by working with the relator’s attorney.

Like the federal FCA, the State FCA provides a financial incentive for relators. If the attorney general intervenes in the suit, the relator is entitled to 15-25 percent of any recovery, based upon the relator’s contribution to the suit. If the attorney general declines to intervene, then the relator is entitled to receive 25-30 percent of any recovery, again based on the relator’s contribution. Successful relators are also entitled to receive reasonable attorney’s fees and expenses from the defendant.

In investigating potential violations of the State FCA, the attorney general has the authority to issue a civil investigative demand (CID). A CID may be issued when there is reason to believe that a person may be in possession of any information relevant to a false claims investigation, and it includes the power to seek documents, interrogatories, or oral testimony. Any CID must state the nature of the conduct that constitutes the alleged violation, as well as the applicable provision of law that has been allegedly violated. Materials obtained by the attorney general from a CID may be used for any lawful purpose in investigating or prosecuting a claim under the act, and may be shared with a relator as necessary as part of the investigation.

**Damages and Penalties.**
State FCA violations carry stiff penalties and damages. The act provides for the following mandatory penalties upon a finding of liability:

- A civil penalty of not less than $5,500 and not more than $11,000 per false claim;
- Three times the amount of damages that the State sustained in connection with the false claim(s); and
- The costs of investigation and prosecution of the false claims(s).

A person can limit his or her exposure to damages and penalties under the State FCA by self-reporting. A person who self-reports a false claim prior to the commencement of any action regarding the violation will not be subject to civil penalties under the law and will be subject to reduced damages of two times the amount of damages sustained by the State in connection with the false claims(s).

*The Criminal False Representation Law.*
In addition to the State FCA, which is a civil law, Vermont also has a criminal false representation law, 33 V.S.A. § 141, that prohibits a person from making any false or fraudulent material statements, or falsifying, concealing, or covering up with any trick, scheme or device,
any material facts, or falsifying any documents or writings knowing they contain a material fact, regarding any matter within the jurisdiction of a state or local government body.

The penalty for a violation depends on the amount of the loss sustained by the government entity and the benefit gained by the person. If there is no loss to the government or gain to the person, or if the loss or gain is less than $500, the violator can be imprisoned for up to two years and fined up to $5,000, or both. If the government’s loss or the person’s gain exceeds $500, the person can be imprisoned for up to five years, fined up to $10,000, or both.

A provider may be subject to both the State FCA and criminal false representation law, as well as the federal FCA. However, a provider who commits an act punishable under the Medicaid fraud statute discussed below may not be prosecuted under the state criminal false representation law.

**The Medicaid Fraud Statute**

Medicaid fraud is committed when a provider is untruthful regarding services provided to Medicaid beneficiaries to obtain improper payment. A provider may commit Medicaid fraud, for instance, by submitting false claims and records for services or supplies that were not provided, billing twice or more for the same services or supplies, or billing for services or supplies that were not medically necessary. The Medicaid fraud statute, 33 V.S.A. § 14, specifically defines fraudulent conduct as:

- knowingly filing, attempting to file, or aiding and abetting in filing a claim for services to a Medicaid beneficiary that were not rendered; or
- knowingly filing a false claim, or a claim for unauthorized items or services under the Medicaid program; or
- knowingly billing the Medicaid recipient or family for amounts in excess of the amount allowable by law; or
- failing to credit the State for amounts received from Social Security, insurance or other sources; or
- in any way knowingly receiving, attempting to receive, or helping to receive unauthorized payments from Medicaid.

**Investigation and Enforcement.**

Medicaid is a joint federal-state program, and states administer their own Medicaid programs within the context of minimum federal requirements. Congress enacted the Medicare and Medicaid Fraud and Anti-abuse Amendments in 1977 to authorize and substantially fund Medicaid Fraud Control Units in the states.

The Medicaid Fraud and Residential Abuse Unit of the Vermont attorney general’s office is responsible for investigating and prosecuting providers who commit fraud against the Medicaid program. It also is responsible for investigating and prosecuting instances of residential patient abuse and neglect.

**Penalties.**

Medicaid fraud is a felony and conviction can lead to substantial penalties including but not limited to imprisonment for up to 10 years, a fine of up to $1,000 or an amount equal to twice the amount of the assistance or benefits wrongfully obtained, or both a fine and imprisonment.
Additionally, individuals convicted of Medicaid fraud will be excluded from participating in Medicaid for four years unless the state secretary of human services waives the suspension after finding that the recipients the provider serves would suffer a substantial hardship by being denied medical services that cannot reasonably be obtained through another provider. 33 V.S.A. § 143a.

There is also a provision that allows the attorney general to bring a civil action against a provider who knowingly violates the fraud statute. 13 V.S.A. § 3016. The penalties for civil violations include restitution of any amounts wrongfully obtained plus interest, and a penalty of up to three times the amount wrongfully obtained, or $500 per false claim, or $500 for each false document submitted in support of a false claim, whichever is greatest.

**Healthcare Whistleblower’s Protection Act.**

The act, 21 V.S.A. § 507 - 509, makes it illegal for a hospital or nursing home to fire, threaten, or take any adverse employment action (such as demotion, suspension, failure to make a promotion, discrimination) against an employee because the employee:

- Discloses or threatens to disclose what he or she reasonably believes is a violation of law or improper quality of patient care by the employer or by an agent of the employer;
- Testifies or provides information to a public body that is investigating whether the employer violated any law or engaged in behavior constituting improper patient care; or
- Objects to or refuses to participate in any activity, policy, or practice of the employer’s that the employee reasonably believes is a violation of law or constitutes improper quality of patient care.

To receive protection under the act (other than protection for testimony), the employee must first report the alleged violation of law or improper care to the employer, supervisor or other person designated by the employer to address such reports, unless the employee reasonably believes that doing so would be futile. After receiving the employee’s report, the employer must then have a reasonable opportunity to address the violation. Any violation must be addressed under the employer’s compliance plan, if it has one.

The law gives aggrieved employees the right to bring an action in superior court seeking such relief as reinstatement, back pay, lost wages and benefits, punitive and compensatory damages, and attorney’s fees.

Hospitals are required to have internal processes that reflect the Magnet Recognition Program quality care and professional standards developed by the American Nurses Credentialing Center. Hospitals and nursing homes are also required to post a notice about the Healthcare Whistleblower Protection Act. The notice is available on the Vermont Department of Labor’s website, labor.vermont.gov. An employer’s willful failure to post the notice can lead to a fine of up to $100 per day.

**Preventative Action Compliance Programs**

Every office should voluntarily adopt and scrupulously adhere to a compliance program. In October 2000, OIG issued a document entitled “Compliance Program Guidance for Individual and Small Group Physician Practices.” OIG has issued a number of such compliance program
guidelines for different segments of the health care industry, including clinical laboratories, hospitals, home health agencies, third-party medical billing companies, durable medical equipment suppliers, hospices, pharmaceutical manufacturers, ambulance suppliers, Medicare+Choice organizations, and nursing facilities. The most recent guideline issued is a “Supplemental Compliance Program Guidance for Nursing Facilities,” which was published in September 2008. OIG’s compliance program guidance documents are posted on its website at https://oig.hhs.gov/compliance/compliance-guidance/index.asp.

OIG’s motive in issuing compliance program guidance is to engage the industry in avoiding the submission of erroneous claims and ferreting out fraudulent conduct through internal controls that monitor adherence to the laws and program requirements. Having a compliance program in place is one of the factors that enforcement authorities will take into account when assessing some violations.

It is important, however, that the compliance policy not be simply a volume on a shelf, but a discernible good faith attitude and set of procedures that are integrated into the fabric of the practice.

**Basic Recommendations for Individual and Small Group Physician Practices.**

The OIG Compliance Program for Individual and Small Group Physician Practice guidance sets forth seven basic elements that should be addressed in every compliance program. OIG acknowledges that small and solo practices have limited resources to devote to a compliance program. Therefore, the fundamental principle in adopting a compliance program is showing a good faith commitment to compliance, best exhibited by taking reasonable steps to address each of the following seven elements.

1. Establish written practice standards and procedures – after doing an initial audit to understand risk areas.
2. Audit and monitor – periodically review the office’s standards and procedures and audit the claims submission process.
3. Designate a compliance officer or compliance contacts.
4. Conduct appropriate training and education in coding, billing and compliance.
5. Respond to detected offenses and develop corrective action initiatives.
6. Develop open lines of communication.
7. Enforce disciplinary standards through well-publicized guidelines.

**Other Useful Materials for Compliance with Fraud and Abuse Issues.**

OIG publishes a Work Plan for each fiscal year that lists the areas it intends to focus on during the year to protect the integrity of all of the Department of Health and Human Services programs, including Medicaid and Medicare. The Work Plan can serve as an annual guide for practitioners for purposes of reviewing compliance issues and risks. Each year, starting in early October, practitioners can review the compliance areas of interest to OIG for the coming year with respect to hospitals, physician practices, home health agencies, and other health care providers. The current and past Work Plans can be found at https://oig.hhs.gov/reports-and-publications/workplan/index.asp#current.
In addition, OIG issues special fraud alerts and special advisory bulletins to inform the health care industry of particular practices that it deems suspect. As mentioned earlier, OIG responds to requests for formal advisory opinions on the application of the anti-kickback statute and other fraud and abuse statutes to particular business arrangements. All of these materials are found on OIG’s website, http://oig.hhs.gov.

Finally, OIG provides compliance education resources and materials aimed at a fairly basic level at its “Compliance 101” website at https://oig.hhs.gov/compliance/101/index.asp. In addition to links to the compliance guidance documents described above, the website features a variety of educational materials aimed at providers including videos, podcasts, bulletins, transcripts and more.

CONCLUSION

As you can see, there are many laws regulating fraud and abuse in the health care field, many of which are quite complex. Moreover, violations of these laws can lead to substantial penalties. Accordingly, if someone alleges that you or your practice has committed fraud or abuse, or if you discover potential fraud or abuse, you should contact competent fraud and abuse counsel right away.

ABOUT THE AUTHORS

Eileen Elliott is an attorney focusing on health care and human services legal issues with the Burlington law firm of Dunkiel Saunders Elliott Raubvogel & Hand, PLLC. She was the deputy secretary of Vermont’s Agency of Human Services from 2003-2005 and the commissioner of Vermont’s Social Welfare Department from 1999-2003. From 1993 until 1999, she worked for the Office of the Attorney General, serving as chief of its Human Services Division, and before that as counsel to the Agriculture Department. She spent the first decade of her professional career in private and corporate practice. Eileen graduated with distinction from the University of Colorado with a B.A. in Environmental Conservation and she has a J.D. from the University of Denver.

Drew Kervick is a health care and business law attorney with Dunkiel Saunders Elliott Raubvogel & Hand, PLLC. Before joining the firm in 2014, Drew worked in private practice both in Burlington, Vermont and in Boston Massachusetts. He also clerked for Justice John Dooley of the Vermont Supreme Court. Drew graduated summa cum laude from Boston College Law School, and holds a master’s degree from University of North Carolina at Chapel Hill and a bachelor’s degree from Stanford University, both in Political Science.

Jon Rose is a regulatory and civil litigation attorney at Dunkiel Saunders Elliott Raubvogel & Hand, PLLC. Prior to joining Dunkiel Saunders, Jon served as an Assistant Attorney General in the Civil Division of the Vermont Attorney General’s Office, and before that, practiced at another prominent Burlington law firm advising a wide range of institutional and corporate clients on labor and employment, health care, and regulatory issues. Jon graduated first in his class from Vermont Law School, and holds a B.S. in Mechanical Engineering from Penn State University.
Topics Covered in this Chapter:
- The Vermont Marijuana Registry
- Legal and Regulatory Considerations for Clinicians
- Resources
- About the Authors

By Lindsey Wells, B.S & Meredith Bullock
Vermont Department of Public Safety

In the United States, marijuana for is legal in over half the states and Washington, D.C. for medicinal purposes. In 2004, the Vermont Legislature passed Act 135, “An Act Relating to Marijuana Use by Persons with Severe Illness.” The Department of Public Safety was designated to administer the Vermont Marijuana Registry (VMR). There have been various amendments passed since 2004, the most notable being the 2011 amendment authorizing the VMR to register four dispensaries throughout the state.

In Vermont, marijuana for symptom relief may be used to alleviate the symptoms or effects of a registered patient’s debilitating medical condition as permitted by Vermont statute and the Rules Regulating Cannabis for Symptom Relief. More information about the clinical applications of cannabis for therapeutic use can be found from the University of Vermont Medical Cannabis Program.

THE VERMONT MARIJUANA REGISTRY

What is the Vermont Marijuana Registry?
The Vermont Marijuana Registry (VMR) is a program located within the Department of Public Safety whose purpose is to implement the provisions of 18 V.S.A. Chapter 86, Therapeutic Use of Cannabis, as they pertain to registered patients, caregivers, and the creation and operation of four dispensaries. The VMR’s primary purpose is to assist individuals applying for a registry identification card and oversee the operations of the four registered dispensaries in Vermont that provide marijuana for symptom relief to registered patients. The VMR reviews and processes applications, issues registry identification cards to residents of Vermont with verified debilitating medical conditions and evaluates the compliance of registered dispensaries. Additionally, the VMR strives to ensure registered patients are able to access cannabis for symptom relief in a timely manner, protect confidentiality, and prevent the diversion and theft of cannabis. The VMR website, including links to statutes, rules and application forms can be found here.

Who is eligible to register with the VMR?
The act creating the VMR was designed to establish a registration process for those with a severe illness who wish to use marijuana for symptom relief. To become a registered patient, a person must be diagnosed with a debilitating medical condition by a health care professional in the course of a bona fide health care professional-patient relationship. A registered patient must also be a resident of Vermont and issued a registry identification card by the VMR. "Resident of
Vermont” means a person whose domicile is Vermont. A “debilitating medical condition” means reasonable medical efforts have been made over a reasonable amount of time to relieve the symptoms associated with:

(A) cancer, multiple sclerosis, positive status for human immunodeficiency virus, acquired immune deficiency syndrome, glaucoma, or the treatment of these conditions, if the disease or the treatment results in severe, persistent, and intractable symptoms; or

(B) a disease, medical condition, or its treatment that is chronic, debilitating, and produces one or more of the following intractable symptoms: cachexia or wasting syndrome; chronic pain; severe nausea; or seizures.

How does an individual apply for a registry identification card with the Vermont Marijuana Registry?
To apply a patient must submit:
- A completed Department-approved patient application;
- A Health Care Professional Verification Form completed by a health care professional;
- The required $50 non-refundable fee payable to the Department of Public Safety; and,
- An electronic color photo of themselves from the shoulders up. Photos may be submitted on a CD or emailed to the VMR at DPS.MJRegistry@vermont.gov.

The VMR must approve or deny an application in writing within 30 days of receipt of a completed application. If the application is approved, the VMR will issue the applicant a registry identification card. A registry identification card expires one year after its issue date, and may be renewed by completing and submitting the required forms and fee listed above.

A registered patient may designate a caregiver to assist with marijuana for symptom relief. A caregiver is a person who has agreed to undertake responsibility for managing the well-being of a registered patient with respect to the use of cannabis for symptom relief. A registered caregiver must be a person who is a resident of Vermont, at least 21 years of age, and has met the criminal history record requirements contained in the Rules Regulating Cannabis for Symptom Relief. A registered caregiver cannot be a currently registered patient and may only assist one registered patient. To apply as a caregiver and individual must submit:
- A completed Department-approved caregiver application;
- The required $50 non-refundable fee payable to the Department of Public Safety; and,
- An electronic color photo of themselves from the shoulders up. Photos may be submitted on a CD or emailed to the VMR at DPS.MJRegistry@vermont.gov.

What is the definition of a Health Care Professional?
A health care professional is an individual licensed as a:
- Doctor of Medicine (M.D.);
- Osteopathic Physician (D.O.);
- Naturopathic Physician (N.P);
- Physician Assistant (P.A.); or,
- Advanced Practice Registered Nurse (APRN).
This definition includes individuals licensed in Vermont and under substantially equivalent provisions in New York, New Hampshire and Massachusetts.

**What are the responsibilities of the patient’s Health Care Professional?**

A health care professional verifies that the patient has a debilitating medical condition and whether or not the health care professional has a “bona fide health care professional-patient relationship” with the applicant. The phrase “bona fide health care professional-patient relationship” means a treating or consulting relationship of not less than three months’ duration, in the course of which the health care professional has completed a full assessment of the registered patient’s medical history and current medical condition, including a personal physical examination. 18 VSA § 4472 (1)(A).

The three-month requirement does not apply if a patient has been diagnosed within the past six months or diagnosed with:
- A terminal illness;
- Cancer;
- Acquired immune deficiency syndrome; or
- Is currently under hospice care.

Two additional exceptions to the three-month requirement were added in 2016:

1. When a patient been diagnosed with a debilitating medical condition by a health care professional in another jurisdiction in which the patient had been formerly a resident and the patient is now a resident of Vermont and has his or her diagnosis confirmed by a health care professional in Vermont or a neighboring state. The new health care professional must still complete a full assessment of the patient’s medical history and current medical condition, including a personal physical examination. 18 VSA § 4472 (1)(B)(ii).

2. When a patient who is already registered with the VMR changes health care professionals three months or less prior to the annual renewal of the patient’s registration, provided the patient’s new health care professional has completed a full assessment of the patient’s medical history and current medical condition, including a personal physical examination. 18 VSA § 4472 (1)(B)(iii).

The health care professional must also certify that reasonable medical efforts have been made over a reasonable amount of time without success to relieve the patient’s symptoms. See VMR Rules Regulating Cannabis for Symptom Relief § 3.3.

The Health Care Professional Verification Form can be found here.

**Can a Health Care Professional inactivate a patient’s registry identification card?**

Yes, a registered patient’s verifying health care professional may void a registered patient’s registry identification card by notifying the VMR in writing. The VMR will issue a written notice informing the patient his or her registry identification card has been inactivated. The patient’s designated dispensaries will also be informed when a registered patient has been inactivated. See VMR Rules Regulating Cannabis for Symptom Relief § 11.9.5.
Where can I obtain information about the dispensaries and the types of products they offer?
The VMR website has information available from the dispensaries in a document titled
Dispensary Selection Information. Additional information may be obtained from the individual
dispensary’s websites listed under Resources.

What protections are there for Health Care Professionals?
A health care professional who has participated in a patient’s application process under the
statute, rules, policies or procedures of the VMR are not be subject to arrest, prosecution, or
disciplinary action under 26 V.S.A. Chapter 23, penalized in any manner, or denied any right or
privilege under state law, except for giving false information, pursuant to 18 V.S.A. § 4474c(f).
18 VSA § 4474b (b). Health Care Professionals are not asked to prescribe or recommend the use
of marijuana. They are verifying the nature of the disease and its symptoms. All information
received by the VMR is confidential, including identifying information pertaining to health care
professionals.

Have questions about the Vermont Marijuana Registry?
Contact the VMR at:
45 State Drive
Waterbury, VT  05671-1300
Tel: 802.241.5115
Fax: 802.241.5230
Email: DPS.MJRegistry@vermont.gov

OTHER LEGAL AND REGULATORY CONSIDERATIONS FOR CLINICIANS

This section has been prepared by the Vermont Medical Society to provide additional
information to clinicians considering whether to participate in a patient’s application under
Vermont law. For further information, contact the Vermont Medical Society at 802-223-7898.

Despite state laws legalizing marijuana for therapeutic and recreational use, the federal
government continues to classify marijuana as a Schedule I drug with no currently accepted
medical use and a high potential for abuse. Federal law prohibits knowingly or intentionally
distributing, dispensing, or possessing marijuana. 21 U.S.C. §§841–44. Additionally, a person
who aids and abets another in violating federal law or engages in a conspiracy to purchase,
cultivate, or possess marijuana may be punished to the same extent as the individual who
violate federal law are also possible, such as revocation of a prescriber’s DEA registration or
exclusion from participation in the Medicare and Medicaid programs.

Based on the increasing number of states legalizing marijuana for therapeutic and recreational
use, the U.S. Department of Justice has issued several memoranda regarding its marijuana
enforcement policy. An August 2013 updated policy reiterates marijuana’s classification as an
illegal substance under federal law, but states that it is not an efficient use of resources to focus
federal enforcement efforts on seriously ill individuals and that federal enforcement activities
will focus on addressing threats such as distribution to minors, revenue going to criminal
enterprises and state-authorized marijuana activity from being used as a cover for illegal activity.
Participating in a patient’s application process under a state medical marijuana law is generally considered low risk for federal enforcement or other disciplinary or legal action. However, the continuing inconsistency between federal and state law, and lack of clarity in case law around the country regarding the extent to which clinicians are protected from federal prosecution when discussing marijuana with patients, raises questions for medical professionals. The Vermont Medical Society recommends that physicians and other clinicians consider the following:

- If you assist a patient with the application process, you should comply with all state statutory requirements for establishing a bona fide professional-patient relationship under the Vermont program and meet the same standard of care as in other types of patient encounters. Disciplinary actions from other states that have been reported in the press relate to issues such as physicians completing a high number of forms without establishing the required physician-patient relationship, not performing required medical exams and/or maintaining inadequate patient records. See the model guidelines created by the Federation of State Medical Boards discussing best practices regarding patient evaluations, medical record keeping, informed decision making and written treatment agreements.

- Limit the information provided to the Department of Public Safety to that required to complete and confirm the accuracy of the information contained on the Health Care Professional Verification Form. Do not prescribe, and avoid “recommending,” marijuana to patients. While health care professionals may discuss relevant information regarding possible health risks and therapeutic benefits of cannabis with patients, “recommending” marijuana may be seen as akin to “prescribing” and abetting the patient in obtaining an illegal substance in violation of federal law. (See e.g. Conant v. Walters (9th Cir. 2002) 309 F.3d 629, affirming Conant v. McCaffrey (N.D.Cal. Sept. 7, 2000) 2000 WL 1281174). Also, avoid offering individualized patient advice concerning appropriate cannabis strains, dosage, timing, amount and route of administration as this may also be interpreted as prescribing.

- Do not dispense or otherwise provide marijuana to patients.

- Discuss your participation with your liability carrier to ensure that you would have coverage for any harms resulting from medications, including cannabis, that are not approved by the FDA.

- Discuss your participation with any employer or facility for which you work. Health care facilities and employers may have to comply with additional federal regulations that impact the ability of a physician to participate in the application process.

- The statute does not require a health care professional to participate in the application process.

RESOURCES

- Department of Public Safety, Vermont Marijuana Registry: http://vcic.vermont.gov/marijuana-registry

- Vermont Marijuana Registry, Document Library (Forms and other useful information) http://vcic.vermont.gov/marijuana-registry/library
- Vermont Statutes Online, Title 18, Chapter 86
  http://legislature.vermont.gov/statutes/fullchapter/18/086

- Federation of State Medical Boards, Model Guidelines for the Recommendation of
  Marijuana in Patient Care (April 2016)

- Vermont Department of Health, Board of Medical Practice
  http://healthvermont.gov/systems/medical-practice-board

- Mayo Clinic, Medical Marijuana
  http://www.mayoclinic.org/healthy-lifestyle/consumer-health/in-depth/medical-marijuana/art-20137855

- Champlain Valley Dispensary and Southern Vermont Wellness: http://www.cvdvt.org/

- Vermont Patients Alliance: http://www.vtpatientsalliance.org/

- Grassroots Vermont: http://www.grassrootsvermont.com/

ABOUT THE AUTHORS

Lindsey Wells, B.S., received her Bachelor of Science Degree in Business Administration in 2005 from Norwich University. She has an extensive work history with state and federal regulations. She has been employed by the Department of Public Safety as the Marijuana Program Administrator for the Vermont Marijuana Registry since 2012. Her duties include overseeing and performing assessments of Vermont’s four registered dispensaries, overseeing the Registry’s applicant process, participating in the legislative process, and amending the rules governing the program to facilitate the implementation of various statutory changes.

Meredith Bullock has been employed by the Department of Public Safety as a Program Technician for the Vermont Marijuana Registry (VMR) since August 2013. Her job duties include assisting applicants and the general public with inquiries, interacting with Health Care Professionals, and processing applications for patients, caregivers, and dispensary personnel. Since she began working for the VMR she has strived to build the VMR’s rapport with the general public, applicants, dispensary personnel, and health care professionals.
NON-DISCRIMINATION IN HEALTH CARE

Topics Covered in this Chapter:
General Non-Discrimination Requirements/Section 1557
Disability Discrimination & ADA Accommodations
Limited English Proficiency & Interpreter Requirements
About the Author

By Jessa Barnard, Esq.
Vermont Medical Society

GENERAL NON-DISCRIMINATION REQUIREMENTS/SECTION 1557

In May 2016, the US Department of Health and Human Services (HHS) issued a Final Rule implementing a prohibition of discrimination by health care services found in Section 1557 of the Affordable Care Act (ACA). Section 1557 builds on long-standing Federal civil rights laws that have already applied to most health care practices, such as Title VI of the Civil Rights Act of 1964 (requiring, among other things, access for those with limited English proficiency) and the Americans with Disabilities Act. While many of the provisions of the Final Rule are not new to physicians, there are several new procedural requirements and clarifications of existing requirements.

Generally, Section 1557 and the Final Rule prohibit “discrimination on the basis of race, color, national origin, sex, age or disability [by] any health program or activity, any part of which is receiving Federal financial assistance.”

Does the new rule apply to me?
The Final Rule applies to “every health program or activity, any part of which receives Federal financial assistance.” HHS expects that this applies to almost all physician practices in the nation as Federal financial assistance includes, but is not limited to: state Medicaid payments, “meaningful use” payments, National Health Service Corp funding, NIH research funding and CMS gain-sharning payments. Physician practices that exclusively receive Medicare Part B payments are not included, but practices should conduct a thorough review of their funding sources and consult an attorney before deciding they are exempt.

When does the law go into effect?
Section 1557 has been in effect since the passage of the ACA and the Final Rules went into effect July 18, 2016 – with a limited exception for practices to have until October 16, 2016 for posting required notices and taglines, discussed further below.

What does the law require?
The law clarifies and extends discrimination protections on the basis of national origin, sex, and disability and adds several new procedural requirements.
What are the new procedural requirements?

- **Designate responsible employee:** Practices employing 15 or more individuals need to designate at least one employee to coordinate its efforts to comply with and carry out the responsibilities under Section 1557, including investigating any grievance alleging noncompliance. (Effective July 18, 2016)

- **Adopt grievance procedures:** Practices employing 15 or more individuals need to adopt a grievance procedure that provides for the prompt and equitable resolution of any grievance alleging discrimination in violation of the law. HHS has provided a model procedure that meets the requirements [here](#). (Effective July 18, 2016).

- **Post notice:** Beginning 90 days after the effective date of the rule (October 16, 2016) practices will need to post a notice of non-discrimination in English AND shorter “taglines” in the top 15 non-English languages spoken in Vermont about the availability of free language assistance services. The long form English notice and the 15 taglines need to be posted in a visible font size in:
  - A conspicuous physical location where the practice interacts with the public (e.g. the waiting room); AND
  - In a conspicuous location on the practice website (if the practices has one), accessible from the homepage of the website - meaning a link to the notice must appear on the homepage. Links to the 15 non-English taglines must also appear on the homepage and must be provided in the non-English language (e.g. “Español”, not “Spanish”). You may have to work with your web designer to ensure these links appear appropriately.

  HHS has provided a [sample notice in English](#) and [sample taglines](#). Practices in Vermont will need the following 15 taglines: French, Spanish, Vietnamese, Chinese/Mandarin, Nepali, Serbocroatian (Serbian language), German, Cushite (Oromo language - African), Arabic, Russian, Tagalog, Italian, Thai, Japanese, and Portuguese.

  The Vermont Medical Society has created an editable document that contains the Notice and 15 Taglines [here](#).

- **Include notice in communications:** Beginning October 16, 2016, practices must also include the notice and taglines in “significant publications” or “significant communications” targeted at patients or the public. HHS will interpret this broadly and does not specify which communications are included but provides examples such as patient handbooks, consent and complaint forms, initial patient paperwork, outreach publications, marketing materials and any notice “requiring a response from an individual” – which could include patient bills or notices to contact the practice to schedule an appointment. The notice/taglines can be part of the publication or a separate insert. While the compliance date is October, you can continue to exhaust your current stock of materials before you reprint with the necessary notices.
  - The full notice and 15 taglines (links above) must be included in any larger format publications.
A shortened “statement of non-discrimination” in English and taglines in the top 2 non-English languages in Vermont (Spanish, French) must be included in significant publications or communications that are small-sized, such as postcards or tri-fold brochures.

**What changes should I know about to non-discrimination/Language Access Laws**

The Final Rule also clarifies and strengthens various non-discrimination protections that apply in the health care setting. Among other things, the law addresses:

- **Limited English Proficiency/Requirements for Interpretation Services.** Most health care providers have already been required to provide language access services to patients or prospective patients with limited English proficiency. Under this Final Rule, all covered practices are required to “take reasonable steps to provide meaningful access to each individual with limited English proficiency eligible to be served.” See more details in the section on Limited English Proficiency, below.

- **Disability Discrimination:** The final rule reiterates the requirement that physician practices not discriminate on the basis of disability. See more details in the Disability Discrimination section, below.

- **Sex Discrimination**
  - The final rule establishes that individuals cannot be denied health care or services based on their sex. Sex-specific health programs are permissible only if the entity can demonstrate an exceedingly persuasive justification (e.g. most sex-specific clinical trials would still be considered appropriate.)
  - As adopted, the final rule also prohibited discrimination on the basis of gender identity. This required individual to be treated consistent with their gender identity and prohibited providers from denying or limiting services ordinarily available to individual of one gender based on the fact that a person seeking such services identifies as belonging to another gender. (E.g. a provider may not deny treatment for ovarian cancer to an individual who identifies as a transgender male where the treatment is medically indicated). On December 31, 2016, the U.S. District Court for the Northern District of Texas issued an opinion in *Franciscan Alliance, Inc. et al v. Burwell*, enjoining the Section 1557 regulation’s prohibitions against discrimination on the basis of gender identity and termination of pregnancy on a nationwide basis. Accordingly, HHS’ Office for Civil Rights (HHS OCR) is not enforcing these same provisions at the time of publication of this Guide.

**Where can I find more information?**

HHS has a website dedicated to Section 1557, available here, that includes a summary of the rule, fact sheets and an FAQ page.

**DISABILITY DISCRIMINATION & ADA ACCOMMODATIONS**

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.

What laws address providing health care services to those with disabilities?
The [Americans with Disabilities Act of 1990 (ADA)](https://www.gpo.gov/fdsys/pkg/PLAW-101STAT254/pdf/PLAW-101STAT254.pdf) is a federal civil rights law that prohibits discrimination against individuals with disabilities in every-day activities, including medical services. Private hospitals or medical offices are covered by [Title III of the ADA](https://www.gpo.gov/fdsys/pkg/PLAW-101STAT254/pdf/PLAW-101STAT254.pdf) as places of public accommodation. [Section 504 of the Rehabilitation Act of 1973](https://www.gpo.gov/fdsys/pkg/PLAW-101STAT254/pdf/PLAW-101STAT254.pdf) (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 that receive federal financial assistance, which can include Medicare and Medicaid reimbursements. [Section 1557](https://www.gpo.gov/fdsys/pkg/PLAW-113STAT589/pdf/PLAW-113STAT589.pdf) of the Affordable Care Act, and accompanying regulations, reiterate the requirement that physician practices not discriminate on the basis of disability. These statutes require medical care providers to make their services available in an accessible manner.

Vermont public accommodation law, [9 VSA §4502](https://statutes.vermont.gov/dig/95/4502.html), contains substantially the same requirements as the ADA and Section 504 and requires that places of public accommodation, including medical practices, provide an individual with a disability the opportunity to participate in its services, facilities, privileges, advantages, benefits and accommodations. Failure to comply may result in an action for civil penalties either through the Vermont Human Rights Commission or a private court action.

What accommodations must be provided?
Federal and State public accommodations law require that medical care providers provide individuals with disabilities:

- Full and equal access to their health care services and facilities; and
- 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 or create an undue burden.

Regarding physical modifications/accommodations for those with mobility disabilities, the ADA sets [requirements](https://www.gpo.gov/fdsys/pkg/PLAW-101STAT254/pdf/PLAW-101STAT254.pdf) for new construction of and alterations to buildings and facilities, including health care facilities. Most physician practices have already been required to meet this standard. For those few that did not, under [Section 1557](https://www.gpo.gov/fdsys/pkg/PLAW-113STAT589/pdf/PLAW-113STAT589.pdf), facilities or programs that are constructed or altered after January 18, 2018 must comply with the [2010 ADA Standards for Accessible Design](https://www.ada.gov/2010standards.pdf).

In addition, all buildings, including those built before the ADA went into effect, are subject to accessibility requirements for existing facilities. Under Title III and Vermont Public Accommodations law, existing facilities are required to remove architectural barriers where such removal is readily achievable. Examples of architectural barriers include inaccessible parking spaces, entrances, door handles and registration counters. 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. This same program accessibility standard generally also applies under Section 504 and Vermont public accommodations law.

Under Section 1557, online services must also be accessible, unless doing so would result in an undue financial burden. This means that patient portals, online forms, online appointment systems, electronic billing and other online services must be accessible to individuals with disabilities, such as vision impairments, unless doing so would impose an undue financial burden to the practice. See Section 1557 Summary, Ensuring Effective Communication with and Accessibility for Individuals with Disabilities.

For more information, See the ADA Guidance Access to Medical Care for Individuals with Mobility Disabilities. This technical assistance publication provides guidance for medical care providers on the requirements of the ADA in medical settings with respect to people with mobility disabilities, which include, for example, those who use wheelchairs, scooters, walkers, crutches, or no mobility devices at all. See also the ADA Guide for Small Businesses for more information about making parking lots, entrances and other structures accessible.

**LIMITED ENGLISH PROFICIENCY & INTERPRETER REQUIREMENTS**

Vermont physician practices are required under federal and state laws to provide interpreters for patients with limited English proficiency and for those who are deaf or hard of hearing. There are very limited exceptions, discussed below. Use of interpreters or translation services may be necessary to ensure that patients can give informed consent. There are several sources for interpreters that physicians can use, including phone-based interpreters, in-person interpreters and online translation services. Vermont’s Medicaid program does reimburse physicians for the cost of providing an interpreter to patients.

*What are the legal requirements for a physician practice to provide language interpretation services for patients with limited English proficiency (LEP)?*

The following is a summary of applicable federal and state laws.

**Civil Rights Act/Affordable Care Act**

*Title VI of the Civil Rights Act of 1964* states that no person shall be subjected to discrimination on the basis of race, color or national origin under any program or activity that receives federal financial assistance. 42 U.S.C.A. § 2000d. In 2003 the Department of Health and Human Services issued *Guidance to Federal Financial Assistance Recipients* Regarding Title VI Prohibition Against National Origin Discrimination Affecting Limited English Proficient Persons to clarify the responsibilities of health care providers under Title VI.

In May 2016, the US Department of Health and Human Services (HHS) issued a Final Rule implementing a prohibition on discrimination by health care services found in *Section 1557* of the Affordable Care Act (ACA). Section 1557 builds on long-standing Federal civil rights laws that have already applied to most health care practices, such as Title VI of the Civil Rights Act. The requirements of the Section 1557 Final Rule and Title VI are consistent and should generally be reviewed together – though Section 1557 provides further detail in several areas.
The Section 1557 Final Rule applies to “every health program or activity, any part of which receives Federal financial assistance.” HHS expects that this applies to almost all physician practices in the nation as Federal financial assistance includes, but is not limited to: state Medicaid payments, “meaningful use” payments, National Health Service Corp funding, NIH research funding and CMS gain-sharing payments. Physician practices that exclusively receive Medicare Part B payments are not included, but practices should conduct a thorough review of their funding sources and consult an attorney before deciding they are exempt.

Under Title VI and Section 1557, practices must ensure that they take reasonable steps to provide meaningful access to each individual with limited English proficiency eligible to be served.

There is no one-size-fits all solution for the type of interpretation (verbal) or translation (written) services that must be provided by the practice and HHS acknowledges that the services provided by a large facility may not be appropriate for a small practice. For verbal interpretation, practices can consider options including hiring bilingual staff, using in-person interpreters or contracting with phone or video interpretation services. Written translation can range from translation of an entire document to translation of a short description of the document to an oral translation by an interpreter. Practices should conduct a review to determine if specific “vital” documents or portions of documents should be translated into the language of any frequently-encountered LEP groups.

Language access services must be provided free of charge, be accurate and timely, and protect the privacy and independence of the patient; a patient cannot be required to provide his or her own interpreter.

Interpreters and translators must also be qualified. Bilingual staff, interpreters and translators must all adhere to ethical principles, including confidentiality, demonstrate proficiency in the appropriate languages and be effective, accurate and impartial. Adult family can only be used in emergencies involving an imminent threat to health or safety and when no other interpreter is available OR if the patients specifically requests the family member and it is appropriate under the circumstances. In many circumstances family members are NOT competent to provide quality, accurate interpretation. Minor children can only be relied on if there is an imminent threat to safety or health. Interpretation over video must be capable of providing high quality images.

In examining compliance on a case-by-case basis, the Office of Civil Rights will examine a number of factors to determine if reasonable steps are being taken to ensure meaningful access to LEP patients. The factors may include: the length and complexity of a given communication; the prevalence of the language spoken by the patient; the frequency the provider encounters the language; whether the provider weighed the patient’s preferences; and the cost of language assistance services/resources available to the practice. The primary factor OCR will consider is whether the practice has created a language access plan. For more information on a language access plan, see the section on Non-Discrimination/Section 1557, above.
Under Section 1557, practices also have procedural requirements regarding non-discrimination. Beginning October 16, 2016 practices need to post a notice of non-discrimination in English and shorter “taglines” in non-English languages regarding the availability of language assistance services (1) in their practice, (2) on the practice website and (3) in any significant publications or communications. Practices employing 15 or more individuals also need to create a grievance procedure and name a compliance coordinator for Section 1557’s requirements, effective July 2016. For more information, see the section on Non-Discrimination/Section 1557, above.

Enforcement by the Office of Civil Rights is carried out through compliance reviews, complaint investigations and providing technical assistance and guidance.

To help understand how to comply with the Title IV Guidance, HHS published a Summary, Fact Sheet and FAQs here. For more information about Section 1557, see the OCR website here.

**Vermont Patients’ Bill of Rights**
In a hospital inpatient setting, “a patient who does not speak or understand English has a right to an interpreter if the language barrier presents a continuing problem to patient understanding of care and treatment.” Failure to comply with any provision of the Patients’ Bill of Rights may constitute a basis for disciplinary action against a physician by the Board of Medicine. 18 VSA § 1852 (a)(15).

**Vermont Public Accommodations**
Vermont law states that “An owner or operator of a place of public accommodation or an agent or employee of such owner or operator shall not, because of the race, creed, color, national origin…of any person, refuse, withhold from or deny to that person any of the accommodations, advantages, facilities and privileges of the place of public accommodation.” Public accommodations include any facility in which services are offered to the general public. Failure to comply may result in an action for civil penalties either through the Vermont Human Rights Commission or a private court action. 9 VSA § 4502.

*What are the legal requirements for a physician practice to provide language interpretation services for patients who are deaf of hard of hearing?*
The following is a summary of applicable federal and state laws.

**Americans with Disabilities Act (42 USC §§ 12131-12134)/Affordable Care Act**
The ADA prohibits public accommodations from discriminating against people with disabilities. This entails furnishing auxiliary aids and services when necessary to ensure effective communication with individuals with hearing impairments, including in certain situations, providing an interpreter who is able to interpret sign language effectively, accurately, and impartially.

The ADA provides an exception for services that would impose an undue burden or would fundamentally alter your offered services. The fact that an interpreter’s charge exceeds the fee for the visit is not alone considered an undue burden.
Private individuals may bring lawsuits in which they can obtain court orders to stop discrimination. Individuals may also file complaints with the attorney general, who is authorized to bring lawsuits in cases of general public importance or where a pattern or practice of discrimination is alleged. In these cases, the attorney general may seek monetary damages and civil penalties.

For a helpful summary of physician practice obligations under the ADA, see the National Association for the Deaf Questions and Answers for health care providers.

Section 1557 of the Affordable Care Act, and accompanying regulations, reiterate the requirement that physician practices ensure that communications with individuals with disabilities are as effective as communications with others. In addition, physician practices covered by the law must ensure that any online services available to patients are accessible to individuals with disabilities. For more information, see the section on Non-Discrimination/Section 1557, above.

Vermont Patients’ Bill of Rights
In a hospital inpatient setting, “a patient who is hearing impaired has a right to an interpreter if the impairment presents a continuing problem to patient understanding of care and treatment.” Failure to comply with any provision of the Patients’ Bill of Rights may constitute a basis for disciplinary action against a physician by the Board of Medicine. 18 VSA § 1852 (a)(15).

Vermont Public Accommodations
The law says that, “A public accommodation shall provide an individual with a disability the opportunity to participate in its services, facilities, privileges, advantages, benefits and accommodations [and] shall make reasonable modifications in policies, practices, or procedures when those modifications are necessary to offer goods, services, facilities, privileges, advantages, or accommodations to individuals with disabilities….“ 9 VSA § 4502 (c)(1), (c)(5) Failure to comply may result in an action for civil penalties either through the Vermont Human Rights Commission or a private court action.

Do the interpretation requirements affect the requirement to provide informed consent to patients?
The Vermont Patients’ Bill of Rights provides that “the patient has the right to receive from the patients’ physician information necessary to give informed consent prior to the start of any procedure or treatment.” 18 VSA §1852 (a)(4).

Additionally, failing to obtain informed consent may be a factor in medical malpractice litigation, although there are some exceptions. For the purposes of medical malpractice actions, "lack of informed consent" is defined as a failure to disclose to the patient reasonably foreseeable risks, benefits, and alternatives to the proposed treatment, in a manner permitting the patient to make a knowledgeable evaluation. In addition, patients are entitled to reasonable answers to specific questions about foreseeable risks and benefits. 12 V.S.A. §1909.

Using interpreters, translation services or other communication aids and services may be necessary to ensure that patients with limited English proficiency (LEP), deaf, or hard-of-hearing
patients receive appropriate information about the proposed treatment to enable them to give informed consent to treatment.

**Is a written authorization required to disclose health information to interpreters?**

An interpreter/interpreter service or a bilingual employee is covered under the health care operations exception for purposes of HIPAA, and the patient’s written authorization to disclose protected health information is not required. Providers who utilize a private company for interpretation or translation on an ongoing contractual basis should put in place a HIPAA Privacy Rule business associate agreement.

In other situations, with disclosures to family members, friends, or other persons identified by an individual as involved in his or her care, when the individual is present, the health care professional or facility may obtain the individual’s agreement or reasonably infer, based on the exercise of professional judgment, that the individual does not object to the disclosure of protected health information to the interpreter.

Authorization is also not required when using a TRS phone service for an individual who has a hearing impairment.

For more information, see the HHS FAQs on disclosing protected health information to an interpreter or when using a TRS service.

**Is there a need for language interpretation services in the Vermont population?**

According to US Census data released in October 2015, Vermont had roughly 31,000 people over the age of 5 in Vermont listed as speaking a language other than English and close to 9,000 who spoke English “less than very well.” The top 15 language spoken “less than very well” are: French (1,806 speakers), Spanish (1,473), Vietnamese (752), Chinese/Mandarin (650), Nepali (570), Serbocroatian (Serbian language) (455), German (305), Cushite (Oromo language - African) (275), Arabic (168), Russian (168), Tagalog (131), Italian (124), Thai (123), Japanese (121) and Portuguese (120). They are followed by Korean (114), Serbian (110) and Russian (103).

**Is there a need for interpretation services for patients who are deaf or hard of hearing in Vermont?**

According to the 2014 American Community Survey, conducted by the US Census Bureau, of the approximately 620,000 people living in Vermont, 181 under the age of 5, 10,000 between 18-64 and 15,000 over the age of 65 have a hearing difficulty.

**Where can I find more information about working with interpreters?**

There are a number of resources available for practices seeking more information about working with interpreters, including:

- AMA Office Guide to Communicating With Limited English Proficient Patients [PDF]
- Agency of Human Services Division of Vocational Rehabilitation page on hiring and working with sign language interpreters.
AAFP, Family Practice Management article, Getting the Most from Language Interpreters and American Family Physician article, Appropriate Use of Medical Interpreters

Where can I find interpreter services?
Below is a non-exclusive list of service providers and contact information. Please note that the Vermont Medical Society does not endorse any of these services and this information is subject to change.

**Limited English Proficiency**

Organization: Language Line Services  
Phone: 1-877-866-3885  
Web: [www.languageline.com](http://www.languageline.com)  
Fee Structure: Per minute charges  
Type of Service: Over-the-phone interpretation, online document translation

Organization: Cryacom Language Solutions  
Phone: 1-800-481-3289  
Web: [http://www.cyracom.com/](http://www.cyracom.com/)  
Fee Structure: Per minute charges  
Type of Service: Over-the-phone interpretation, online document translation

Organization: Vermont Refugee Resettlement Program  
Phone: 1-802-654-1706  
Web: [http://refugees.org/serving-the-uprooted/services/interpreting-services/](http://refugees.org/serving-the-uprooted/services/interpreting-services/)  
Fee Structure: Varies based on services provided  
Type of Service: In-person interpretation, phone/audio, translation services

**Sign Language Specific**

Organization: Vermont Relay Service  
Phone: 1-866-931-9028  
Web: [http://www.vermontrelay.com/](http://www.vermontrelay.com/)  
Fee Structure: The Vermont Telecommunications Relay Service is a free service for all Vermonters, connecting deaf, hard-of hearing, deaf-blind and speech-disabled individuals with users of regular telephones.

Organization: Vermont Interpreter Referral Service  
Phone: 1-802-254-3920  
Web: [www.virs.org](http://www.virs.org)  
Fee Structure: Finder fee for each interpreter in addition to negotiated interpreter fee  
Type of Service: In-person interpretation

Organization: Vermont Division of Vocational Rehabilitation  
Web: [http://vocrehab.vermont.gov/programs/rcd/interpreters](http://vocrehab.vermont.gov/programs/rcd/interpreters)
Can I charge patients for the cost of using an interpreter?
No, the patient cannot be held responsible for the additional cost of using an interpreter to access services.

Do insurers reimburse for interpreter services?
To our knowledge, only Medicaid reimburses for interpreter services at this time. A provider who pays for interpreter services for Vermont Medicaid members may bill procedure code T1013 for each 15 minutes of paid interpreter services provided, on site or via telephone. The rate on file for the billing code is $15 per 15-minute increment. This may include interpreter service outside of the actual healthcare provider encounter in order to fill out forms or review information/instructions. The provider may not bill Vermont Medicaid or the member for a missed appointment per federal policy. Claims are submitted using the CMS 1500 claim form with HCPCS code T1013.

Is there any way to bill for the extra time spent with a patient in the office when a translator is involved?
The only way to account for this extra time is to submit one of the prolonged services codes (99354-99357), which requires that the face-to-face time spent with the patient extend at least 30 minutes beyond the typical time associated with the appropriate CPT services. Note that Medicare and most other payers will not pay for the services of the translator even if they are willing to pay for the extra visit time associated with using a translator.

About the Author

Jessa Barnard is the Vermont Medical Society’s General Counsel and Vice President for Policy. She is a native of Bennington and holds a Bachelor’s degree from Dartmouth College and a law degree from Stanford University School of Law. She served as VMS’ policy specialist from 2002 to 2005. Following her graduation from law school, she founded a program in San Jose, California to address the legal barriers to health stability facing low income individuals living with diabetes. She then spent four years with the Maine Medical Association, most recently as their Associate General Counsel, representing physicians in Augusta and addressing their legal and regulatory concerns. She is experienced in health care law, policy and regulation and is a frequent speaker on topics including health reform, advocacy and issues in health law.
PATIENT-PHYSICIAN RELATIONSHIPS

Topics Covered in this Chapter:
Initiation & Termination of The Patient-Physician Relationship
Treating Self or Family
Patients’ Rights: Residents in Long-Term Care Setting
Patients’ Rights: Hospital Setting
Patients’ Rights: Palliative Care and Pain Management
About the Author

Professional Courtesy [To Linda?]

By Jessa Barnard, Esq.
Vermont Medical Society

INITIATION AND TERMINATION OF THE PATIENT-PHYSICIAN RELATIONSHIP

What constitutes the beginning of a patient-physician relationship?
Vermont law does not specify what constitutes the beginning of a patient-physician relationship. The CPT Manual defines a new patient as a patient who has not received any professional services from the physician or another physician of the same specialty who belongs to the same group practice within the past three years. Professional services are defined as face-to-face professional services or evaluation and management services. See CMS FAQ 1969. The AMA Code of Ethics states that a “patient-physician relationship exists when a physician serves a patient’s medical needs.” Opinion 1.1.1.

As a general precaution, a physician should consider that the relationship has begun at any point when the physician or a member of the physician’s practice has offered to provide a medical service, unless it has been explicitly stated in writing to the patient that the physician is providing care only on an interim or emergency basis and that the patient should continue looking for another physician. If, after offering to provide services to a patient, the physician decides that he or she does not want to or is unable to serve that patient, the physician should follow the process for discharging a patient as described below.

May physicians refuse to see a specific patient even if the practice is accepting other new patients?
Yes, according to AMA Ethics Opinion 1.1.2, regarding prospective patients, physicians have the prerogative to choose whether to enter into a patient-physician relationship with any individual. However, a physician must respond in cases of emergency and cannot discriminate based on race, color, national origin, sex (including sexual orientation or gender identity), age, disability or other personal or social characteristics that are not clinically relevant to the individual’s care. See Chapter on Non-Discrimination, above. The physician should also be aware of any contractual relationship - for example, a participation agreement with an insurer - that requires him or her to treat certain patients.

As a general guideline, when considering his or her ability to accept additional patients, the physician should balance a prospective new patient’s medical needs with the physician’s skills...
and capacity and the needs of the physician’s other patients. According to the AMA’s Code of Medical Ethics, “greater medical necessity of a service engenders a greater obligation to treat.”

Potential reasons to refuse to initiate treatment may include:

- the treatment request is beyond the physician’s competence;
- the treatment request is scientifically or medically invalid or cannot reasonably be expected to achieve the intended clinical benefit;
- the treatment request is incompatible with the physician’s personal, religious, or moral beliefs, in keeping with ethical guidelines on exercise of conscience;
- the physician lacks the resources to provide safe and competent care; or
- the individual is abusive or threatens the physician, staff or other patients and it is not an emergency situation.

See [AMA Ethics Opinion 1.1.2](#).

A practice should consider developing an office policy on acceptance of patients, including examples of reasons for refusing to accept a patient, the process for informing the patient that they will not be accepted into the practice and a statement of nondiscrimination.

### May physicians discharge patients from their care?

Yes, physicians are allowed to end the physician-patient relationship if it is the best option. Such situations may include a patient who is consistently noncompliant, unreasonably demanding, or threatening to you or your staff. It may also be necessary to transition a patient due to the relocation or retirement of a health care professional.

A 1999 Vermont Board of Medical Practice [advisory](#) states that the Board recognizes that the physician-patient relationship may be terminated by either party and that a physician “has the right to terminate the relationship when he or she believes it is best to do so. However, termination... must be done in compliance with the physician’s obligation to support continuity of care for the patient.”

### What are the risks of discharging a patient from care?

If not done properly, a patient can claim that a physician has abandoned his or her care. Abandonment means the termination of a professional relationship between physician and patient at an unreasonable time and without giving the patient the chance to find an equally qualified replacement. Under Vermont law abandoning a patient is considered unprofessional conduct, [26 VSA § 1354 (a)(4)](#), and if reported to the Vermont Board of Medical Practice can trigger an investigation. The Board has the authority to issue warnings, impose fines, or condition, suspend or revoke licenses to practice medicine. A physician may also be at risk for medical malpractice claims if the termination of care occurs at a critical state of treatment and an injury results.

### Are there guidelines to follow when discharging a patient from the care of a practice?

The Vermont Board of Medical Practice and the American Medical Association offer guidance on the appropriate process physicians should follow when discharging a patient from their practice.
In assessing whether a physician has abandoned a patient, the Board will consider the following factors:

- Whether the physician gave the patient at least 30-days notice - in writing - if ending a treatment relationship. The physician does not have to state a reason for ending the relationship, only his or her belief that it is best to do so. The notice should also state clearly whether termination involves only an individual physician or an entire group practices and written notice of termination should be “presented to the patient by a method to ensure that the patient received the notice.” (For example, via certified, return receipt mail).
- Whether the physician provided necessary ongoing and/or emergency treatment during the transition period to a new physician - at least 30 days; and
- Whether the physician promptly transferred records to a new physician, as chosen by the patient, regardless of whether the patient had any outstanding bills with the practice. (It is best practice to provide resources and/or recommendations to help a patient locate another physician of appropriate specialty.)

Note that while 30 days is the minimum transition period advised by the Board of Medical Practice, more time may be necessary based on patient circumstances, such as the complexity of the patient’s ongoing medical needs. AMA Ethics Opinion 1.1.5, states that physicians “must notify the patient (or authorized decision maker) long enough in advance to permit the patient to secure another physician [and] must facilitate transfer of care when appropriate.” See also Ethics Opinion 1.1.3: “the physician will not discontinue treating [a patient] when further treatment is medically indicated without giving them sufficient notice and reasonable assistance in making alternative arrangements for care.”

A practice should consider developing an office policy on termination of patients. The policy should include examples of reasons for terminating a patient such as:

- The patient is displaying hostility or inappropriate behavior to staff, physicians or other patients;
- The patient is refusing to cooperate with staff in diagnosis or treatment, or refusing to follow treatment instructions; or
- The patient is making unreasonable demands on staff or physicians.

The policy should clarify the process for providing notice of termination to patients and should also clarify that the practice will continue to comply with ethical and legal requirements including:

- Providing care in an emergency;
- Continuity of care; and
- Prohibition of discrimination on the basis of race, color, national origin, sex (including sexual orientation or gender identity), age, disability or other personal or social characteristics that are not clinically relevant to the individual’s care. See more information in the Chapter on Non-Discrimination.
For more practical guidance on terminating the physician-patient relationship, including a sample termination policy, see Medical Mutual Insurance Company of Maine’s Practice Tip on Termination of the Physician/Patient Relationship.

What immediate and longer-term steps can a practice take when a patient is dangerous, threatening or inappropriate for the practice?

If a patient is dangerous or threatening to staff or other patients you can ask him or her to leave your office immediately, and call law enforcement if the patient refuses. If necessary, you may obtain a court order such as a “no trespass” order or a restraining order that will bar the patient from your practice. The practice can also initiate steps to terminate the patient from the practice, as discussed above. During the notice period while the patient finds another physician, you may need to make arrangements to see the patient at a safe location such as an emergency room, should he or she require treatment.

If the patient is served by a mental health agency or other social support agency, you can request that the patient not come to your office without one or more staff persons from the agency accompanying the patient.

TREATING SELF OR FAMILY

AMA Ethical Opinion 1.2.1 advises physicians generally not to treat themselves or family members as it “poses several challenges for physicians, including concerns about professional objectivity, patient autonomy and informed consent.” It may be acceptable in limited circumstances, such as emergency situations or isolated settings when there is no other qualified physician available or for short-term, minor problems. When treating self or family members, physicians have an obligation to document treatment and care and convey information to the patient’s primary care physician, avoid providing sensitive or intimate care, especially to a minor, and recognize that family may be reluctant to state a preference for another physician. See Opinion 1.2.1 for further guidance.

Under Vermont Board of Medical Practice rules, it also is unacceptable medical practice and unprofessional conduct for a licensee to prescribe Schedule II, III, and IV controlled substances to him or herself or to a member of his or her immediate family. There is an exception to allow prescribing to family members in a bona fide emergency, of short-term and unforeseeable character. "Immediate family" includes the following: a spouse (or spousal equivalent), parent, grandparent, child, sibling, parent-in-law, son/daughter-in-law, brother/sister-in-law, step-parent, step-child, step-sibling, or any other person who is permanently residing in the same residence as the licensee. See Board Rules Part 4.3

PATIENTS’ RIGHTS: RESIDENTS IN LONG-TERM CARE SETTING

Both the federal and Vermont governments have adopted certain regulations to promote quality care in nursing facilities. Under these regulations, all nursing facilities are required by law to have written policies called the Nursing Home Residents’ Bill of Rights. These rights were implemented to protect the resident from abuse and neglect and to provide the resident with an opportunity to participate in his or her care.
What are some of the individual rights specific to a long-term care setting?

Vermont’s Nursing Home Residents’ Bill of Rights is found at 33 VSA § 7301. Further detail can be found in the Department of Disabilities, Aging and Independent Living Licensing and Operating Rules for Nursing Homes. Patients’ rights include, but are not limited to:

- The right to be fully informed, prior to or upon admission and during the stay, of the rules of the facility, his or her rights as a resident in the facility, the services with related charges available in the facility, about Medicare and Medicaid eligibility and about eligibility for hospice services.
- The right to choose one’s own personal physician, be fully informed about his or her care and treatment, to participate in planning or changing the care or treatment, and to refuse to participate in experimental research.
- The right to refuse care or treatment, including the right to discharge him or herself from the facility.
- The right to be transferred or discharged only for medical reasons or for the resident’s welfare or that of other residents or for nonpayment.
- The right to return to the first available bed after hospitalization if the facility is able to meet his or her needs, and to retain his or her bed in the facility while absent due to hospitalization not exceeding ten successive days.
- The right to professional assessment and management of pain.
- The right to be free from mental and physical abuse and free from chemical and physical restraints except as authorized in writing by a physician.
- The right to confidentiality of personal and clinical information, and to approve or refuse their release to any individual outside the facility, except, in case of his or her transfer to another health care institution, or as required by law or third-party payment contract.
- The right to send and receive mail unopened, to have access to a private use of a telephone, to receive visitors, to voice grievances, to vote, to participate in council meetings, and to manage his or her personal financial affairs.
- The right to review current and past state and federal survey and inspection reports of the facility.

Examples of violations include, but are not limited to:

- Failure to provide the resident with a copy of his or her rights and responsibilities or any changes made to such rights.
- Failure to disclose all cost and charges to the resident.
- Not permitting the resident to manage his or her finances. Restraining a resident, except in an emergency, without an order from a physician.
- Discussing a resident’s medical condition with someone who is not involved with the resident’s care.
- Not providing the resident an opportunity to review the latest inspection report.

These State-established rights generally follow the rights established in the Federal 1987 Nursing Home Reform Act (42 USC 1396r(c) and 42 USC §1395i-5(c)) and federal regulations for skilled nursing facilities, 42 CFR Part 483. Federal law includes: the right to freedom from abuse, mistreatment, and neglect; the right to freedom from physical restraints; the right to
privacy and confidentiality; the right to reasonable accommodation of medical, physical, psychological, and social needs; the right to participate in resident and family groups; the right to be treated with dignity; the right to exercise self-determination; the right to communicate freely; the right to participate in the review of one's care plan, and to be fully informed in advance about any changes in care, treatment, or change of status in the facility; and the right to voice grievances without discrimination or reprisal.

**How are these rights implemented?**
The governing body of the facility establishes written policies regarding these rights and responsibilities of residents and, through the administrator, is responsible for the development of and adherence to procedures implementing such policies. All staff of the facility ensures compliance with these rights. 33 VSA § 7301 (1).

**How are these rights communicated to the residents?**
The facility is required to post a summary of the obligations of the facility to residents in clear language, in easily readable print and posted conspicuously in a public place on each floor of the facility. The notice shall summarize the facility’s grievance procedure and directions for contacting the ombudsman program. A readable copy of the notice shall be presented to each resident on admission together with an oral explanation of the rights, grievance procedure, and directions for contacting the ombudsman program. 33 VSA § 7303. (Under the federal Older Americans Act, every state is required to have an Ombudsman Program that addresses complaints and advocates for improvements in the long term care system.)

**What if a right is violated?**
Every nursing home must establish a grievance mechanism that allows for residents to file a grievance. Complaints that cannot be resolved by the facility grievance procedure within 7 days must be referred to the Long-Term Care Ombudsman program. 33 VSA § 7302.

A resident can also file complaints with a variety of state and federal agencies, including the Vermont Department of Disabilities, Aging and Independent Living, which regulates and surveys nursing homes: 1-800-564-1612 or 280 State Drive HC 2 South Waterbury, Vermont 05671-2060. Residents can also contact the Vermont Long Term Care Ombudsman’s Project at 1-800-889-2047.

To establish compliance with the 1987 Nursing Home Reform Act, the law established a state-based certification and survey process. If found in violation of the law, nursing homes, their owners, and administrators can be fined, have their licenses suspended or revoked, and lose their right to payment by Medicaid or Medicare.

**If the resident is not capable of understanding these rights, who may exercise these rights on his or her behalf?**
Under 33 VSA §7306, the rights and obligations under the Nursing Home Residents’ Bill of Rights devolve to a resident’s reciprocal beneficiary, guardian, next of kin, sponsoring agency, or representative payee (except when the facility itself is a representative payee) if the resident:

- has been adjudicated incompetent;
• has been found by his or her physician to be medically incapable of understanding or exercising the right; or
• exhibits a communication barrier.
The facility however will make every reasonable effort to communicate the rights and obligations directly to the resident.

**PATIENTS’ RIGHTS: HOSPITAL SETTING**

Vermont has adopted a Bill of Rights for Hospital Patients that establishes certain rights for an individual who is an inpatient at a Vermont hospital. See 18 VSA §1852.

**What are some of these rights?**

These include, but are not limited to, the right to:

- Considerate and respectful care.
- Obtain current and understandable information about his or her diagnosis, treatment and prognosis.
- Be informed of the name and position of the doctor in charge of the care.
- Receive all the information necessary to give informed consent for any proposed procedure or treatment, including risks and alternative options.
- Refuse treatment and be told what effect this may have on the patient’s health.
- Privacy while in the hospital and confidentiality of all information and records regarding the care.
- Receive all reasonable medical services provided by the hospital, at the request of the physician.
- Know the identity and professional status of individuals providing services.
- Whenever possible, parents/guardians of children and family members of terminally ill patients have the right to remain with the patient 24 hours per day.
- Receive an itemized bill and explanation of all charges.
- Know what hospital rules and regulations apply to his or her conduct as a patient.
- Professional assessment and management of pain.
- Written information about availability of and eligibility for hospice services.
- To an interpreter if the patient’s predominant language is not English or is hard of hearing.

Hospitals are also required to make public the maximum patient census and the number of registered nurses, licensed practical nurses, and licensed nursing assistants providing direct patient care in each unit during each shift. The information must be posted in a prominent place that is readily accessible to patients and visitors in that unit. 18 VSA §1854.

**How are these rights communicated to patients?**

A summary of the hospital’s obligation written in clear language and readable print must be distributed to each patient upon admission and posted conspicuously at each nurse’s station throughout the hospital. 18 VSA §1852(c). The notice must inform patients of the hospital grievance procedure and that patients may also contact the licensing agency or the Board of Medical Practice with a complaint.
What if a right is violated?
Failure to comply with any of these rights may constitute a basis for disciplinary action against a physician. A complaint may be filed with the Board of Medical Practice. 18 VSA §1852(b).

PATIENTS’ RIGHTS: PALLIATIVE CARE AND PAIN MANAGEMENT

18 VSA §1871 establishes a patient's bill of rights for palliative care and pain management. Under the statute:

- A patient has the right to be informed of all evidence-based options for care and treatment, including palliative care, in order to make a fully informed patient choice.
- A patient with a terminal illness has the right to be informed by a clinician of all available options related to terminal care; to be able to request any, all, or none of these options; and to expect and receive supportive care for the specific option or options available.
- A patient with pain has the right to request or reject the use of any or all treatments in order to relieve his or her pain.
- A patient with a chronic condition has the right to competent and compassionate medical assistance in managing his or her physical and emotional symptoms.
- A pediatric patient with a serious or life-limiting illness or condition has the right to receive palliative care while seeking and undergoing potentially curative treatment

ABOUT THE AUTHOR

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PHARMACEUTICAL ISSUES

Topics Covered in this Chapter:
Drug Diversion
About the Author

By: Jason Turner
Office of the Attorney General

DRUG DIVERSION: PATIENT FRAUD

What steps can I take to reduce drug diversion and protect against drug fraud?
You can develop an office or institutional policy on prescribing controlled substances for treatment of chronic conditions and apply it consistently. The policy could address the following components:

- Initial patient evaluation and risk stratification
- Use of an informed consent form for treatment with controlled substances;
- Use of agreements for long-term controlled substances therapy;
- Individualized treatment plans;
- Objectives for evaluation of treatment;
- Regular monitoring of treatment progress;
- Periodic Drug Testing;
- Plan for response to aberrant behavior
- Consultation with pain or addiction specialist;
- Numbered and watermarked prescription pads;
- Copies of prescriptions kept in the chart;
- Collaboration with pharmacists, hospitals and other physicians treating the same patient; and
- Requiring a photo ID and maintaining a copy in the chart.

Are sample opiate treatment contracts and opiate treatment consent forms for pain and addiction treatment available?

The Nation Institute on Drug Abuse at the National Institutes of Health has sample agreements for treatment with Opioid Medications and for the long-term use of controlled substances: https://www.drugabuse.gov/sites/default/files/files/SamplePatientAgreementForms.pdf
WebMD has a sample agreement on its website: [http://www.webmd.com/pain-management/pain-management-pain-treatment-agreement#1](http://www.webmd.com/pain-management/pain-management-pain-treatment-agreement#1)

**How can I know if a patient is taking the controlled substances I prescribe?**
If you have concerns, you can ask the patient to agree to substance monitoring to verify that they are taking the type and amount of narcotic that you are prescribing.

**From a liability perspective, what is the most important step a clinician can take when prescribing controlled substances, particularly to patients who have a history of drug abuse, or you consider at risk for diversion?**
Document evaluation, treatment, instructions to patient, and rationale carefully in the medical record.

**How can I be sure that my patient is only obtaining controlled drugs from one physician and one pharmacy?**
Physicians must register with the Vermont Prescription Monitoring System (VPMS) and check the VPMS database on-line to determine if their patients are obtaining controlled substances from other physicians or more than one pharmacy. (See Vermont Prescription Monitoring System Rules: [http://healthvermont.gov/about-us/laws-regulations/rules-and-regulations](http://healthvermont.gov/about-us/laws-regulations/rules-and-regulations))

Prescribers can also request that patients who are Medicaid Beneficiaries be “locked-in” to one pharmacy or prescriber for controlled substances. To request that a Medicaid, VHAP or Dr. Dynasaur patient be “locked-into” a single pharmacy or physician, a prescriber can contact the Department of Vermont Health Access (DVHA) Clinical Division. The clinical division can provide prescribers with “Beneficiary Lock-In Request” forms and can answer any questions regarding the lock-in procedure and process. Because the physician lock-in procedure requires patients to obtain all drugs from one prescriber, some physicians prefer to use only the pharmacy lock-in.

Prescribers may also check with other insurers or their pharmacy benefit managers (PBMs) to determine if they have this type of information available.

**May I disclose drug fraud to law enforcement?**
HIPAA permits disclosure to law enforcement in limited situations. Disclosure is permitted when it is required by state law, such as disclosure of gun shot wounds, or proper legal process, such as warrants, subpoenas, or administrative requests. 45 CFR 164.512(f)(1). Disclosure is permitted when, consistent with applicable law and ethical standards, the disclosure is designed to prevent or lessen a serious and imminent threat to the health or safety of an individual or the public, such as the public health threat represented by the diversion of drugs. A health care professional may also disclose evidence of criminal conduct on the premises. 45 CFR 164.512(f)(5). Other more limited exceptions may apply for the victim’s of crime or emergencies. Finally, disclosures to law enforcement who are working for a health oversight agency (Medicare/Medicaid agency or investigations, professional boards) for health oversight activities are permitted. 45 CFR 164.512(d). Whether a particular situation meets one of these exemptions is dependent on the specific facts of the situation. We recommend that practitioners consult with an attorney, their
medical malpractice carrier or risk manager in making a determination of whether one of these exceptions apply.

Vermont law has an exception to the patient privilege for drug fraud. It states that information communicated to a physician in an effort to unlawfully procure a regulated drug will not be considered a privileged physician - patient communication. 18 V.S.A. § 4223(b). Misrepresentation, forgery, alteration of prescriptions, concealment of a material fact, or use of a false name or address are all examples of fraud that would be covered under this law. To report information about patients to law enforcement, a HIPAA exception, (see above) is required, in addition to the Vermont law’s waiver of the patient privilege. Once again, we encourage practitioners to consult with risk management, malpractice carriers or an attorney.

Under both HIPAA and Vermont law, disclosure must be limited to the minimum amount of information needed to enable law enforcement to investigate the drug fraud, or prevent the harm from occurring. Detailed information about the patient’s medical condition or treatment should not be disclosed.

Finally, AMA Ethical Opinion E-3.2.1 on Confidentiality provides that the obligation to safeguard patient confidences is subject to certain exceptions which are ethically and legally justified because of overriding considerations. Where a patient threatens to inflict serious bodily harm to another person and there is a reasonable probability that the patient may carry out the threat, the physician should take reasonable precautions for the protection of the intended victim, including notification of law enforcement authorities.

May I disclose suspected diversion or drug abuse to law enforcement if I work in a program that treats patients for drug abuse or addiction?

Programs that provide alcohol or drug abuse diagnosis, treatment or referral for treatment are subject to the federal regulation governing confidentiality of alcohol and drug abuse patient records. 42 CFR Part 2. These rules provide strict protection for patient confidentiality and include specific consent forms that must be used to disclose information about drug treatment.

Under 42 CFR Part 2, you may disclose if the patient has consented in writing and the consent form meets the specific requirements of the federal regulation. The patient may revoke the consent at any point, however. You may also disclose if law enforcement presents a signed court under authorizing disclosure under 42 CFR Part 2.

Under 42 CFR Part 2, you may also disclose certain crimes or threats to commit crimes without patient consent. These include crimes committed on the premises of your office or program, crimes against staff or threats to commit crimes on your premises or against your staff. In these limited circumstances, regardless of patient consent, you may disclose to law enforcement, the patient’s name, address, whereabouts, the circumstances of the incident and the patient status of the patient who committed or threatened to commit a crime.

Substance Abuse Confidentiality Regulations FAQ: https://www.samhsa.gov/about-us/who-we-are/laws/confidentiality-regulations-faqs
What steps can I take if I believe a patient whom I am treating for drug abuse, dependence or addiction is diverting drugs?

You can request that the patient sign a written treatment agreement, specifying that if the patient fails to comply with the contract, you will no longer prescribe controlled substances for the patient. You can also require the patient to agree to urine monitoring as a condition of continuing to receive treatment with controlled substances from your practice. Finally, you can discharge the patient from your practice provided the requirements of the VBMP policy on Termination of the Physician-Patient Relationship are met.

- See also the section of this Guide on Terminating the Physician-Patient Relationship.

ABOUT THE AUTHOR

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PRACTITIONER HEALTH & RECOVERY PROGRAMS

Topics Covered in this Chapter:
Vermont Practitioner Health Program
Alternative Program for Nurses
Joint Commission Requirements
About the Author

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VERMONT PRACTITIONER HEALTH PROGRAM

Is there a statewide recovery program for physicians and other Board of Medical Practice licensees in Vermont?
The Vermont Practitioner Health Program (VPHP) was established by the Vermont Medical Society in 1999. It offers a monitoring and recovery program for Vermont physicians and other practitioners licensed by the Vermont Board of Medical Practice (VBMP) who are impaired or at risk of impairment by the use of drugs, including alcohol. Eligible clinicians include physicians (MDs), physician assistants, podiatrists, radiologist assistants and anesthesiologist assistants. Osteopathic physicians (DOs) are also eligible. Between 2000 and 2016, VPHP has served 245 participants.

See the program brochure here.

How is the VPHP program operated?
The VPHP program was established by the Vermont Medical Society in 1999 as a peer review committee of the VMS. The Program is managed by a physician medical director and a case management committee, made up of physicians, physician assistants and podiatrists -- a number of whom have personal experience with recovery. Day to day operations are handled by Vermont Medical Society staff.

What services does VPHP offer?
VPHP will assign a case management team to a participant, made up of the medical director and one other member of the case management committee. The team works with the participant to design an individual monitoring and recovery program for the participant. VPHP does not provide individual treatment services directly, but supervises the development of individual recovery and monitoring programs for participants. The elements of a participant’s program are spelled out in a monitoring agreement entered between the participant and the VPHP program.

Typical programs will include evaluation, addiction treatment, ongoing testing for substances and participation in support groups such as A.A., N.A. or Caduceus. Participants are required to disclose their substance-related illness to treatment professionals and those who are working are required to have a workplace monitor.
VHP also offers other services as needed, including intervention, intake evaluation, and information and referral, and also conducts regular outreach to educate the medical profession about the program and issues of impairment.

**What is the relationship between VPHP and the Vermont Board of Medical Practice?**

VPHP is a separate entity from the VBMP. Under a protocol developed by the two organizations, VPHP only discloses the identity of self-referred/non-Board involved participants to the VBMP when there is patient injury, risk of patient injury, unprofessional conduct impacting fitness to practice, criminal acts, uncorrected failure to abide by the terms of the monitoring agreement, or relapse to the use of alcohol or drugs with risk of harm to patients.

In cases where a participant is already subject to a VBMP investigation, order or stipulation, and is referred to the program by the Board, the VPHP program works in coordination with the VBMP to design and implement the recovery program for the health care practitioner.

**How is the confidentiality of the program’s participants protected?**

A number of laws protect the confidentiality of the program’s participants. VPHP has been established as a peer review program of the VMS. (See 26 VSA § 1441). Under Vermont, law peer review activities of the program are confidential and privileged. They are not subject to discovery or introduction into evidence in any civil action. (Information, documents, or records otherwise available from original sources may be discoverable.) 26 VSA § 1443

VPHP also complies with the Health Insurance Portability and Accountability Act of 1996 (HIPAA) and accompanying regulations (45 CFR Part 164) and the Public Health Services Act/42 CFR Part 2 (Confidentiality of Alcohol and Drug Abuse Patient Records).

Under 42 CFR Part 2, records relating to the identity, diagnosis, prognosis, or treatment of any patient maintained in connection with a program relating to substance abuse treatment or rehabilitation may not be disclosed without patient consent, a court order or to medical personnel in a medical emergency. 42 CFR §§ 2.33, 2.51, 2.61. With the exception of the circumstances noted above that trigger reporting to the VBMP, VPHP will not release any information concerning participants to an employer, health plan, hospital or any other party without a signed release from the participant. Participants are typically required to sign authorizations allowing the program to discuss program status and recovery with treating providers, workplace monitors and others involved in the recovery process.

**How is the VPHP program funded?**

The administrative costs of the VPHP program are funded by a surcharge on licensing fees of those professionals licensed by the Vermont Board of Medical Practice, by a VMS foundation dedicated to the health of physicians and their families, by donations from malpractice carriers and by the VMS general operating fund. Participating health care professionals are responsible for their own evaluation, treatment, and monitoring expenses, although some of these expenses may be covered by the participant’s health insurance plan.

**Where can I find more information about the VPHP recovery program?**

The program is operated by the Vermont Medical Society and can be reached at:
ALTERNATIVE PROGRAM FOR NURSES

Is there a similar program for nurses?
In 2004 the Vermont State Board of Nursing established by rule a non-disciplinary Alternative Program for eligible nurses and nursing assistants who are physically or psychologically dependent on alcohol or other drugs. See Vermont Board of Nursing Rules, Part 11. See the program brochure here.

In order to participate, nurses must voluntarily request admission to the program, if requested agree to undergo a comprehensive assessment at their own expense, and agree to comply with a contract prepared by the alternative committee of the nursing board. Part 11.3. The contract sets forth the terms, conditions, costs, and restrictions which the committee deems appropriate for the individual participant. The contract is an Order of the Board with which the individual must comply. Part 11.7(a).

Contract conditions may include: substance abuse counseling and treating professional reports; participation in recovery group meetings; random drug and alcohol testing; abstinence from drug and alcohol use; professional practice only with Program approval, work restrictions and supervision; six-month prohibition on administering controlled substances; and self-assessment reports. See Board of Nursing Alternative Program Purpose and Objectives.

Records pertaining to a nurse’s participation are confidential, except as necessary to ensure compliance with the program requirements, such as to the nurse’s employer to ensure work site monitoring. See Part 11.1(e).

Nurses are not eligible to participate in the program if they: have a pending felony charge or felony conviction related to chemical dependency; have a restricted license for reasons of unprofessional conduct; have diverted controlled substances; have taken or disregarded a substantial risk of harm; present an imminent danger to the public; or have a recent history of chemical dependency with failed treatment. See Part 11.4.

Where can I find more information about the Board of Nursing program?
Information is available from the Vermont Board of Nursing Alternative Program website and confidential phone number: 802-828-1635.
**Does The Joint Commission require hospitals to establish practitioner health programs?**

The Joint Commission (formerly JCAHO) requires each hospital medical staff to implement a process to identify and manage matters of individual health for licensed independent practitioners that is separate from the medical staff disciplinary function. See Standard MS 11.01.01.

According to Standard MS 11.01.01, the purpose of the process is to facilitate rehabilitation, rather than discipline, and to aid a practitioner in retaining or regaining optimal professional functioning, consistent with protection of patients. However, if the practitioner is unable to safely perform the privileges he or she has been granted, the medical staff leadership must take appropriate corrective action.

The process should include the following elements:

- Education of staff about illness and impairment recognition issues;
- Accepting self-referral or referral by other staff;
- Providing referral of the practitioner to internal or external resources for evaluation, diagnosis and treatment of the condition or concern;
- Maintenance of the confidentiality of the practitioner, except as limited by applicable law, ethical obligation, or when the safety of a patient is threatened;
- Evaluation of the credibility of a complaint, allegation or concern;
- Monitoring of the affected practitioner and the safety of patients until the rehabilitation process is complete;
- Reporting to medical staff leadership when a licensed independent practitioner is providing unsafe treatment;
- Initiating appropriate actions when a practitioner fails to complete a required rehabilitation program.

**How are hospitals in Vermont meeting this standard?**

Hospitals may develop their own education and monitoring programs to address licensed practitioner health, or they may utilize statewide programs, such as the Vermont Practitioner Health Program, to meet this requirement. For example, the University of Vermont Medical Center has its own long-standing Practitioner Health and Advocacy Program. More information about forming a medical staff health committee that complies with MS 11.01.01 can be found from the Massachusetts Medical Society Physician Health Services program.

**ABOUT THE AUTHOR**

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their legal and regulatory concerns. She is experienced in health care law, policy and regulation and is a frequent speaker on topics including health reform, advocacy and issues in health law.
PROFESSIONAL LIABILITY

Topics Covered in This Chapter:
- Medical Malpractice
- Statutes of Limitation
- Liability with Respect to Informed Consent
- Liability with Respect to Advance Directives for Health Care, DNR Orders, and COLST Orders
- Liability with Respect to the Duty to Protect the Endangered Act
- Alternative Dispute Resolution
- National Practitioner Data Bank
- Safe Apology Law
- Patient Safety Surveillance and Improvement System

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MEDICAL MALPRACTICE

The vast majority of legal claims filed against health care professionals are those based on allegations of medical malpractice, that is, a claim that a health care professional was negligent in his or her provision of care. However, patients may also assert other types of claims based on the practice of medicine. These claims are infrequent but could include: battery (unauthorized touching of another person resulting in harm); libel or slander (for untruthful reporting of a physical or mental condition); duress or false imprisonment (for detaining a patient in a hospital or medical facility without just cause); and invasion of privacy (such as unlawful use of bodily tissue, etc.).

What do I do if am being sued?
Step 1: Take a deep breath and try to relax. Not to minimize a medical malpractice suit and the stress inherent in being named in one, but life will go on and worse things can and do happen in life.

Step 2: Immediately upon receiving notice of a lawsuit or the potential for a suit, mail a copy of whatever papers you have received to your medical malpractice carrier and call the carrier to advise it of the claim. If you are employed by a health care organization or hospital, the risk manager of that organization should be immediately notified and he or she will assist you.

What is the relationship between my insurance carrier and me?
Fortunately, in Vermont there is almost always an excellent working relationship between insurance carriers, defense counsel, and the physician. The cornerstone of that relationship is trust; the physician must be comfortable with and confident in his or her defense team. In that regard, most medical malpractice insurance carriers will consider a physician’s request for a specific attorney, provided that attorney has expertise in medical malpractice litigation. An important factor in this relationship is an early meeting between the physician and defense
counsel not only to discuss the substance of the case, but also to address all questions and concerns the physician may have.

Must the carrier have my best interest in mind? What happens if our interests diverge?
The carrier has fiduciary and contractual obligations to act in the best interest of its insured. In the rare instances where a divergence of interests develops, e.g., a dispute as to coverage, it is important for the physician to consider retaining personal counsel to guarantee that his or her interests are protected.

Should a physician ever talk to the plaintiff-patient after a lawsuit has begun?
Like it or not, once a lawsuit has been filed the physician and patient/former patient are in an adversarial posture. As a result, there should be no communication between the physician and that former patient, particularly with respect to any subject implicated by the lawsuit. The safest course of action before taking any significant steps after the filing of the lawsuit is for the physician to speak to his or her attorney to obtain guidance and direction.

On a related note, the physician should keep separate from the patient’s chart and clearly marked as “attorney/client communications not for release,” all communications to and from the physician from his or her insurance carrier and/or legal counsel in order to protect the attorney client privilege.

What must a plaintiff-patient prove to recover on a claim of medical malpractice?
In a medical malpractice action brought against a health care professional in Vermont, the plaintiff-patient has the burden of proving:

- The degree of knowledge or skill possessed or the degree of care ordinarily exercised by a reasonably skillful, careful, and prudent health care professional engaged in a similar practice under the same or similar circumstances whether or not within the state of Vermont;
- That the defendant/health care professional either lacked this degree of knowledge or skill or failed to exercise this degree of care; and
- That as a proximate result of this lack of knowledge or skill or the failure to exercise this degree of care the plaintiff suffered injuries that would not otherwise have been incurred.

12 V.S.A. § 1908.

Section 1908 creates an objective, national standard of care in which the defendant’s conduct is to be measured against what a reasonable health care professional in a similar practice would have done in the same or similar circumstances. Stated differently, health care professionals are expected to deliver health care with the same degree of care and skill that is ordinarily possessed and exercised in like cases by professionals in the same general line of practice. The failure to do so is medical malpractice.

This does not mean that a physician is required to be infallible. Utzler v. Medical Center Hosp., 149 Vt. 126, 127 (1987). A physician will not be held liable for malpractice as a result of a “mere error in judgment,” meaning that a physician may choose from several appropriate treatment alternatives and the mere fact that harm results from the physician’s choice of one alternative over the other is not necessarily malpractice. Rooney v. Medical Center Hosp. of Vermont, Inc., 162 Vt. 513, 521 (1994). This also means that the standard of care does not
require a health care professional to guarantee a good result. If the provider meets the standard of care for his or her profession, then he or she will not be found liable for malpractice regardless of the result of the treatment. *Lockwood v. Lord,* 163 Vt. 210, 217 (1994).

**What type of evidence must a plaintiff-patient produce in order to meet her burden of proof on a medical malpractice claim?**

Generally, to meet his or her burden of proof in a medical malpractice action, the plaintiff-patient must present expert medical testimony setting forth:

- The proper standard of medical skill and care;
- That the defendant/health care professional’s conduct departed from that standard; and
- That this conduct was the proximate cause of the harm complained of.

*Senesac v. Associates in Obstetrics and Gynecology,* 141 Vt. 310, 313 (1982); *Utzler v. Medical Center Hosp.,* 149 Vt. 126 (1987). This expert should be another health care professional who is familiar with the specialty and/or type of practice in which the defendant/health care professional is engaged and will offer testimony on the three previously enumerated issues.

The reason expert testimony is generally required is that “the human body and its treatment are extraordinarily complex subjects requiring a level of education, training and skill not generally within our common understanding.” *Taylor v. Fletcher Allen Health Care,* 2012 VT 86, ¶ 9 (2012). An exception to this general rule exists, however, in those cases where the violation of the standard of medical care is so apparent to be comprehensible to any ordinary lay person. *Largess v. Tatem,* 130 Vt. 271, 278-79 (1972); *Larson v. Candlish,* 144 Vt. 499, 502 (1984). For instance, when a health care professional treats the wrong patient or body part, the plaintiff is not required to present expert testimony to support his or her claim since the violation of the standard of care is obvious to anyone.

**What types of damages can a plaintiff-patient recover in a medical malpractice action?**

Both compensatory and punitive damages may be awarded in a medical malpractice case. Compensatory damages are actual damages incurred by the plaintiff-patient, which can include, but is not limited to, lost wages, medical expenses (regardless of whether they are paid for by insurance), impairment of earning capacity, pain and suffering, emotional distress and other related provable damages. Pain and suffering includes compensation for any pain, discomfort, fears, anxiety, and other mental and emotional distress suffered by the patient as a result of the health care professional’s conduct. The purpose of compensatory damages is not to punish a defendant or to reward a plaintiff, but rather to compensate the plaintiff for the injuries he or she has suffered.

Punitive damages, on the other hand, are not awarded to compensate for any injury, but are awarded to punish a party for morally culpable conduct and to deter that party and others from acting in the same way in the future. To recover punitive damages, two elements must be proven: (1) wrongful conduct that is outrageously reprehensible; and (2) malice. With respect to the first element, Vermont law limits the recovery of punitive damages to only those cases where the evidence shows that defendant’s wrongdoing has the character of outrage frequently associated with crime. The second element requires evidence of malice which has been defined as bad motive, ill will, personal spite or hatred, and reckless disregard. It is not enough for the plaintiff to show that the defendant’s acts were wrongful or unlawful; there must be proof of the

Negligence resulting from mere inadvertence, incompetence, unskillfulness, or a failure to take precautions is not enough to establish liability for punitive damages because it lacks the element of malice. *Id.* at ¶ 24. Thus, medical malpractice alone is insufficient to establish liability for punitive damages, since medical malpractice is nothing more than professional negligence in the provision of health care.

**What is joint and several liability?**
Joint and several liability refers to the situation in which two or more individuals may be liable for the same harm. For example, if a patient sues three (3) doctors and the jury finds that each doctor was negligent and awards damages, the patient can choose to collect the entire verdict from any one of the three doctors. In Vermont, if the patient collects the entire award from Doctor A, Doctor A cannot seek reimbursement or contribution from Doctors B and/or C.

**What is vicarious liability?**
Generally, vicarious liability or respondeat superior refers to the imposition to liability on one person for the legally actionable conduct of another person based on the relationship between the two persons. A common relationship to which vicarious liability applies is that of employer and employee; employers are legally responsible for the acts of their employees. *In re Desautels Real Estate, Inc.*, 142 Vt. 326, 337 (1982). Thus, if a physician employs a nurse and that nurse commits malpractice, the physician can be found liable for that malpractice based on his or her status as the nurse’s employer.

**Are there “caps” on damages in Vermont?**
There are no “caps” on compensatory or punitive damages in Vermont in medical malpractice cases.

**STATUTES OF LIMITATION**

**What is a statute of limitation?**
A statute of limitation sets forth the maximum time period in which a lawsuit may be brought. Once that time period expires, no suit may be filed, regardless of the validity of the claims asserted.

**What is the statute of limitation on claims of medical malpractice in Vermont?**
Generally, actions for medical malpractice must be brought either within three years of the date of the incident or two years from the date of discovery of (a) the injury, and (b) the fact that it may have been caused by the negligence of a health care provider. 12 V.S.A. § 521; *Lillicrap v. Martin*, 156 Vt. 165, 175-76 (1990). In any case, however, a medical malpractice action may not be filed more than seven years from the date of the incident unless:

- fraudulent concealment has prevented the patient’s discovery of the medical malpractice; or
the action is based upon the discovery of a foreign object in the patient’s body, in which case, the action may be commenced within two years of the date of the discovery of the foreign object. 12 V.S.A. § 521.

Vermont law sets forth a separate statute of limitations for ionizing radiation injuries and injuries from other noxious agents (i.e., a substance originating outside of the body that acts upon the body when exposed to the substance) which are medically recognized as having a prolonged latent development. 12 V.S.A. § 518; Campbell v. Stafford, 2011 VT 11, ¶ 14-15. An action to recover for these types of injuries must be commenced within three years after the person suffering the injury knew or ought reasonably to have known that an injury has been suffered, but in no event can the action be commenced more than twenty years after the date of the last occurrence to which the injury is attributed.

What is the statute of limitations on a wrongful death claim?
Actions for wrongful death must be commenced within two years from the discovery of the death of the person. 14 V.S.A. § 1492(a).

What is the statute of limitations on a survival action?
A survival action is a cause of action that may be prosecuted even though one of the parties has since died. That cause of action is said to “survive” and may be prosecuted by or against the executors or administrators of the estate of the deceased party. 14 V.S.A. § 1452.

The types of actions that will survive are set forth by statute and include an action for the recovery of damages for bodily hurt or injury, such as medical malpractice. 14 V.S.A. § 1452. For example, if a patient dies sometime after suffering an injury believed to be caused by malpractice, but before filing a lawsuit, his or her executor or administrator may file a lawsuit within the time period outlined below. Similarly, if a physician dies, a legal action against the physician or his estate may be filed or continued after his or her death.

To pursue a survival action, the party must have died before the expiration of the applicable statute of limitation (such as the medical malpractice statute of limitation) or within 30 days of the expiration of that statute of limitation. 12 V.S.A. § 557(a). The cause of action must then be commenced by or against the executor or administrator of the decedent within two years of the issuance of the “letters testamentary” or “letter of administration” by the Probate Court. 12 V.S.A. § 557(a).

What is the statute of limitation on other types of claims?
In Vermont, except as provided otherwise, most civil actions, including contract actions, must be commenced within six years after the cause of action “accrues”; that is, after the injured party discovers or reasonably should discover the injury and its cause. 12 V.S.A. § 511.

There is a specific statute of limitation for claims of assault and battery, false imprisonment, slander and libel, personal injuries, and damage to personal property. These actions must be brought within three years from the date of accrual of the cause of action. 12 V.S.A. § 512. As
with the general statute of limitations, a cause of action for these claims accrues when the injured party discovers or reasonably should discover the injury and its cause.

*What does it mean when a statute of limitations is tolled?*

When a statute of limitations is “toll ed,” it means that the time in which a person has to file a suit is temporarily paused or delayed. In Vermont, statutes of limitation are tolled on a variety of grounds, the most common of which are when the person entitled to bring an action is a minor, lacks capacity to protect his or her interests due to a mental condition or psychiatric disability, or is imprisoned at the time the cause of action accrues. The statute of limitation will begin to run only after the “disability is removed”; that is, when the minor turns 18, the person lacking capacity regains capacity, or the imprisoned person is released from prison. 12 V.S.A. § 551(a).

If a person entitled to bring an action becomes unable to protect his or her interests due to a mental condition or psychiatric disability after the cause of action accrues but before the statute has run, the time during which the person cannot protect his or her interests due to a mental condition or psychiatric disability shall not be included as a part of the time allowed for commencement of the action. 12 V.S.A. § 551(b).

**LIABILITY WITH RESPECT TO INFORMED CONSENT**

For specific information regarding consent to treatment, see the section “Consent to Treatment” in the chapter, Consent, Privacy and Medical Records. This section addresses civil liability for failure to obtain informed consent for treatment.

*When may a physician be held liable for failure to obtain informed consent?*

In Vermont, a health care practitioner may be held liable for medical malpractice when he or she fails to obtain informed consent for treatment. Lack of informed consent is defined as:

- The failure to disclose to the patient treatment alternatives and the reasonably foreseeable risks and benefits involved as a reasonable medical practitioner would have disclosed under similar circumstances, in a manner permitting the patient to make a knowledgeable evaluation; or
- The failure to provide a patient with a reasonable answer to any specific question about foreseeable risks and benefits, or the withholding of requested information.

12 V.S.A. § 1909(a).

With respect to the first category, the patient asserting a claim of failure to obtain informed consent must produce expert medical testimony to establish that he or she was not properly informed of the treatment alternatives and the reasonably foreseeable risks and benefits involved therewith, as a reasonable medical professional under similar circumstances would have disclosed. Without that expert testimony, the plaintiff cannot meet his or her burden of proof.

In assessing whether a physician has obtained informed consent, the focus is on whether the physician advised the patient of the treatment alternatives and reasonably foreseeable risks and benefits which were commonly known to the profession at the time the informed consent discussion did or could have occurred, as well as at the time the treatment was given. In other
words, the scope of the information that should be provided in obtaining informed consent is based on the information commonly known to the profession at that time.

Although there is no specific requirement that informed consent be acknowledged in writing, obtaining such written acknowledgement is common and prudent. In any case, the practitioner should document the content of the discussion and the patient’s consent in the medical record.

Under what circumstances, if any, does a physician not have to obtain informed consent?
Civil liability for medical malpractice based on a lack of informed consent does not apply when the provision of medical care and treatment occurred during an emergency. 12 V.S.A. § 1909(b). What constitutes an emergency is not defined in Vermont’s informed consent statute and has not been specifically addressed by the Vermont Supreme Court. In Small v. Gifford Memorial Hospital, 133 Vt. 552, 349 A.2d 703 (1975), which was decided before the informed consent statute was enacted, the Vermont Supreme Court cited with approval to a case in which the emergency exception was described as encompassing a situation where the “patient is unconscious or otherwise incapable of consenting and the harm from non-treatment outweighs the harm threatened by the treatment.”

Whether a particular situation constitutes a medical emergency such that the requirement of informed consent does not apply would be measured by an objective medical standard. In an emergency situation in which the patient is unable to provide consent, if the patient’s spouse, significant other or other close family member is available, then the better practice if time permits would be to seek the consent of the family, although doing so is not required by the informed consent statute. In all circumstances, the physician should document in the patient’s chart the factors which created the emergency situation and which informed his or her decision to render the treatment provided.

What defenses can be asserted in a malpractice action alleging the failure to obtain informed consent?
The informed consent statute in Vermont sets forth four defenses that may be asserted in a suit alleging failure to obtain informed consent. 12 V.S.A. § 1909(c)(1)-(4). They are:

- The risk not disclosed is too commonly known to require disclosure and is not substantial;
- The patient assured the medical practitioner he or she would undergo the treatment, procedure or diagnosis regardless of the risk involved, or the patient indicated to the medical practitioner that he or she did not want to be informed of the matters to which he or she would be entitled to be informed;
- Consent by or on behalf of the patient was not reasonably possible; or
- A reasonably prudent person in the patient’s position would have undergone the treatment or diagnosis if he or she had been fully informed.

A plaintiff cannot establish liability by simply alleging that he or she would not have undergone the treatment received if he or she had been fully informed. Rather, the statute creates an objective standard focused on whether a reasonably prudent person would give his or her consent to treatment if he or she had been fully informed. If so, then there is no liability for failure to obtain informed consent.
LIABILITY WITH RESPECT TO ADVANCE DIRECTIVES FOR HEALTH CARE, DNR ORDERS, AND COLST ORDERS

“The State of Vermont recognizes the fundamental right of an adult to determine the extent of health care the individual will receive, including treatment provided during periods of incapacity and at the end of life.” 18 V.S.A. § 9700. To that end, Vermont allows adults to retain control over their own health care through the use of advance directives.

What is an advance directive?
An advance directive is a written document that may include an appointment of an agent (an adult with capacity to whom authority to make health care decisions for a principal/patient is delegated under an advanced directive), identification of a preferred primary care clinician, instructions on health care desires or treatment goals, an anatomical gift, disposition of remains, and funeral goods and services. It includes documents designated under prior law as a durable power of attorney for health care or a terminal care document. 18 V.S.A. § 9701(1).

In Vermont, health care providers, health care facilities, and residential care facilities shall not provide health care to a patient without capacity, except on an emergency basis, without first attempting to determine whether the patient has an advance directive in effect. 18 V.S.A. § 9707(a).

For specific information regarding advance directives, see the chapter on End of Life Issues (forthcoming).

What is a DNR Order?
A DNR (do-not-resuscitate) order is a written order of the patient’s clinician directing health care providers not to attempt resuscitation. 18 V.S.A. § 9701(8). DNR identification is a necklace, bracelet, or anklet identifying the patient as an individual who has a DNR order. 18 V.S.A. § 9701(9).

For specific information regarding DNR orders, see the chapter on End of Life Issues (forthcoming).

What is a COLST Order?
A COLST (clinician order for life-sustaining treatment) order is a clinician’s order for treatment such as intubation, mechanical ventilation, transfer to a hospital, antibiotics, artificially administered nutrition, or another medical intervention. COLST orders are designed for use in outpatient settings and health care facilities and may include a DNR order. 18 V.S.A. § 9701(6).

For specific information regarding COLST orders, see the chapter on End of Life Issues (forthcoming).
Is a health care professional who complies with the terms of an advance directive, DNR Order, or COLST Order exposed to civil or criminal liability for doing so?

No. If health care professionals, health care facilities, residential care facilities, and their agents comply with the provisions of Chapter 231 of Title 18 governing advance directives for health care, then they are immune from civil and criminal liability when they:

- provide or withhold treatment or services in good faith pursuant to the direction of a principal/patient, the provisions of an advance directive, a DNR order, a COLST order, a DNR identification, the consent of a principal/patient with capacity or the principal/patient’s agent or guardian, or a decision or objection of a principal/patient; or
- rely in good faith on a suspended or revoked advance directive, a DNR order, or a COLST order, unless the provider or facility should have known of the suspension or revocation.

18 V.S.A. § 9713(b)(1). Health care professionals and facilities are not, however, immune from liability for the failure to follow the standards of professional conduct and to exercise due care in the provision of services. 18 V.S.A. § 9713(b)(3).

Additionally, no employee of the previously listed professionals and facilities can be subjected to an adverse employment decision or evaluation for:

- complying with an advance directive, a DNR order, a DNR identification, or a COLST order;
- relying on an amended, suspended or revoked advance directive unless the employee knew or should have known of the amendment, suspension or revocation; or
- for providing notice to his or her employer that he or she is unwilling to comply with an instruction in an advance directive due to a moral or other conflict, so long as the employee has provided ongoing healthcare until a new employee has been found to provide the services.

18 V.S.A. § 9713(c).

When may a health care provider be exposed to civil or criminal liability?

Health care providers, health care facilities, residential care facilities, and their agents having actual knowledge of an advance directive or an instruction of the principal, agent or guardian, are subject to review and disciplinary action by the appropriate licensing entity, and/or civil or criminal liability for failing to comply with the terms of a known advance directive or failing to follow the instructions of a duly appointed agent or guardian who has the authority to make health care decisions for a principal/patient. 18 V.S.A. § 9714.

There are, however, circumstances when health care providers may properly refuse to comply with the terms of an advance directive or the instructions of a duly appointed agent or guardian. See 18 V.S.A. § 9707 and chapter on End of Life Issues. In those circumstances, the professional must still comply with the procedures set forth in 18 V.S.A. § 9707 in order to be protected from civil and/or criminal liability or disciplinary action.

**LIABILITY WITH RESPECT TO THE DUTY TO PROTECT THE ENDANGERED ACT**
What is the Duty to Aid the Endangered Act?

The purpose of the Duty to Aid the Endangered Act, 12 V.S.A. § 519, is to encourage rescuers to assist others in danger by penalizing them for not acting, while at the same time shielding them from civil liability for acts of ordinary negligence committed during the rescue. *Hardingham v. United Counseling Serv.*, 164 Vt. 478, 483 (1995).

The Vermont Legislature enacted this statute largely due to its concern that medical personnel were reluctant to help those in need for fear of malpractice suits. The statute imposes an affirmative duty on everyone, including health care professionals, to provide reasonable assistance to individuals who are known to be exposed to “grave physical harm.” Grave physical harm is not limited to a single, traumatic event such as a car accident, but rather encompasses all situations in which a rescuer knows that someone is exposed to serious harm.

A person who provides reasonable assistance to an individual exposed to grave physical harm shall not be held liable for civil damages arising out of his or her conduct unless his or her acts constitute gross negligence, or he or she will receive or expects to receive remuneration for his or her services. 12 V.S.A. § 519(b). Gross negligence is “more than an error of judgment, momentary inattention, or loss of presence of mind, rather, it amounts to a failure to exercise even a slight degree of care and an indifference to the duty owed to another.” *Hardingham v. United Counseling Serv.*, 164 Vt. 478, 482 (1995).

With respect to the receipt of remuneration, the mere fact that a rescuer is paid a regular salary during the time period in which he or she comes to the assistance of an individual does not remove the immunity created by the statute. Rather, it is when the rescuer charges the victim for the services rendered that the statutory immunity becomes unavailable. Nothing in this statute alters the liability of a health care provider for acts committed in the ordinary course of his or her practice.

**ALTERNATIVE DISPUTE RESOLUTION**

Does Vermont law require screening or arbitration of medical malpractice claims prior to the commencement of a lawsuit?

No. In 1991, Vermont adopted a system for mandatory arbitration of medical malpractice claims; however, implementation of that system was tied to the enactment of a “universal access health care system” by the General Assembly. To date, that has not happened. Instead, the Legislature enacted a procedure for voluntarily submitting medical malpractice claims to arbitration, which has seldom been used. Both the physician and patient must agree to submit a claim to arbitration after discovery of the alleged injury and prior to the commencement of any trial on that claim. 12 V.S.A. §§ 7001-7009.

The arbitration panel consists of a judicial referee selected by the court administrator, and a layperson and member of the same profession as the respondent-doctor, both chosen by lot. In addition to challenges for cause, the parties have the right to one preemptory challenge with respect to the judicial referee and three such challenges with respect to the lay and professional panel members. The law sets forth the specific procedures to be followed in the arbitration process. Claims are to be submitted to the arbitration panel in an “informal matter;” strict
adherence to the technical rules of procedure and evidence, as in a civil litigation, is not required. Discovery under the Vermont Rules of Civil Procedure is allowed and the parties have the right to present testimony and cross-examine witnesses. Notably, unlike in medical malpractice claims filed in Superior Court, no expert testimony is required during arbitration. The decision of the arbitration panel may be appealed directly to the Vermont Supreme Court.

_Does Vermont law require participation in alternative dispute resolution after a medical malpractice lawsuit is commenced?_

Yes. Parties to medical malpractice actions (and most other civil actions) filed in Vermont, whether in state or federal court, are required to participate in alternative dispute resolution prior to going to trial. V.R.C.P. 16.3.

**NATIONAL PRACTITIONER DATA BANK**

_What is the National Practitioner Data Bank?_

The National Practitioner Data Bank (NPDB) is a federal data bank which was created to serve as a repository of information on medical malpractice payments and other adverse actions related to health care providers in the United States. Federal law determines the types of actions reported to the NPDB, who submits the reports, and who may query the data bank to obtain information on a health care provider. The mission of the NPDB is to improve health care quality, protect the public, and reduce health care fraud and abuse in the United States.

The NPDB’s website, www.npdb.hrsa.gov, has many resources for practitioners regarding the data bank, including this guidebook: [https://www.npdb.hrsa.gov/resources/NPDBGuidebook.pdf](https://www.npdb.hrsa.gov/resources/NPDBGuidebook.pdf).

_What information must be reported to the National Practitioner Data Bank and by whom?_

There are several categories of information that need to be reported to the NPDB and a variety of sanctions, including civil money penalties, which may be imposed for failure to report required information. Below are some of the more common categories of information that must be reported to the NPDB:

1. **Medical Malpractice Payments**
   Any entity, including an insurance company, that makes a payment for the benefit of a health care practitioner in settlement of a written claim or judgment for medical malpractice against that practitioner must report the payment information to the NPDB and the appropriate state licensing board. The report must be made within 30 days of the date the first payment is made.

   Payments made as a result of a suit or claim asserted solely against an entity (such as a hospital, clinic, or practice group), and not against an identified individual practitioner, are not reportable. Additionally, payments made by individual practitioners for their own benefit do not need to be reported to the NPDB. A business corporation or other business entity comprised of a sole practitioner that makes a payment for the benefit of a named practitioner, however, must report that payment to the NPDB.

2. **Adverse State Licensure and Certification Actions**

State licensing and certification authorities must report all adverse licensure actions (not just those based on professional competence and conduct) taken against all healthcare practitioners, as well as those actions taken against healthcare entities. Such licensure actions include, but are not limited to revocation, suspension, censure, reprimand, probation and surrender of a license or certification; and the dismissal or closure of an action because the health care practitioner or entity surrendered the license or certification, or because the subject of the proceeding left the state or jurisdiction. Reports must be made to the NPDB within 30 days from the date of the action.

(3) Adverse Clinical Privileging and Medical Staff Membership Actions
Hospitals and other eligible health care entities must report to the NPDB and the appropriate state licensing board any professional review actions that adversely affect a physician’s or dentist’s clinical privileges or medical staff membership for a period of more than 30 days. Additionally, they must also report a physician’s or dentist’s surrender or the restriction of clinical privileges or medical staff membership while under investigation for possible professional incompetence or improper professional conduct, or in return for not conducting an investigation or professional review action. The report must be made within 30 days from the date adverse action was taken.

In addition, hospitals and other health care entities may (and are encouraged to) report adverse actions taken against the clinical privileges or medical staff membership of licensed health care practitioners other than physicians and dentists when those actions are based on the practitioner’s professional competence or professional conduct that adversely affects, or could adversely affect, the health or welfare of a patient.

(4) Adverse Professional Society Membership Actions
Professional societies must report to the NPDB and the appropriate state licensing board any professional review action based on professional competence or professional conduct that adversely affects or may adversely affect a physician or dentist’s membership. The report must be made within 30 days from the date adverse action was taken.

Professional societies may report similar adverse actions taken against the membership of health care practitioners other than physicians and dentists.

(5) Exclusions from Medicare/Medicaid
Federal agencies, state law enforcement agencies, state Medicaid fraud control units, and state agencies administering or supervising the administration of a state health care program must report health care practitioners who have been excluded from participating in federal or state health care programs. Exclusion from such a program means a temporary or permanent disqualification of an individual or entity from participation in the program, such that the individual or entity will not be reimbursed under any federal or state health-related program for any services that are provided.
Are health care practitioners notified when reports concerning them are made to the NPDB?

Yes, the subject of a report made to the NPDB is notified of the report by the NPDB. The notification will include instructions for obtaining an official copy of the report through the Report Response Service on the NPDB website.

What recourse do health care practitioners have when they dispute some or all of the information contained in a report to the NPDB?

The NPDB is prohibited by law from modifying the information in the reports. If the information in a report is inaccurate, the subject of that report has several options. First, he or she can add a statement (“Subject Statement”) to a report at any time. The statement will be appended to the report and sent with the report when queries are made. There are specific requirements that must be complied with when submitting a Subject Statement; they can be found on the NPDB’s website.

Second, the subject of the report can contact the reporting entity to request that it voluntarily correct the information by filing a correction to the report.

Third, at any time, the subject of a report may dispute the report and enter the report into “Dispute Status” to disagree with either the factual accuracy of the report or whether the report was properly submitted under the NPDB’s reporting requirements. After a report has been entered into Dispute Status, the subject of the report may do any of the following:

- Leave the report in Dispute Status and the NPDB will take no further action;
- Withdraw the report from Dispute Status; or
- Request that the report be elevated to “Dispute Resolution.”

The NPDB will not review a report entered into Dispute Status; such a review will only occur after the report has been elevated to Dispute Resolution. The subject of the report must meet two prerequisites in order to have a report elevated to Dispute Resolution:

- During the 60 days after a report has been entered into Dispute Status, the subject of the report must contact the reporting entity in an attempt to resolve the issues raised by the report; and
- The subject of the report must submit to the NPDB proof of his/her attempt to resolve the issues with the reporting entity (such as correspondence to the reporting entity outlining the issues with the report and the entity’s response to that correspondence).

The scope of a review once a report is elevated to Dispute Resolution is very narrow. Specifically, the subject of the report may only dispute (1) whether the report was submitted in compliance with the NPDB reporting requirements, and (2) the factual accuracy of the information reported. Thus, the subject cannot seek review of issues such as the merits of a medical malpractice claim or whether the reporting entities complied with due process in connection with its internal processes.

The NPDB website and NPDB Guidebook explain the specific procedures which must be followed when a report is elevated to Dispute Resolution and reviewed by the NPDB, and the relief potentially available to the subject following such a review.
Who can access the information maintained by the NPDB?

Information reported to the NPDB is considered confidential and shall not be disclosed except as specifically provided in the NPDB regulations. Under those regulations, certain information in the NPDB may, and in some cases shall, be requested by a number of entities and individuals including, but not limited to, the following:

1. **Hospitals**
   Hospitals are the only health care entities with mandatory requirements for querying the NPDB. Failure to do so will subject the hospital to sanctions. Hospitals must query the NPDB:
   - When a physician, dentist, or other licensed health care practitioner applies for a position on its medical staff (courtesy or otherwise) or for clinical privileges;
   - Every two years on every physician, dentist, or other licensed health care practitioner who is a member of the medical staff or has clinical privileges;
   - When a physician, dentist, or other licensed health care practitioner wishes to add to or expand existing privileges; and
   - Each time a physician, dentist, or other licensed health care practitioner submits an application for temporary privileges.
   Hospitals may query at other times as they deem necessary.

2. **Physicians, dentists and other (licensed) health care practitioners**
   All health care practitioners can self-query the NPDB regarding himself or herself at any time. This can be done through the NPDB website.

3. **State licensing and certification agencies**
   These agencies may query the NPDB on physicians, dentists and other licensed health care practitioners when they are:
   - determining the fitness of individuals to provide health care services,
   - protecting the health and safety of individuals receiving health care through programs that they administer, or
   - protecting the fiscal integrity of the programs that they administer.

4. **Plaintiffs’ Attorneys**
   Plaintiffs’ attorneys or plaintiffs acting on their own behalf may query the NPDB concerning a practitioner only if they have:
   - filed a medical malpractice claim in state or federal court or other adjudicative body against a hospital,
   - the practitioner whose information is requested is named in that action, and
   - the requester must be able to demonstrate that the hospital failed to make a required mandatory query with evidence not obtained from the NPDB.

   Plaintiffs and their attorneys may not query the NPDB for information to be used in suits against practitioners. Defense attorneys are not allowed to query the NPDB, even though the defendant-practitioner is allowed to self-query.

5. **Other health care entities and professional societies**
Health care entities, other than hospitals, that provide health care services and follow a formal peer review process for the purpose of furthering health care, may query when entering an employment or affiliation relationship with a health care practitioner, when health care practitioners apply for clinical privileges or medical staff appointments, or when they are engaging in professional review activities.

(6) Professional societies
Professional societies may query when entering an employment or affiliation (membership) relationship with a health care practitioner or in conjunction with professional review activities.

SAFE APOLOGY LAW

What is the safe apology law?
In 2006, the Vermont Legislature enacted the so-called safe apology law, which provides that an oral expression of regret or apology, including any oral good faith explanation of how a medical error occurred, made by or on behalf of a health care provider or health care facility (1) does not constitute a legal admission of liability for any purpose, and (2) is inadmissible in any civil or administrative proceeding against the provider or facility. 12 V.S.A. § 1912(a). The statute only applies to medical errors that occurred on or after July 1, 2006.

The person making the apology may not be questioned at deposition or otherwise with respect to the apology. The apology must be made within 30 days of when the provider or facility knew or should have known of the consequences of the error in order to come within the protections of this law. Failure to comply with the requirements of the statute will result in a waiver of the protections it offers.

PATIENT SAFETY SURVEILLANCE AND IMPROVEMENT SYSTEM

What is the Patient Safety Surveillance and Improvement System (PSSIS)?
The Patient Safety Surveillance and Improvement System (PSSIS) was created in 2006 for the purpose of improving patient safety, eliminating adverse events in Vermont hospitals, and supporting and facilitating quality improvement efforts by hospitals. 18 V.S.A. §§ 1912-1919.

Under the PSSIS, hospitals are required to do, among other things, the following:

- Develop, maintain and implement internal policies and procedures to:
  - Identify, track, and analyze reportable adverse events, adverse events, and near misses;
  - Determine what type of causal analysis, if any, is appropriate;
  - Conduct causal analyses and develop corrective action plans; and
  - Disclose to patients, or; in the case of a patient death, an adult member of the immediate family, at a minimum, adverse events that cause death or serious bodily injury;
- Report reportable adverse events to the Department of Health, including providing the department with copies of the hospital’s causal analysis and corrective action plan in connection with each reportable adverse event;
For reportable adverse events that must also by law be reported to other departments or agencies, notify the Department of Health or provide a copy of any written report and provide any causal analysis information required by the department; and

Provide the commissioner and his/her designees reasonable access to (1) confidential patient health information under 12 V.S.A. § 1612(a), and (2) the minutes and records of any peer review committee and any other information subject to peer review protection under 26 V.S.A. § 1443, for the purpose of evaluating compliance with this law.

18 V.S.A. § 1915.

All of the information made available to the Department of Health as part of the PSSIS is confidential and privileged, and exempt from the public access to records law. 18 V.S.A. § 1917(a). In any civil or administrative proceeding against a health care provider arising out of the matters which were evaluated and reviewed by the department:

- The information made available to the Department of Health as part of the PSSIS is immune from subpoena, not subject to discovery and is not admissible into evidence; and
- No person with access to information made available to the commissioner or his/her designees shall be permitted or required to testify as to any findings, recommendations, evaluations, opinions, or other actions of the department in any civil or administrative proceeding against a health care provider arising out of the matters which were evaluated and reviewed by the department.

18 V.S.A. § 1917(a), (b).

Hospitals are permitted to replace health care provider identifying information in peer review materials with a surrogate identifier that allows for tracking of adverse events involving the same provider without disclosing the provider’s identity. 18 V.S.A. § 1917(e).

Hospitals that fail to comply with any of the requirements of the PSSIS are subject to monetary penalties. 18 V.S.A. § 1918.

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REGULATION OF PHYSICIANS

Topics Covered in this Chapter:
Overview
Licensing
Standard of Conduct
Complaint Process
Discipline
Appellate Avenues
Public Access to Information
About the Author

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Note: This information is provided to assist interested persons in becoming familiar with the law pertaining to the regulation of physicians in Vermont. It is not an official interpretation or statement of policy by the Department of Health or the Vermont Board of Medical Practice and does not constitute legal advice.

OVERVIEW

What state agency is responsible for licensing and disciplining physicians, podiatrists, physician assistants, and anesthesiologist assistants in Vermont?
The Vermont Board of Medical Practice (“Board”) licenses physicians and podiatrists, and certifies physician assistants, radiologist assistants, and anesthesiologist assistants. The Board also investigates complaints and issues findings and actions regarding unprofessional conduct. The Board’s stated mission is to evaluate the fitness of professionals to practice in Vermont, and to take action where needed to protect the public health and safety.

The Board consists of 17 members: nine licensed physicians, one certified physician assistant, one licensed podiatrist, and six public members. Members are appointed by the governor, with the advice and consent of the Senate. They serve for up to two consecutive five-year terms. See http://healthvermont.gov/health-professionals-systems/board-medical-practice/board-members.

The Board operates under administrative rules adopted under the Vermont Administrative Procedure Act. As a result, these rules carry the same force of law as a statute. The Board, however, last adopted rules in 2001. Since 2002, it has advised the public that it was in the process of revising its rules in light of legislative changes to governing statutes, but it has yet to do so. As a result, to the extent any rule conflicts with a current statute, the statute controls. See http://healthvermont.gov/health-professionals-systems/board-medical-practice/statutes-laws-rules-and-policies.
What state agency is responsible for licensing and disciplining doctors of osteopathy, advance practice registered nurses, and other health care professionals in Vermont?

The Board of Osteopathic Physicians and Surgeons is a five-member board that regulates osteopathic physicians through the Vermont Office of Professional Regulation, a branch of the Secretary of State’s office. The Board of Nursing is an eleven-member board that regulates nurses through the Vermont Office of Professional Regulation. Other certified and registered health care professionals are similarly regulated by other boards within the Office of Professional Regulation. These boards function similarly to the Board, but each generally operates under different laws and has adopted its own administrative rules. A list of professions that provide links to each board and its rules can be found at: https://www.sec.state.vt.us/professional-regulation/list-of-professions.aspx.

Note: the discussion of licensing and discipline below applies to the physicians and other health care professionals licensed by the Board of Medical Practice.

LICENSING

How do I obtain a Vermont Medical License from the Vermont Board of Medical Practice?
In order to be granted a license to practice medicine the applicant must present evidence satisfactory to the Board that the applicant:

- Is at least 18 years of age;
- Is competent in speaking, writing and reading the English language;
- Has completed high school and two years of college or the equivalent;
- Is a graduate of a Board-approved medical school, or a medical school accredited by the LCME or CACMS;
- Has met the Board’s criteria for postgraduate training;
- Has met the Board’s criteria for license by examination, see Rules of the Vermont Board of Medical Practice Rules (“Board Rules”) 2.3; license by reciprocity, see Board Rules 2.4; or license by appointment to the faculty of a Vermont medical college, see Board Rules 2.5;
- Has presented reference forms as to moral character and professional competence; and
- Has been interviewed by a Board member, the licensing committee, and/or the Board. Board Rules 2.2; see also, 26 V.S.A. §§ 1391, 1393, 1395, 1396.

The Board now uses the Uniform Application for Physician State Licensure, portions of which must be completed online. All application material can be found at: http://healthvermont.gov/health-professionals-systems/board-medical-practice/application-licensing-and-fees. It is the applicant’s personal responsibility to ensure complete and accurate responses to all application questions.

What does the renewal process consist of?
Licenses are renewed on a fixed biennial schedule. It is the responsibility of the licensee to renew his or her license before it lapses. The date on which a license shall lapse is printed on the license. The Board is required by statute to send each licensee a notice and renewal application
form to the address last provided to the Board at least a month prior to the physician’s license expiration date. Board Rules 3.1; 26 V.S.A. § 1400.

The licensee must ensure complete and accurate responses to all renewal application questions. If a physician does not return the completed renewal application and fee to the Board by the date on which the license lapses, the physician’s license will automatically lapse.

Licensees have a continuing obligation during each two-year renewal period to promptly notify the Board of any change or new information regarding disciplinary or other action limiting or conditioning their license or ability to practice in any licensing jurisdiction. Failure to do so may subject the licensee to disciplinary action by the Board. Board Rules 3.1.

If a license has not been renewed by the required date, it has lapsed. Therefore, a licensee may not legally practice in Vermont after a license has lapsed. The licensee must halt the practice of medicine until the license has been reinstated. Board Rules 3.2.

What are the continuing medical education requirements?
In response to a legislative change in 2011, the Board has adopted Continuing Medical Education (“CME”) Rules. These can be found at: http://healthvermont.gov/health-professionals-systems/board-medical-practice/statutes-laws-rules-and-policies.

These rules require a physician seeking to renew his or her license to certify that he or she has completed at least thirty (30) hours of qualifying CME during the most recent two-year licensing period. Continuing Medical Education Rules 22.2(a).

There are some exceptions to the 30-hour requirement for new licensees and active or deployed members of the military. Continuing Medical Education Rules 22.2(b) & (h). For physicians licensed in Vermont for the first time during the most recent two-year licensing period, if licensed in Vermont for less than one year, there is no requirement for CME at the time of the first renewal. If licensed for one year or more during that initial period of Vermont licensure, the licensee shall complete at least 15 hours of approved CME activity and those 15 hours shall include any subject-specific CME required by Board rules. Continuing Medical Education Rules 22.2(b).

Licensees who are members of the armed forces and who are subject to a mobilization and/or deployment for all or part of a licensing cycle will be treated the same as licensees who are licensed for the first time during a licensing cycle. Therefore, a licensee whose military mobilization/deployment covers a year or more is not required to complete CME for that cycle. A licensee whose military duties during the two-year cycle total less than one year must meet the CME requirement of at least 15 hours, including any required subjects. Continuing Medical Education Rules 22.2(h).

A CME activity qualifies only if it has been approved by the American Medical Association Physician’s Recognition Category 1 Credit. Id. at 22.3. The licensee need not file proof of completion of the CME requirement, but he or she should maintain such proof for at least four
years after submission of compliance certification, as the Board may audit such records during this time. Id. at 22.2(a).

CME Rules also require that CME hours assure that the licensee has updated his or her knowledge and skills in his or her own specialties and also has kept abreast of advances in other fields for which patient referrals may be appropriate. Id. at 22.2(d). The Board broadly interprets a licensee’s “own area of practice,” and it acknowledges that training in many other fields may be reasonably related to a licensee’s own specialties. Id.

The CME Rules also require a licensee to complete CME hours on other subjects. This includes at least one CME hour on hospice, palliative, and pain management. Id. at 22.2(e). All licensee who prescribe controlled substances must also complete one CME hour on prescribing controlled substances. Id. at 22.2(f). Any licensee registered with the U.S. Drug Enforcement Agency (“DEA”), holding a DEA number to prescribe controlled substances, or has submitted a pending application for one is presumed to prescribe controlled substances. Id. Due to the passage of Act 173, beginning with license renewals due Nov. 2018, all physicians with a DEA number, pending application for a DEA number, or who dispense controlled substances will be required to show they have completed a total of at least 2 hours over the course of the previous 2-year licensing cycle - rather than the current one hour - of continuing education on the topic of prescribing controlled substances.

If a licensee fails to complete the CME requirements by the time of his or her license renewal, the renewal application must be accompanied by an acceptable make-up plan signed by the licensee, or the Board will reject the application. Id. at 22.4(a). An acceptable plan must be signed and include a timeline for completing the requirements within 120 days after his or her license expiration date, an indication of his or her good faith intent to complete the requirements as indicated, and identify the activities the licensee will attend, although the licensee may later substitute activities. Id. at 22.5(c). Any licensee not in good standing with respect to CME requirements is subject to investigation by the Board for unprofessional conduct. Id. at 22.5(d).

**How do I reinstate my lapsed license?**

To seek reinstatement after failing to renew within a year of the expiration of the license, a physician must complete a renewal application and tender it to the Board with a late fee, in addition to the fee required for renewal. The Board may stay the decision on the application pending investigation of charges or allegations of unprofessional conduct against the renewal applicant. The Board may seek or request such additional information as it deems needed to make a determination as to the renewal application. The Board may deny the renewal of a license on grounds of unprofessional conduct as set forth under Vermont law after notice and opportunity to be heard has been provided to the physician. Board Rules 3.3.

If a license is lapsed for one year or more the physician must complete a reinstatement application in full and pay the associated application fee. The reinstatement application requires additional information beyond that required in the standard renewal application. A chronological accounting of the physician’s professional activities in other jurisdictions during the period the license was lapsed in Vermont must be presented. The physician must include:

- A letter from the chief of staff of each hospital at which he or she held privileges during the period in which the Vermont license was lapsed; and
• A license verification from each state in which he or she held an active license during the period in which the Vermont license was lapsed.

In addition, he or she must appear for a personal interview. Reinstatement may be denied on grounds of unprofessional conduct as set forth under Vermont law or for other good cause, after notice and opportunity to be heard has been provided to the physician. Board Rules 3.4. The reinstatement application can be found here: http://healthvermont.gov/health-professionals-systems/board-medical-practice/application-licensing-and-fees.

How are physician assistants certified?
Physician assistants receive a certification that authorizes them to practice only within the employment contract and scope of practice submitted and approved by the Board. Section II of the Board Rules provides the rules governing physician assistants, including their certification. Physician assistants must file the necessary documents and obtain Board approval in advance of practicing, when a change of employer occurs, to change supervising physician(s), to add new practice sites, or to otherwise make any changes to their scope of practice as approved by the Board. Physician assistant certification and authority to practice terminates immediately upon dissolution of the employment contract that was approved for a particular certification and does not resume unless and until a new certification is issued by the Board. Board Rules Sec. II.

STANDARD OF CONDUCT

What actions constitute unprofessional conduct?
Per statute, 26 V.S.A. § 1354, the following actions constitute unprofessional conduct:
• Fraud or misrepresentation in applying for or procuring a medical license or in connection with applying for or procuring periodic renewal of a medical license;
• All advertising of medical business which is intended or has a tendency to deceive the public or impose upon credulous or ignorant persons and so be harmful or injurious to public morals or safety;
• Abandonment of a patient;
• Habitual or excessive use or abuse of drugs, alcohol, or other substances that impair the licensee's ability to practice medicine;
• Promotion by a physician of the sale of drugs, devices, appliances, or goods provided for a patient in such a manner as to exploit the patient for financial gain of the physician or selling, prescribing, giving away, or administering drugs for other than legal and legitimate therapeutic purposes;
• Conduct which evidences unfitness to practice medicine;
• Willfully making and filing false reports or records in his or her practice as a physician;
• Willful omission to file or record, or willfully impeding or obstructing a filing or recording, or inducing another person to omit to file or record medical reports required by law;
• Failure to make available promptly to a person using professional health care services, that person’s representative, succeeding health care professionals or institutions, when given proper written request and direction of the person using professional health care services, copies of that person's records in the possession or under the control of the licensed practitioner;
Solicitation of professional patronage by agents or persons or profiting from the acts of those representing themselves to be agents of the licensed physician;
Division of fees or agreeing to split or divide the fees received for professional services for any person for bringing to or referring a patient;
Agreeing with clinical or bio-analytical laboratories to make payments to such laboratories for individual tests or test series for patients, unless the physician discloses on the bills to patients or third party payors the name of such laboratory, the amount or amounts to such laboratory for individual tests or test series and the amount of his or her processing charge or procurement, if any, for each specimen taken;
Willful misrepresentation in treatments;
Practicing medicine with a physician who is not legally practicing within the state, or aiding or abetting such physician in the practice of medicine; except that it shall be legal to practice in an accredited preceptorship or residency training program;
Gross overcharging for professional services on repeated occasions, including filing of false statements for collection of fees for which services are not rendered;
Offering, undertaking, or agreeing to cure or treat disease by a secret method, procedure, treatment, or medicine;
Consistent improper utilization of services;
Consistent use of nonaccepted procedures which have a consistent detrimental effect upon patients;
Professional incompetency resulting from physical or mental impairment;
Permitting one’s name or license to be used by a person, group, or corporation when not actually in charge of, or responsible for, the treatment given;
In the course of practice, gross failure to use and exercise on a particular occasion or the failure to use and exercise on repeated occasions, that degree of care, skill, and proficiency which is commonly exercised by the ordinary skillful, careful, and prudent physician engaged in similar practice under the same or similar conditions, whether or not actual injury to a patient has occurred;
Revocation of a license to practice medicine or surgery in another jurisdiction;
Failure to comply with the provisions of the Vermont Bill of Rights for Hospital Patients;
Failure to comply with an order of the Board or violation of any term or condition of a license which is restricted or conditioned by the Board;
Any physician who, in the course of a collaborative agreement with a nurse practitioner allows the nurse practitioner to perform a medical act which is outside the usual scope of the physician’s own practice or which the nurse practitioner is not qualified to perform by training or experience, or which the ordinary reasonable and prudent physician engaged in a similar practice would not agree should be written into the scope of the nurse practitioner’s practice;
Failure to comply with provisions of federal or state statutes or rules governing the practice of medicine or surgery;
Practice of profession when medically or psychologically unfit to do so;
Delegation of professional responsibilities to a person whom the licensed professional knows, or has reason to know, is not qualified by training, experience, education, or licensing credentials to perform them;
Conviction of a crime related to the practice of the profession or conviction of a felony, whether or not related to the practice of the profession, or failure to report to the board a
conviction of any crime related to the practice of the profession or any felony in any court within 30 days of the conviction;

- Use of the services of an anesthesiologist assistant by an anesthesiologist that is inconsistent with the assistants’ certification;
- Use of the services of a radiologist assistant by a radiologist in a manner that is inconsistent with the assistant’s certification;
- Providing, prescribing, dispensing, or furnishing medical services or prescription medication or prescription-only devices to a person in response to any communication transmitted or received by computer or other electronic means, when the licensee fails to take the following actions to establish and maintain a proper physician-patient relationship:
  - a reasonable effort to verify that the person requesting medication is in fact the patient, and is in fact who the person claims to be;
  - establishment of documented diagnosis through the use of accepted medical practices; and
  - maintenance of a current medical record.
- With respect to this conduct, an electronic, on-line, or telephonic evaluation by questionnaire is inadequate for the initial evaluation of the patient.
- With respect to this conduct, the following would not be in violation if transmitted or received by computer or other electronic means:
  - initial admission orders for newly hospitalized patients;
  - prescribing for a patient of another physician for whom the prescriber has taken the call;
  - prescribing for a patient examined by a licensed advanced practice registered nurse, physician assistant, or other advanced practitioner authorized by law and supported by the physician;
  - continuing medication on a short-term basis for a new patient, prior to the patient's first appointment; or
  - emergency situations where life or health of the patient is in imminent danger;
- Failure to provide to the Board such information it may reasonably request;
- Disruptive behavior which involves interaction with physicians, hospital personnel, office staff, patients, or support persons of the patient or others that interferes with patient care or could reasonably be expected to adversely affect the quality of care rendered to a patient;
- Commission of any sexual misconduct which exploits the physician-patient relationship, including sexual contact with a patient, surrogates, or key third parties;
- Prescribing, selling, administering, distributing, ordering, or dispensing any drug legally classified as a controlled substance for the licensee’s own use or to an immediate family member as defined by rule;
- Signing a blank or undated prescription form;
- Use of the services of a physician assistant by a physician in a manner which is inconsistent with the assistant’s certification, see 26 V.S.A. § 1739a;
- Failure to practice competently by reason of any cause on a single occasion or on multiple occasions; Failure to practice competently includes, as determined by the Board:
  - Performance of unsafe or unacceptable patient care; or
• Failure to conform to the essential standards of acceptable and prevailing practice.

Additionally, the Board may refuse to issue a license to a physician who, by false or fraudulent representation, has obtained or sought to obtain practice in their profession, or by false or fraudulent representation of his or her profession, has obtained or sought to obtain money or any other thing of value, or who assumes names other than his or her own, or for any other immoral, unprofessional, or dishonorable conduct. 26 V.S.A. § 1398. However, a certificate shall not be suspended, revoked, or refused until the holder or applicant is given a hearing before the Board. In the event of revocation, the holder of any certificate so revoked shall forthwith relinquish the same to the secretary of the board. Id.

When does a physician have to report an impaired, incompetent, or unethical colleague?
The Board follows Opinion 9.031 (recently renumbered to 9.4.2 and 9.3.2) of the American Medical Association Code of Medical Ethics that emphasizes that a physician has an ethical obligation to report colleagues who are impaired, incompetent, or unethical. See Board of Medical Practice, Reporting Impaired, Incompetent, or Unethical Colleagues. A failure to do so could lead to charges of unprofessional conduct. 26 V.S.A. §§ 1317(a) & 1354(b). Opinions 9.4.2 and 9.3.2 can be found here.

According to the Board’s interpretation of Opinion 9.031, incompetence that poses an immediate threat to the health and safety of patients should be reported directly to the Board. Id.

Can a failure to comply with the Hospital Patients’ Bill of Rights result in disciplinary action against physicians?
The Vermont Patients’ Bill of Rights, 18 V.S.A. § 1852(b), also creates standards of conduct that physicians must follow when treating patients admitted to hospitals on an inpatient basis. The violation of the following patients’ rights may be reported to the Board and constitute unprofessional conduct:
• The patient has the right to considerate and respectful care at all times and under all circumstances with recognition of his or her personal dignity.
• The patient shall have an attending physician who is responsible for coordinating a patient’s care.
• The patient has the right to obtain, from the physician coordinating his or her care, complete and current information concerning diagnosis, treatment, and any known prognosis in terms the patient can reasonably be expected to understand. If the patient consents or if the patient is incompetent or unable to understand, immediate family members, a reciprocal beneficiary, or a guardian may also obtain this information. The patient has the right to know by name the attending physician primarily responsible for coordinating his or her care.
• Except in emergencies, the patient has the right to receive from the patient's physician information necessary to give informed consent prior to the start of any procedure or treatment, or both. Such information for informed consent should include but not necessarily be limited to the specific procedure or treatment, or both, the medically significant risks involved, and the probable duration of incapacitation. Where medically significant alternatives for care or treatment exist, or when the patient requests information concerning medical alternatives, the patient has the right to such information.
The patient also has the right to know the name of the person responsible for the procedures or treatment, or both.

- The patient has the right to refuse treatment to the extent permitted by law. In the event the patient refuses treatment, the patient shall be informed of the medical consequences of that action and the hospital shall be relieved of any further responsibility for that refusal.

- The patient has the right to every consideration of privacy concerning the patient's own medical care program. Case discussion, consultation, examination, and treatment are confidential and shall be conducted discreetly. Those not directly involved in the patient's care must have the permission of the patient to be present. This right includes the right, upon request, to have a person of one's own sex present during certain parts of a physical examination, treatment, or procedure performed by a health care professional of the opposite sex; and the right not to remain disrobed any longer than is required for accomplishing the medical purpose for which the patient was asked to disrobe. The patient has the right to wear appropriate personal clothing and religious or other symbolic items so long as they do not interfere with diagnostic procedures or treatment.

- The patient has the right to expect that all communications and records pertaining to his or her care shall be treated as confidential. Only medical personnel, or individuals under the supervision of medical personnel, directly treating the patient, or those persons monitoring the quality of that treatment, or researching the effectiveness of that treatment, shall have access to the patient's medical records. Others may have access to those records only with the patient’s written authorization.

- The patient has the right to expect that within its capacity a hospital shall respond reasonably to the request of a patient for services. The right shall include if physically possible a transfer to another room or place if another person in that room or place is disturbing the patient by smoking or other unreasonable actions. When medically permissible a patient may be transferred to another facility only after receiving complete information and explanation concerning the needs for and alternatives to such a transfer. The institution to which the patient is to be transferred must first have accepted the patient for transfer.

- The patient has the right to know the identity and professional status of individuals providing service to him or her, and to know which physician or other practitioner is primarily responsible for his or her care. This includes the patient’s right to know of the existence of any professional relationship among individuals who are treating him or her, as well as the relationship to any other health care or educational institutions involved in his or her care.

- The patient has the right to be advised if the hospital proposes to engage in or perform human experimentation affecting the patient’s care or treatment. Participation by patients in clinical training programs or in the gathering of data for research purposes shall be voluntary. The patient has the right to refuse to participate in such research projects.

- The patient has the right to expect reasonable continuity of care. The patient has the right to be informed by the attending physician of any continuing health care requirements following discharge.

- The patient has the right to receive an itemized, detailed, and understandable explanation of charges regardless of the source of payment.
• The patient has the right to know what hospital rules and regulations apply to his or her conduct as a patient.
• Whenever possible, guardians or parents have the right to stay with their children 24 hours per day. Whenever possible, agents, guardians, reciprocal beneficiaries, or immediate family members have the right to stay with terminally ill patients 24 hours a day.
• A patient who does not speak or understand the predominant language of the community has a right to an interpreter if the language barrier presents a continuing problem to patient understanding of the care and treatment being provided. A patient who is hard of hearing has a right to an interpreter if the impairment presents a continuing problem to patient understanding of the care and treatments being provided.
• The patient has the right to receive professional assessment of pain and professional pain management.
• The patient has the right to be informed in writing of the availability of hospice services and the eligibility criteria for those services.
• The patient has the right to know the maximum patient census and the full-time equivalent numbers of registered nurses, licensed practical nurses, and licensed nursing assistants who provide direct care for each shift on the unit where the patient is receiving care.
• A summary of the hospital’s obligations under this section, written in clear language and in easily readable print, shall be distributed to patients upon admission and posted conspicuously at each nurse’s station. Such notice shall also indicate that as an alternative or in addition to the hospital’s complaint procedures, the patient may directly contact the licensing agency or the Board of Medical Practice. The address and phone number of the licensing agency and Board of Medical Practice shall be included in the notice.

Does the Board have a policy on the termination of the physician-patient relationship?
The Board issued a policy statement in 1999 to provide clarification on the termination of the physician-patient relationship. Abandonment of a patient constitutes unprofessional conduct, and the Board has stated that when presented with a complaint of abandonment, the Board will consider:
• Whether the physician gave the patient timely notice of the termination (at least 30 days);
• Whether the physician provided necessary treatment for an existing problem and/or emergency care during the transition period (at least 30 days); and
• Whether the physician diligently transferred records to another physician chosen by the patient.
The notice of termination should be in writing and delivered to ensure that the patient receives the notice, and all records should be transferred promptly regardless of any outstanding bills.

Does the Board have a policy on the use of controlled substances for the treatment of pain?
In 2014, the Board adopted a Policy on the Use of Opioid Analgesics in the Treatment of Chronic Pain, which updated a previous policy regarding the use of controlled substances for the treatment of pain.

This policy pertains only to the treatment of chronic pain. It largely follows the revised model policy of the Federation of State Medical Boards. It constitutes the Board’s view as to how to
best meet the standard of care when engaged in this aspect of medical practice. As noted in the
document itself, the policy provides guidelines, but on its own, will not be the basis for an
allegation of unprofessional conduct. See Policy at page 4. That being said, the guidelines, in
part, reflect Vermont and federal laws and regulations. \textit{Id.}

Therefore, any physician who treats chronic pain with controlled substances should be familiar
with and follow these guidelines. These guidelines cover many pages and a complete review is
not provided herein. Generally, the guidelines address responsibility for appropriate pain
management, preventing opioid diversion and abuse, patient evaluation and risk stratification,
development of a treatment plan, initiating an opioid trial, monitoring and adapting the treatment
plan, consultation and referral, medical records, and compliance with other regulations. \textit{Id.}

In addition, in 2014, the Board has adopted a policy on the \textit{Drug Addiction Treatment Act of
2000 (“DATA 2000”) and Treatment of Opioid Addiction in the Medical Office.}

This policy provides guidelines addressing treating opioid addiction. It too largely follows the
revised model policy of the Federation of State Medical Boards and constitutes the Board’s view
as to how to best meet the standard of care when engaged in this aspect of medical practice. It
provides guidelines, but on its own, will not be the basis for an allegation of unprofessional
conduct. See Policy at Page 4. Again, however, the guidelines, in part, reflect Vermont and
federal laws and regulations. \textit{Id.}

Therefore, any physician who treats opioid addiction should be familiar with and follow these
guidelines. They cover many pages and a complete review in not provided herein. Generally, the
guidelines address federal requirements for prescribing buprenorphine for addiction, prescription
requirements, Board requirements, State of Vermont opioid addiction treatment programs and
standards, physician qualifications, patient assessment, treatment planning, patient education,
preventing and managing relapse, and medical records. \textit{Id.}

When prescribing controlled substance, physicians should also be aware of the Vermont
Department of Public Health \textit{Rule Governing the Prescribing of Opioids for Chronic Pain} and
the \textit{Vermont Prescription Monitoring System Rule}. Both of these Rules have been revised, and
the new rules, which go into effect July 1, 2017, can be found \textit{here.}

\textit{Does the Board have a policy on the use of telemedicine technologies in the practice of medicine?}

In 2015, the Board adopted a policy on the \textit{Appropriate Use of Telemedicine Technologies in the
Practice of Medicine}. It largely follows the model policy of the Federation of State Medical
Boards. The policy addresses how to determine when physician-patient relationship is
established, assuring privacy of patient data, guaranteeing proper evaluation and treatment of the
patient, and limiting the prescribing and dispensing of certain medications. Generally, a
physician using telemedicine technologies in the provision of medical services to a patient
(whether existing or new) must take appropriate steps to establish the physician-patient
relationship and conduct all appropriate evaluations and history of the patient consistent with
traditional standards of care for the particular patient presentation, and as required by Vermont
law. \textit{Id} at 2. Any physician who uses telemedicine technologies should be familiar with this
policy.
Does the Board have a policy on the treatment of Lyme disease and other tick-borne illness?

By an act of the Legislature, Act 134 of 2014, the Board has adopted a policy addressing the treatment of Lyme disease and other tick-borne illnesses. It can be found here. The policy provides that a health care professional licensed by the Board who diagnoses a patient as having Lyme disease, another tick-borne illness, or a related coinfection:

- Must document the basis for the diagnosis and the treatment for Lyme disease, other tick-borne illness, or coinfection in the patient’s medical record;
- Must provide information to assist patients’ understanding of available Lyme disease tests, the meaning of a diagnostic Lyme disease test result, and any limitations to that test result; and
- Must obtain a patient’s informed consent in writing prior to ordering or administering any proposed long-term treatment for Lyme disease, other tick-borne illness, or coinfection. The Board considers long-term treatment to be a planned course of treatment expected to extend for more than 28 days.

If a physician meets these requirements, the Board will not pursue disciplinary action solely for the use of medical care recognized by the guidelines of the Centers for Disease Control and Prevention, Infectious Diseases Society of America, or International Lyme and Associated Diseases Society for the treatment of a patient’s symptoms when the patient is clinically diagnosed with Lyme disease or other tick-borne illness; however, this does not preclude discipline for errors, omissions, or other unprofessional conduct when practicing within such guidelines.

COMPLAINT PROCESS

What kind of complaints does the Board investigate?

The Board investigates all complaints and charges of unprofessional conduct against any holder of a license or certificate, and holds hearings to determine whether such charges are substantiated or unsubstantiated. 26 V.S.A. § 1353(2). Anyone wishing to make a complaint of unprofessional conduct against a physician, podiatrist, physician assistant, or anesthesiologist assistant may file a written complaint with the Board. 26 V.S.A. § 1355. The Board can also open an investigation on its own initiative to evaluate instances of possible unprofessional conduct. 26 V.S.A. § 1355(a).

A complaint must be signed and include a release of medical records form. Additionally, any hospital, clinic, community mental health center, or other health care institution in which a licensee performs professional services shall report to the Board any disciplinary action taken by it or its staff which significantly limits the licensee’s privilege to practice or leads to suspension or expulsion from the institution, a nonrenewal of medical staff membership, or the restrictions of privileges at a hospital taken in lieu of, or in settlement of, a pending disciplinary case related to unprofessional conduct. 26 V.S.A. § 1317(a).

In addition, within 30 days of any judgment or settlements involving a claim of professional negligence by a licensee, any insurer of the licensee shall report the information to the Commissioner of Health and, to the extent the claim relates to the provision of mental health services, to the Commissioner of Mental Health. 26 V.S.A. § 1317(b).
A person who fails to make one of the mandatory reports above shall be subject to a civil penalty of not more than $10,000.00. 26 V.S.A. § 1317(c).

In addition, the Board has authority to undertake any actions and procedures to carry out its duties, including, but not limited to the following:

- Issue subpoenas and administer oaths;
- Take or cause depositions to be taken;
- Require a licensee or applicant to submit to a mental or physical examination, and an evaluation of medical knowledge and skill by individuals or entities designated by the Board if the Board has a reasonable basis to believe a licensee or applicant may be incompetent or unable to practice medicine with reasonable skill and safety;
- Investigate all complaints of illegal practice of medicine and refer any substantiated illegal practice of medicine to the Office of the Attorney General or the State’s Attorney in the county in which the violation occurred;
- Obtain from the Vermont Crime Information Center a Vermont criminal history record, an out-of-state criminal history record, and a criminal history record from the Federal Bureau of Investigation, for any applicant, licensee, or holder of certification. The Board may also inquire of Interpol for any information on criminal history records of an applicant, licensee, or holder of certification; and
- Inquire of the Vermont Department for Children and Families or of the Vermont Department of Disabilities, Aging, and Independent Living to determine whether any applicant, licensee, or holder of certification who may provide care or treatment to a child or a vulnerable adult is listed on the Child Protection Registry or the vulnerable adult abuse, neglect, and exploitation registry.

**How does the Board conduct its investigation?**

As an initial matter, the Board will send the respondent a copy of the complaint, a copy of the release of medical records signed by the patient or other authorized person, a copy of the grounds of unprofessional conduct, and a standard letter stating that:

- The complaint has been lodged against him or her;
- The letter is not a notice of a formal hearing;
- The matter will be investigated by a committee of the Board working with the Attorney General’s Office; and
- Respondent’s answer should be addressed to the North, Central or South Committee at the address of the Board and filed with the Board within 10 days of the date of the letter.

Board Rules 13.3. While the Rules state 10 days, in practice, the Board generally allows for 21 days to respond and grants extensions when necessary.

One of the three geographic investigating committees, or one specially appointed, and an assistant attorney general will investigate each complaint and recommend disposition to the Board. An investigator from the Board will assist the committee. Board Rules 14.1. Each committee consists of Board members, including at least one public member. V.S.A. §1355(a). After the file is received, the committee will discuss the complaint and plan the investigation. All complaints are investigated. Board Rules 14.2.
What happens when the Board finishes its investigation?
Once the committee is satisfied that the investigation is complete, it shall present its recommendation for final disposition to the Board. The committee may recommend one of five possible dispositions depending on the results of the investigation:

- Concluding the investigation;
- Settlement;
- Specification of charges;
- Interim suspension; or
- Summary suspension. See Board Rules 15.1.

If, after investigating the complaint, the committee and the assistant attorney general are convinced that the alleged misconduct does not constitute unprofessional conduct, then the committee must recommend that the Board conclude the investigation. A concluded investigation may be reopened if new evidence is received or an additional complaint is made. Board Rules 15.1(a).

Under what circumstances might the Board issue a stipulated settlement and consent order?
When an investigation demonstrates a case of unprofessional conduct, the committee may recommend disposition, including the possibility of stipulated settlements and consent orders. Board Rules 15.1(b).

Recommended settlements include a concession of wrongdoing by the licensee, terms and conditions, an understanding that the concession may be relied on by the Board in case the licensee is later found to have engaged in unprofessional conduct, and an understanding that this final disposition of the complaint is public and that the Board shall notify the Federation of State Medical Boards Board Action Data Bank, the National Practitioner Data Bank and the Healthcare Integrity and Protection Data Bank, and may notify other states of its contents. The entire agreement as drafted by the committee and the assistant attorney general will be expressly conditioned on acceptance by the Board. Board Rules 15.1(b).

Under what circumstances might the Board summarily suspend a physician’s license?
The committee may find that certain alleged misconduct poses so grave a threat to the public health, safety or welfare that emergency action must be taken. In such a case, the committee will promptly schedule a hearing and recommend that the Board order summary suspension of the respondent's license pending a hearing under the authority of 3 V.S.A. § 814(c). If the Board orders summary suspension, a hearing will be scheduled as soon as practical and the assistant attorney general will present the case against the suspended licensee. Board Rules 15.1(d).

Under what circumstances might the Board file a formal specification of charges?
If the complaint alleges unprofessional conduct and the committee believes a settlement cannot be reached or is not warranted on the facts, the committee shall recommend the filing of a specification of charges with the Board, setting out the allegations against the licensee in accordance with the provisions of the Administrative Procedure Act, 3 V.S.A. § 809. Board Rules 15.1(c).
The assistant attorney general will draft a specification of the charge or charges of unprofessional conduct. The charges, together with a notice of hearing, shall be served upon the respondent. 26 V.S.A. § 1356.

What happens once a formal specification of charges is filed?
The Board commences disciplinary proceedings by serving a specification of charges and a notice of hearing upon the respondent. The hearing is scheduled no sooner than 30 days after service. The notice shall tell the respondent that he or she may file a response within 20 days of service and state that the respondent has a right to appear at the hearing with counsel and produce their own witnesses and evidence. 26 V.S.A. § 1357, Board Rules 16.1.

If the respondent does not respond to charges or appear at a hearing, after proper notice, the allegations of the charges shall be treated as proven and the Board may take disciplinary action. Upon a request by the respondent and a showing of good cause, the Board may remove a default judgment and schedule a new hearing. Board Rules 16.1.

After a specification of charges has been filed, the Board, or its legal counsel on its behalf, shall have authority to conduct a prehearing conference or discovery conference and to issue orders regulating discovery and depositions, scheduling, motions by the parties, and such other matters as may be necessary to ensure orderly preparation for hearing. Board Rules 16.2.

The hearing will be conducted according to the contested case provisions of the Administrative Procedure Act, 3 V.S.A. §809-815. The Board may authorize its legal counsel to preside at hearings for the purpose of making procedural and evidentiary rulings. A presiding officer may administer oaths and affirmations, rule on offers of proof and receive relevant evidence, regulate the course of the hearing, convene and conduct prehearing conferences, dispose of procedural requests and similar matters, and take any other action authorized by the Administrative Procedure Act. Board Rules 16.3. A hearing panel consists of five members of the Board, including at least one public member, excluding Board members who participated in the investigation of the complaint. 26 V.S.A. § 1360.

The burden of proof in a disciplinary action shall be on the state to show by a preponderance of the evidence that the person has engaged in unprofessional conduct. 26 V.S.A. § 1354(c).

If a majority of the members of the board vote in favor of finding the person complained against guilty of unprofessional conduct as specified in the charges, or any of them, the Board shall prepare written findings of fact, conclusions and order within a reasonable time of the closing of the record in the case. A decision and order is effective upon entry. Notice of the decision and order will be sent to the respondent by certified mail and to the respondent’s attorney, the complainant, and the prosecuting attorney by regular mail. 26 V.S.A. § 1361, Board Rules 16.4.

On what other basis might the Board bring unprofessional conduct charges against a physician?
Upon receipt of the certified copy of the judgment of conviction of a crime for which a licensee may be disciplined for unprofessional conduct, the Board may immediately suspend that person's license until the time for appeal has elapsed and no appeal has been taken, or until the judgment of conviction has been affirmed on appeal or has otherwise become final, and until further order.
of the Board. The Board shall notify the licensee whose license has been suspended under this section and advise the licensee of his or her right to request a hearing, within 90 days. At such hearing, the licensee shall have the burden of showing why the suspension should not remain in effect pending appeal. The disciplinary hearing shall not be commenced until all appeals from the conviction are concluded unless the licensee requests that the matter not be deferred. The sole issue to be determined at such hearing shall be the nature of the disciplinary action to be taken by the Board. An interim suspension shall automatically terminate if the licensee demonstrates that the conviction which served as the basis of the interim suspension has been reversed or vacated. However, a reversal or vacated conviction shall not prohibit the Board from pursuing disciplinary action based on any cause other than the overturned conviction. 26 V.S.A. § 1365.

**DISCIPLINE**

*What are potential board actions for unprofessional conduct?*
Physicians found guilty of unprofessional conduct either after a hearing or by entering into a settlement can face a range of actions that the Board determines proper, including but not limited to:

- Reprimands;
- Conditioning of license;
- Limiting of license;
- Suspension of license; and
- Revocation of license.

See 26 V.S.A. § 1361(b).

**APPELLATE AVENUES**

*What does an appeal consist of?*
A party aggrieved by a final order of the Board may, within 30 days of the order, appeal that order to the Vermont Supreme Court on the basis of the record created before the Board. 26 V.S.A. § 1367; Board Rules 18.1.

**PUBLIC ACCESS TO DISCIPLINARY AND LICENSING INFORMATION**

*What information about physicians is published by the Vermont Department of Health on the Department website?*
Vermont requires the Department of Health to maintain a data repository and to publish profiles of all health care professionals licensed, certified or registered by the department. The information is collected through the physicians’ license renewal applications, and physicians must update the Department of Health with any changes. 26 V.S.A. § 1368. The Vermont Physician Profile, which can be viewed at [http://healthvermont.gov/health-professionals-systems/board-medical-practice/look-license](http://healthvermont.gov/health-professionals-systems/board-medical-practice/look-license), is comprised of the following information provided by physicians:

- A description of any criminal convictions for felonies and serious misdemeanors, as determined by the commissioner of health, within the most recent 10 years. A person
shall be deemed to be convicted of a crime if he or she pleaded guilty or was found or adjudged guilty by a court of competent jurisdiction.

- A description of any charges to which a health care professional pleads nolo contendere or where sufficient facts of guilt were found and the matter was continued without a finding by a court of competent jurisdiction.
- A description of any formal charges served, findings, conclusions, and orders of the licensing authority, and final disposition of matters by the courts within the most recent 10 years, and a summary of the final disposition of such matters indicating any charges that were dismissed and any charges resulting in a finding of unprofessional conduct.
- A description of any formal charges served by licensing authorities, findings, conclusions, and orders of such licensing authorities, and final disposition of matters by the courts in other states within the most recent 10 years.
- A description of revocation or involuntary restriction of hospital privileges for reasons related to competence or character that has been issued by the hospital's governing body or any other official of the hospital after procedural due process has been afforded, or the resignation from, or nonrenewal of, medical staff membership or the restriction of privileges at a hospital taken in lieu of, or in settlement of, a pending disciplinary case related to competence or character in that hospital. Only cases which have occurred within the most recent 10 years shall be disclosed by the Board to the public.
- All medical malpractice court judgments and all medical malpractice arbitration awards in which a payment is awarded to a complaining party during the last 10 years, and all settlements of medical malpractice claims in which a payment is made to a complaining party within the last 10 years. The following statement shall accompany information concerning all settlements: “Settlement of a claim may occur for a variety of reasons which do not necessarily reflect negatively on the professional competence or conduct of the health care professional. A payment in settlement of a medical malpractice action or claim should not be construed as creating a presumption that medical malpractice has occurred.”
- The names of medical professional schools and dates of graduation.
- Graduate medical education.
- Specialty board certification.
- The number of years in practice.
- The names of the hospitals where the health care professional has privileges.
- Appointments to medical school or professional school faculties, and indication as to whether the health care professional has had a responsibility for teaching graduate medical education within the last 10 years.
- Information regarding publications in peer-reviewed medical literature within the last 10 years.
- Information regarding professional or community service activities and awards.
- The location of the health care professional’s primary practice setting.
- The identification of any translating services that may be available at the health care professional’s primary practice location.
- An indication of whether the health care professional participates in the Medicaid program and is currently accepting new patients.
The Department of Health shall provide individual health care professionals with a copy of their profiles prior to the initial release to the public and each time a physician’s profile is modified or amended. A health care professional shall be provided a reasonable time to correct factual inaccuracies that appear in such profile, and may elect to have his or her profile omit items 12 through 14 above. 26 V.S.A. §1368(b).

What other information about physicians is available on the Internet?
Physician profiles, similar to those provided by the Vermont Department of Health, can be found on most state Medical Board web sites. These state web sites contain various information ranging from demographic profiles to malpractice settlements. In addition to the state funded profiles, many private organizations provide information about their members. Each individual organization should be contacted to correct any information. See also the section on the National Practitioner Data Bank.

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REPORTING & DISCLOSURE REQUIREMENTS

Topics Covered in this Chapter:
Reports of Abuse, Possible Criminal Activity or Potential Harm to Third Persons
Reports of Specific Health Conditions or Treatments
Reporting to the Department of Motor Vehicles
Reports Relating to Licensed Health Care Providers and Facilities
Disclosures Related to Identification of Patients and Deceased Patients
Confidentiality of Health Care Records After Disclosure
Enforcement Authority
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Vermont law contains numerous provisions that require or permit health care providers and other professionals to report or disclose particular information to certain state or local government entities. This page of the Health Law Guide highlights most of these reporting or disclosure requirements.

REPORTS OF ABUSE, POSSIBLE CRIMINAL ACTIVITY OR POTENTIAL HARM TO THIRD PERSONS

How must information about the abuse of a vulnerable adult be reported?
A mandatory reporter who suspects, knows of, or has information relating to the abuse, neglect or exploitation of a vulnerable adult is required to report that information within 48 hours to the commissioner of the Department of Aging and Independent Living (DAIL). 33 V.S.A. §§ 6903-6904. A report may also be made to a law enforcement officer, if desired by the reporter.

Under the law “mandatory reporters” include physicians, osteopaths, chiropractors, physician assistants, nurses, licensed nurse assistants, emergency medical services personnel, dentists, psychologists, hospitals, nursing homes, residential care homes, home health agencies, school administrators, teachers, other school employees, social workers, mental health professionals and others. 33 V.S.A. § 6903.

A vulnerable adult is defined as a person who is 18 or older, and:
• Is a resident of a nursing home, psychiatric care hospital or psychiatric care unit within a hospital
• Is receiving personal care services at home for in excess of a month from a home health agency or
• Regardless of residence or services received, has a disability or impairment and requires the assistance of others to meet his or her daily needs or to protect him or herself from abuse, neglect or exploitation.
33 V.S.A. § 6902(14).
The report to the commissioner of DAIL may be in writing or be made orally so long as it is followed within one week by a report in writing. The report must contain the following information:

- The name and address of the reporter, the vulnerable adult and persons responsible for his or her care
- A description of the adult’s age and disability and
- The nature and extent of abuse, neglect or exploitation together with any evidence of previous abuse, neglect, or exploitation of the vulnerable adult.

33 V.S.A. § 6904.

A penalty up to $500 per violation may be imposed on any mandatory reporter who willfully fails to report as required by this law. For purposes of the penalty provision, each 24-hour period that passes beyond the initial 48-hour period, during which the report was due, will constitute a separate violation and additional penalties may be imposed up to a maximum of $5,000 for each reportable incident. 33 V.S.A. § 6913.

Anyone who makes a report in good faith under this statute will be immune from any civil or criminal liability for making the report. 33 V.S.A. § 6908.

**How must information about the abuse of a child be reported?**

Under Vermont’s child abuse law, any mandatory reporter who has reasonable cause to believe that any child has been abused or neglected is required to make a report to the Commissioner of the Department of Children and Families (DCF) within 24 hours of the time information regarding the suspected abuse or neglect was first received or observed. 33 V.S.A. §§ 4913, 4914.

Mandatory reporters include physicians (including residents and interns), surgeons, osteopaths, chiropractors, physician assistants, hospital administrators, nurses, medical examiners, emergency medical personnel, dentists, psychologists, pharmacists, other health care providers, individuals employed by a school district (e.g., school administrators, teachers and other school employees), childcare workers, mental health professionals, social workers, probation officers, police officers, camp counselors and employees, clergy and others. 33 V.S.A. § 4913.

The report to the Commissioner of DCF may be made orally or in writing. Oral reports must be followed-up with a written report upon a request from DCF. The report must contain certain information including the following:

- The name and address or other contact information of the reporter,
- The names and addresses of the child, the parents and/or others responsible for the child’s care
- The child’s age and
- The nature and extent of injury and any evidence of abuse and neglect (current or previous) of the child or the child's siblings.

33 V.S.A. § 4914.

A penalty up to $500 will be imposed on any mandatory reporter who fails to report as required by this law. 33 V.S.A. § 4913.
Anyone, other than someone suspected of child abuse, who makes a report in good faith under this statute will be immune from any civil or criminal liability for making the report. 33 V.S.A. § 4913.

**Do firearm wounds need to be reported?**
Every physician or hospital or other institution attending or treating a case of bullet wound, gunshot wound, powder burn or other injury caused by the discharge of a gun, pistol or other firearm must report such case to local law enforcement officials or the state police, unless the injured person is a member of the armed forces and was on duty when the injury occurred. A penalty of up to $100 will be imposed on anyone who fails to report as required by this law. 13 V.S.A. § 4012.

**Are health care providers required to report blood alcohol levels?**
If a health care provider who is providing health services to a person in the emergency room of a health care facility as a result of a motor vehicle accident becomes aware as a result of any blood test performed in the health care facility that the person's blood alcohol level meets or exceeds the level prohibited by law, the health care provider shall report that fact, as soon as is reasonably possible, to a law enforcement agency having jurisdiction over the location where the accident occurred. A penalty up to $500 will be imposed on anyone who fails to report as required by this law. Anyone who makes a report in good faith under this statute will be immune from any civil or criminal liability for making the report. 23 V.S.A. § 1203b.

**Do health care providers have to report information about crimes against minors?**
Yes, physicians, dentists, chiropractors and nurses are required to disclose information indicating that a patient who is under 16 has been a victim of a crime. Although the statute provides no further provisions, presumably this disclosure would be made to law enforcement or other entities authorized to request such information. 12 V.S.A. § 1612(b).

**Are unusual or suspicious deaths reported?**
When a person dies in any of the following manners, any doctor notified of the death or the superintendent of a psychiatric hospital must immediately notify the medical examiner who resides nearest the town where the death occurred:

- By violence
- Suddenly when in apparent good health
- When unattended by a physician or a recognized practitioner of a well-established church
- By casualty
- By suicide
- As a result of injury
- When in jail or prison
- When in any psychiatric hospital,
- In any unusual, unnatural, or suspicious manner, or
- In circumstances involving a hazard to public health, welfare, or safety

18 V.S.A. § 5205(a).
What information must be disclosed to a medical examiner?
A physician, dentist, chiropractor, mental health professional or nurse is required to produce information as to the mental or physical condition of a deceased individual if requested to do so by the chief medical examiner 12 V.S.A. § 1612 (c)(2).

Who must report possible cases of bioterrorism and what are the requirements for reporting?
Health care providers are required to report in writing within 24 hours to the Commissioner of Health all cases of persons who exhibit any illness, disease, injury or death identified by the Department of Health as likely to be caused by a weapon of mass destruction, which may include illnesses, diseases, injuries or deaths which can result from bioterrorism, epidemic or pandemic disease, or novel and highly fatal infectious agents or biological toxins or which may be caused by specific biological agents. 13 V.S.A. § 3504(a).

For purposes of this bioterrorism reporting, out-of-state medical laboratories that have agreed to the reporting requirements of Vermont are required to comply with the reporting requirements of health care providers. Results must be reported by the laboratory that performs the test, but an in-state laboratory that sends specimens to an out-of-state laboratory is also responsible for reporting results. 13 V.S.A. § 3504(e).

Similarly, pharmacists are required to report in writing within 24 hours to the Commissioner of Health any unusual or increased prescription requests, unusual types of prescriptions, or unusual trends in pharmacy visits that may result from bioterrorist acts, epidemic or pandemic disease, or novel and highly fatal infectious agents or biological toxins, and might pose a substantial risk of a significant number of human fatalities or incidents of permanent or long-term disability. Prescription-related events that require a report include an unusual increase in the number of prescriptions to treat fever, respiratory or gastrointestinal complaints, an unusual increase in the number of prescriptions for antibiotics, an unusual increase in the number of requests for information on over-the-counter pharmaceuticals to treat fever, respiratory or gastrointestinal complaints, and any prescription that treats a disease that is relatively uncommon and may be the result of bioterrorism. 13 V.S.A. § 3504(b).

In addition, veterinarians, livestock owners, veterinary diagnostic laboratory directors or other persons having the care of animals, are required to report within 24 hours to the Commissioner of Health any animals having or suspected of having any disease that can result from bioterrorism, epidemic or pandemic disease, or novel and highly fatal infectious agents or biological toxins, and might pose a risk of a significant number of human and animal fatalities or incidents of permanent or long-term disability. 13 V.S.A. § 3504(d).

The reports required by this law must be made to the Commissioner of Health and, for reports of health care providers and pharmacists, must include all available information such as the patient's name, date of birth, sex, race and current address (including city and county), the name and address of the health care provider and of the reporting individual, if different. Reports from veterinarians and others relating to animals must include all available information such as the location or suspected location of the animal, the name and address of any known owner, and the name and address of the reporting individual. 13 V.S.A. § 3504.
Health care providers who make good faith reports to the Department of Health under this law will be immune from prosecution, suit, administrative or regulatory sanctions for defamation, breach of confidentiality or privacy, or any other cause of action based on such reports or errors contained in such reports.

_Do health care providers have a duty to report, or “warn,” when patients threaten harm to identifiable individuals?_

When a mental health provider has good reason to believe that a patient poses a serious risk of danger to an identifiable individual, the provider has a duty to exercise reasonable care to protect the identifiable victim from that danger even if it requires the disclosure of confidential patient information. This “duty to warn,” which has evolved in case law in Vermont and other states, is an exception to the patient privilege of confidentiality.

The duty to warn exception originated in California with the case of Tarasoff v. Regents of the University of California, 551 P.2d 334 (1976). In Tarasoff, a university hospital psychologist was told by his patient that the patient intended to kill a woman, Tatiana Tarasoff. Two months later, he did so. The Supreme Court of California ruled that the “public policy favoring protection of patient-psychotherapist communications must yield to the extent to which disclosure is essential to avert danger to others.” 551 P.2d at 347.

In Peck v. Counseling Service of Addison County, the Vermont Supreme Court adopted the Tarasoff ruling in a situation where a patient receiving mental health services threatened to burn his father’s barn down. 146 Vt. 61 (1985). The Court held that if a mental health provider knows or should know that his patient poses a serious risk of danger to an identified person, the provider has a duty to take whatever steps are reasonably necessary to protect the identifiable victim from that danger, which could include reporting confidential patient information. Id. The health care provider has a duty to exercise due care in determining what steps may be necessary to protect the identifiable victim of a patient’s threat of harm and what confidential information must be disclosed. Id. at 67. The health care provider must report confidential information discretely, and in a fashion that would preserve the privacy of the patient to the fullest extent compatible with the prevention of the threatened danger, insuring that only that information which is necessary to protect the potential victim is revealed. Peck, 146 Vt. at 67-68, citing Tarasoff, 7 Cal.3d at 441.

Although to date the Vermont courts have not had occasion to extend the waiver of physician-patient confidentiality beyond mental health services, other states have done so when a patient is incapacitated as a result of medical treatment or disease and the person poses an obvious risk of serious harm to others. In these situations, a health care provider may have a duty to warn reasonably identifiable potential victims or a duty to take action to avoid the harm.

_Do health care providers have a duty to report or warn of potential acts of violence?_

The Vermont Supreme Court has recognized that there are additional circumstances where a mental health provider has a duty to warn (and thus disclose confidential patient information) even when there is no identifiable individual to whom the patient poses a risk. In Kuligoski v. Brattleboro Retreat, et al., the Vermont Supreme Court held that mental health providers have a
duty to warn individuals within a “zone of danger” that a patient with dangerous propensities presents a serious risk of harm. 2016 WL 5793088 (September 16, 2016).

The Court did not define the “zone of danger” precisely, but it includes those most likely to be victims if a patient engages in violence. While the Court declined to hold that the duty to warn extends to the general public, it did find that the “zone of danger” includes the identifiable “caretakers” of a potentially violent patient and thus these “caretakers” must be warned.

The Vermont Supreme Court held that the duty to warn applies when: “a caregiver is actively engaged with the patient’s provider in connection with the patient’s care or the patient’s treatment plan (or, in this case, discharge plan), the provider substantially relies on that caregiver’s ongoing participation, and the caregiver is himself or herself within the zone of danger of the patient’s violent propensities.” A “caregiver” does not need to be formally charged with legal responsibility for the patient, as in guardianship.

The Court also held that the “duty to warn” encompasses a “duty to inform,” which the Court described as a duty “to provide sufficient information to [caretakers] so they could fully assume their caretaker responsibilities to assist [the patient] and protect against any harmful conduct in which he might engage.” Mental health providers must not merely warn of potential risks, according to the decision, they must also provide sufficient information to assist the “caretakers” in controlling the patient’s conduct and protecting third parties from harm.

How do I determine if the patient poses a risk of danger to unidentified individuals requiring a warning?

If a patient is considered to have violent propensities, Vermont law places the burden on mental health providers to determine who may be a caregiver or within the “zone of danger” and what information must be disclosed in order to assist the “caretakers” in controlling the patient’s conduct and protecting third parties from harm.

In almost all instances, consultation with a colleague is advised to determine the most appropriate course of action given the conflicting duties imposed on the mental health provider.

How does the duty to warn relate to HIPAA privacy and security rules requirements?

The HIPAA privacy and security rules do not create a duty to warn or mandate disclosure. Rather, under the HIPAA privacy and security rules, a covered entity is permitted to use and disclose protected health information – including psychotherapy notes – if the covered entity believes, in good faith, that the use or disclosure is necessary to prevent or lessen a serious and imminent threat to the health or safety of a person or the public, and the disclosure is to a person or persons reasonably able to prevent or lessen the threat, including the target of the threat. 45 C.F.R. § 164.512(j). A covered entity may also disclose protected health if the covered entity believes, in good faith, that the use or disclosure is necessary for law enforcement to identify or apprehend an individual because of a statement by an individual admitting participation in a violent crime that the covered entity reasonably believes may have caused serious physical harm to the victim, or where it appears from all the circumstances that the individual has escaped from a correctional institution or from lawful custody. Id.
In the above situations, medical records and psychotherapy notes may be disclosed without authorization or consent of the patient. The disclosure must be made to a person or persons reasonably able to prevent or lessen the threat or to the target of the threat. The disclosure must be limited to the minimum amount of information necessary to prevent the harm from occurring.

**Reports of Specific Health Conditions or Treatments**

*What are the requirements for reporting communicable diseases?*

**Reportable Diseases**

A physician, health care provider, infection preventionist, administrator of a long-term care or assisted living facility, laboratory director, nurse practitioner, nurse, physician assistant, or school health official (collectively, “Reporters”) who has reason to believe that a person is sick or has died of a diagnosed or suspected disease that has been identified by the Department of Health as a reportable disease and as dangerous to the public health shall report the diagnosis or suspicion within 24 hours to the Department of Health.

The following diseases must be reported to the Department of Health immediately:

- Anthrax
- Botulism
- Diphtheria
- Individuals cases of influenza due to a novel strain of Influenza A
- Measles
- Meningococcal Disease
- Plague
- Poliovirus infection, including poliomyelitis
- SARS
- Smallpox
- Rabies – Human (post-exposure is reportable irrespective of evidence of disease)
- Tularemia
- Viral Hemorrhagic Fever

Reporters also are required to have policies and procedures in place, which meet a number of criteria set by rule, to ensure confidentiality of the reports. 18 V.S.A. §§ 1001 et seq., VDH Reportable and Communicable Diseases Rule.

The reports are required to include the following information regarding the affected person:

- Name;
- Date of birth;
- Age;
- Sex;
- Address;
- Telephone number;
- Name of health care provider/physician;
- Address of health care provider/physician;
• Disease name;
• Date of onset of disease; and,
• Any other pertinent information.

VDH Reportable and Communicable Diseases Rule, § 5.0.

The report should be made by telephone or in writing or electronically to the Vermont Department of Health, Epidemiology Program (802-863-7240 or 800-640-4374).

**AIDS and HIV Reporting**

HIV and AIDS are both reportable communicable diseases under Vermont law. Prior to performing an HIV test, health care providers must inform the individual to be tested that the provider is required to report a positive result and the individual's name to the Department of Health. 18 VSA § 1001(g). The provider must also tell the patient that there are testing sites that provide anonymous testing that are not required to report positive results. Id. AIDS and HIV reports should be made on the Department of Health’s confidential report form. The Department of Health has established procedures to ensure the confidentiality of the reports it receives related to HIV and AIDS diagnoses. Except for the limited purpose of de-duplication of case records, the Department of Health does not release information relating to HIV and AIDS without prior notice to and written authorization from the individual.


**When must a case of rabies be reported?**

A Reporter is required to report to a local health officer the name, age and address of any person who has been bitten by an animal of a species subject to rabies within 24 hours of notice of the bite. 20 V.S.A. § 3801; VDH Reportable and Communicable Diseases Rule 7.0.

**When must immunizations be reported?**

A health care provider must report to the Department of Health all data regarding immunizations of children under the age of 18 within seven days of the immunization. A health insurer must report to the Department of Health all data regarding immunizations of adults and of children under the age of 18 at least quarterly. In addition, reports of adult immunizations are required commencing within one month after the health care provider has established an electronic health records system. 18 V.S.A. § 1129.

**Who is required to report fetal deaths?**

Physicians, hospitals and funeral directors are required to report certain fetal deaths to the Commissioner of Health within seven days for statistical purposes. 18 V.S.A. § 5222.

Reportable fetal deaths are fetal deaths of 20 or more weeks of gestation, 400 or more grams, 15 or more ounces, or fetal deaths which involve therapeutic or induced abortion regardless of size. If the physician treating a woman for a miscarriage or abortion does not know whether the fetal death has been previously reported, the physician is required to report the death. The reports are not public records and shall be destroyed after five years. Id.
What are the requirements for reporting cancer cases?
Health care facilities and health care providers diagnosing or providing treatment to patients with cancer are required to report each cancer case to the Commissioner of Health within 180 days of admission or diagnosis, except that health care providers are not required to report if the patient has been previously diagnosed or admitted for treatment of cancer at a Vermont facility. 18 V.S.A. § 153. If a health care facility fails to report a cancer diagnosis or admission, the Commissioner has the authority to enter the premises, obtain the information and report it, at the expense of the facility. Id.

Anyone who makes a good faith report in compliance with this law will not be subject to any action for damages. 18 V.S.A. § 156.

Does Vermont allow mammograms to be reported?
Vermont law permits the good faith submission of mammography and pathology data relating to breast cancer to the Vermont Mammography Registry. All information reported to the VMR is confidential and privileged. 18 V.S.A. §§ 154, 155, 157.

Anyone who makes a good faith report in compliance with this law will not be subject to any action for damages. 18 V.S.A. § 156.

Does information have to be reported regarding breast density?
As of January 15, 2017, women in Vermont must receive notification of their breast density classification in the summary of a mammography report. 18 V.S.A. § 158. The law states that all health care facilities that perform mammography examinations must inform all women of their breast tissue classification based on the Breast Imaging Reporting and Data System established by the American College of Radiology. If a woman has heterogeneously dense or extremely dense breasts the report must also contain a notice explaining that dense breast tissue may make it more difficult to detect cancer on a mammogram and may be associated with a slightly increased risk of cancer. The notice must further state that the information is provided to raise awareness and encourage patients to discuss this issue with their health care providers. The full law and language of the notice can be found here.

Do lead poisoning diagnoses have to be reported?
Any laboratory that analyzes blood samples of Vermont residents (includes both adults and children) for lead levels must report to the Department of Health all information required by the Department. All health care providers who analyze blood samples for lead levels or who use laboratories outside Vermont to analyze blood samples for lead levels must report to the Department immediately by telephone if the result of any analysis is 45 micrograms or more of lead per deciliter of blood, or by electronic means within 14 days of analysis if the result of the analysis is less than 45 micrograms of lead per deciliter of blood. All blood lead data reports to the Department must include the name, date of birth, date of blood test, and address of the individual whose blood is analyzed and, if known, the owner of the residence of the individual. 18 V.S.A. § 1755(d).
Are involuntary hospitalizations reported?
The head of the hospital must immediately provide notice to the parents, legal guardian, nearest known relative or interested party (if known), or spouse of an individual who has been involuntarily hospitalized due to a mental health related condition. If the hospital admission was not pursuant to court order, the head of the hospital must also notify the district court judge for the district where the hospital is located. If the hospital admission was pursuant to court order, the head of the hospital must immediately notify the court and the Commissioner of Mental Health of the admission and also of discharge. 18 V.S.A. § 7106.

When an application for involuntary hospitalization is filed with a court, the court will transmit a copy of the application, the physician's certificate (if any) and a notice of hearing to the proposed patient, his or her attorney, guardian, or any person having custody and control of the proposed patient, the state's attorney, or the attorney general, and any other person the court believes has a concern for the proposed patient's welfare. 18 V.S.A. § 7613. A copy of the notice of hearing shall also be transmitted to the applicant and certifying physician. Id. If the court has reason to believe that notice to the proposed patient will be likely to cause injury to the proposed patient or others, it shall direct the proposed patient's counsel to give the proposed patient oral notice prior to written notice under circumstances most likely to reduce likelihood of injury. Id.

REPORTING TO THE DEPARTMENT OF MOTOR VEHICLES

Are physicians obligated to report a patient to the Department of Motor Vehicles if the patient seems unfit to drive?
There is nothing in Vermont law that either authorizes or requires physicians to report drivers to the Department of Motor Vehicles (“DMV”), unless the driver poses such a serious threat that the threshold for the duty to warn is met.

In general, under the case law of Vermont and states throughout the country, a health care provider’s duty to warn requires a health care provider who knows or should know that a patient poses a serious risk of danger to warn potential victims and exercise reasonable care to protect them. Consistent with this duty, a health care provider may have a duty to warn a driver’s “caretakers” that the driver poses a serious risk of harm and to provide sufficient information to assist the “caretakers” in controlling the driver’s conduct and protecting third parties from harm. See Kuligoski v. Brattleboro Retreat, et al., 2016 WL 5793088 (September16, 2016).

Additionally, the HIPAA privacy and security rules permit, but do not require, disclosure of a serious and imminent threat to the health or safety of a person or the public without authorization from the patient. 45 C.F.R. § 164.512(j). The disclosure may only be made to someone who is able to prevent or lessen the threat and disclosure and must be based on a good faith belief that disclosure is necessary to prevent the threatened harm from occurring. Id. Consistent with this, if a health care provider reasonably believes that a patient poses a serious and imminent threat to health and safety by driving, the provider is permitted to disclose the threat to the DMV to prevent or lessen such a threat.

The American Medical Association (“AMA”) provides physicians with ethical guidance about reporting patients who may be impaired drivers. The AMA explains that in deciding whether
and how to intervene when a patient’s medical condition may impair driving, physicians must balance the duty of confidentiality to their patient with the protection of public safety. The AMA advises physicians, within their areas of expertise to:

- Assesses at-risk patient individually for medical conditions that might affect driving, using best professional judgment and keeping in mind that not all physical or mental impairments create an obligation to intervene.
- Before reporting, take initial steps, including a candid discussion with the patient and family and suggesting voluntary steps to improve or limit driving.
- Prior to reporting, physicians should disclose and explain to patients that they may have a responsibility to report a medically at-risk driver.
- Report only the minimal amount of information necessary.

AMA Ethical Opinion 8.2 Impaired Drivers and Their Physicians.

If any person has concerns about an individual’s driving, including a doctor, family member, or neighbor, the Vermont DMV advises them to request, in writing, that a patient be reexamined by the DMV. See more information on the DMV Mature Driver website. A reexamination involves the evaluation of an individual by a DMV examiner and consists of a vision test and a driving test. A written test may be required depending on specific circumstances. Following the reexamination, the examiner will decide whether any action should be taken regarding the individual’s driving privilege, such as restrictions, suspension or revocation of a license or permit.

Reexamination requests can be sent to:
Vermont Department of Motor Vehicles
120 State Street
Montpelier, VT 05603-0001
Attention: DMV Commissioners Office

The request must include the driver’s name, date of birth, address, your name and contact information (address and phone number), your relationship to the driver, and specific reasons for requesting a re-examination (this must be specific and should include details of any personal observations).

See the Vermont DMV Mature Drivers Website: http://dmv.vermont.gov/licenses/renew/mature-drivers; Vermont DMV Universal Medical Evaluation/Progress Report and Driver Eyesight Evaluation.

REPORTS RELATING TO LICENSED HEALTH CARE PROVIDERS AND FACILITIES

What is required for reporting unprofessional conduct?
Hospitals, clinics, community mental health centers and other health care institutions are required to report within 10 days to the Commissioner of Health any disciplinary action taken by it which significantly limits a licensed health care provider’s privilege to practice or leads to suspension or expulsion from the institution. If the disciplinary action taken against the health care provider was based on the provision of mental health services, a copy of the report also must be sent to the Commissioner of Mental Health and the Commissioner of Disabilities, Aging, and
Independent Living. The report must include any supporting information and evidence. Any insurer must report within 30 days of any judgment or settlements involving a claim of professional negligence by a health care provider to the Commissioner of Health and, to the extent the claim relates to the provision of mental health services, to the Commissioner of Mental Health. The reporting requirements of this law do not apply to cases of resignation or separation from service for reasons unrelated to disciplinary action. 26 V.S.A. § 1317.

Anyone who makes a good faith report in compliance with this law will not be liable for damages in any civil action.

**Is reporting of medical errors or unsafe acts required?**

Hospitals are required to report to the Department of Health for the purposes of the Patient Safety Surveillance and Improvement System (“PSSIS”) certain information relating to adverse events and intentional unsafe acts. Each hospital must establish an internal reporting system and develop and implement policies and procedures to identify, track, and analyze reportable adverse events, non-reportable adverse events, and near misses and use that data to improve patient safety. The reports must be submitted using the secure transmission system established by the Department for this purpose. The reportable adverse events are the serious reportable events and specifications as published by the National Quality Forum. The adverse event report must be filed within 7 calendar days of the discovery of the event, and a causal analysis and corrective action plan must be filed no later than 60 calendar days after the initial report. Hospitals are required to disclose to patients, or in the case of death, to an immediate family member, adverse events that cause death or serious bodily injury. 18 V.S.A. §§ 1915, 1916, 1918; Vermont Patient Safety Surveillance and Improvement System Regulation.

Reports of intentional unsafe acts must be filed no later than 7 calendar days after the information available to the hospital supports a reasonable, good faith belief that an intentional unsafe act has occurred. Intentional unsafe acts are defined as adverse events or near misses that result from criminal acts, purposefully unsafe acts, alcohol or substance abuse, or patient neglect, exploitation or abuse. The complete names of the involved in the intentional unsafe act must be provided in the report to the PSSIS. 18 V.S.A. §§ 1915, 1916, 1918; Patient Safety Surveillance and Improvement System Regulation.

All reports made to the Department under the PSSIS have strict confidentiality protections. They are confidential and privileged, exempt from disclosure as a public record and are not subject to discovery or introduction into evidence in civil or administrative proceedings.

**Disclosures related to identification of patients and deceased patients**

**Are providers required to disclose information to identify patients?**

Dentists are obligated to disclose information necessary for the identification of patients. It is presumed that the disclosure would be made to law enforcement or to a medical examiner or other entity authorized to request or receive the information. 12 V.S.A. § 1612(b).
Do health care providers have to disclose information on deceased patients?
Physicians, nurses, and chiropractors are required to disclose information as to the mental or physical condition of deceased patients in certain circumstances (e.g., a will contest) unless the information would be considered “to disgrace the memory of the decedent.” In such circumstances, the privilege must be waived by either the decedent’s personal representative, the surviving spouse of the decedent, or the next of kin of the decedent. In addition, physicians, dentists, chiropractors, mental health professionals, and nurses are required to produce information as to the mental or physical condition of a deceased patient if requested to do so by the chief medical examiner. 12 V.S.A. § 1612(c).

When can the government have access to health care records?
While there are a number of circumstances when government agencies are authorized to review health care records, including patient information or payment information, the following situations are of particular importance to health care providers and facilities:

- Under the federal Medicare program, access to patient information must be granted by a health care provider to the Secretary of the U.S. Department of Health and Human Services, and/or representatives of the Centers of Medicare and Medicaid Services and others involved in enforcement of the requirements of federally funded health programs;
- When seeking to collect reimbursement for the care of a Medicaid beneficiary from a third party, the Department of Vermont Health Access is entitled to obtain any records of the treatment of any individual which are in any way relevant to the treatment paid for through Medicaid. [33 V.S.A. §6705(b)]; and,
- As a condition of participation as a Medicaid provider in Vermont, health care providers must allow the State to review medical records at any time and without advanced notice. Vermont Medicaid Provider Manual, Section 7.2.5.

CONFIDENTIALITY OF HEALTH CARE RECORDS AFTER DISCLOSURE
After health care records have been received by the state agency as permitted or required by state law, the records are generally subject to statutory and regulatory provisions governing their use and further disclosure. These confidentiality provisions vary depending on the reporting requirement. For example, reports received by the Department of Health under the patient safety statute (PSSIS) are confidential and privileged, exempt from disclosure as a public record and are not subject to discovery or introduction into evidence in civil or administrative proceedings. Other of the statutes and regulations may have different requirements permitting or prohibiting further disclosure of the information by the state agency and many of the reports will be subject to the provisions of the HIPAA privacy and security rules. Health care providers are encouraged to review the applicable statutes or regulations or the Consent, Privacy and Medical Records chapter of this Guide to Health Care Law in Vermont for further information about the variety of confidentiality provisions applicable to reports received by state agencies.

ENFORCEMENT AUTHORITY

What enforcement authority does the Department of Health have?
The Vermont Department of Health works collaboratively with health care providers and facilities in public/private partnerships to educate providers and to make them aware of reporting
requirements, and to resolve obstacles to implementation of public health requirements. However, the Department also has significant enforcement authority.

There are a number of specific enforcement provisions related to reporting requirements. For example, a fine of up to $500.00 may be imposed on anyone failing to comply with the fetal death reporting requirements. 18 V.S.A. § 5225.

Similarly, there are also enforcement provisions and authorizations to seek civil penalties related to willful, malicious and negligent disclosure of confidential communicable disease information. Negligent disclosures not authorized by law may be subject to a civil penalty of up to $2500. Penalties range from $10,000 to $25,000 and potential jail time for willful and malicious disclosure.

For further information about these enforcement provisions, health care providers can refer to the specific public health laws. See e.g., 18 V.S.A. §§ 130, 131 (general), §1001 (communicable disease disclosure), §1918 (patient safety reports), §5225 (fetal death); and 26 V.S.A. §1317(e) (unprofessional conduct reports).

ABOUT THE AUTHORS

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Shireen Hart, a shareholder of the firm’s Healthcare Group, draws upon her more than 15 years of experience in the field to provide pragmatic counsel to hospitals, long term care facilities, community mental health agencies, pharmacies, laboratories, individual practitioners and healthcare related vendors on a broad range of matters. Her strong statewide connections and effectiveness working with other attorneys and state agency or department personnel creates a favorable environment in which to deliver positive results for clients.

Lauren Layman’s practice focuses on the areas of health care law and legislative relations. She brings a uniquely effective perspective to her work through her prior experiences as policy advisor for the Centers of Disease Control in Washington, D.C. and a health care staff member for Senator Jim Jeffords. A Vermont native, Lauren is knowledgeable not only about the policies and legal climate affecting the state’s hospitals, but also the patients and communities they work
with on a daily basis. This mix of professional and personal experience enables Lauren to handle cases with a multi-faceted, balanced approach.
REPRODUCTIVE HEALTH/RIGHTS

Topics Covered in this Chapter:
Statewide Health and Sexuality Education
Access to Contraception
Access to Abortion
Non-Traditional Labor and Delivery Care
Coverage for LGBTQ Patients
Assisted Reproductive Technology (ART)
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STATEWIDE HEALTH AND SEXUALITY EDUCATION

Does Vermont require comprehensive sexuality education?
Not exactly, but schools are required to provide Comprehensive Health Education.

What is Comprehensive Health Education?
By definition, Comprehensive Health Education is “systematic and extensive elementary and
secondary educational programs designed to provide a variety of learning experiences based
upon knowledge of the human organism as it functions within its environment.” 16 V.S.A.§131.
In general, the curriculum should include information about body structure and functioning,
sexuality, reproduction, HIV and other sexually transmitted infections (STI), HIV and STI
prevention, how to recognize sexual abuse and sexual violence, and promotion of healthy
relationships. 16 V.S.A.§131.

Is the Health Education Curriculum Standardized?
No, not every student will have the same health education experience. In addition to a religious
exemption, school boards are entitled to create an advisory council for interested community
members; the advisory council is intended to represent various viewpoints while assisting the
school board in curriculum development and implementation. 16 V.S.A §§131, 134, 135. For
more information see: http://education.vermont.gov/student-learning/content-areas/health-
education

ACCESS TO CONTRACEPTION

If my patient has health insurance, what contraceptives are insurance required to cover?
Vermont law requires health insurance plans, including Medicaid and other public assistance
programs to cover outpatient contraceptive services, including voluntary sterilization. Coverage
includes the purchase of all FDA approved prescription contraceptives and contraceptive devices. 8 V.S.A §4099c.

How much of the cost are insurance companies required to cover? 
Annually, an insurance company must provide complete coverage for a contraceptive device, voluntary sterilization, or one year’s worth of prescription contraceptives. 8 V.S.A §4099c.

Are there exceptions to coverage? 
Yes, there are two exceptions to insurance coverage of contraceptives:

1. If a patient is enrolled in an insurance program without prescription drug coverage, the insurance company is not required to cover the cost of prescription contraceptives or prescription contraceptive devices. 8 V.S.A §4099c(b)

2. If a patient with a high deductible insurance plan elects sterilization and insurance coverage of the procedure disqualifies the patient from eligibility for a health savings account under 26 U.S.C §223. 8 V.S.A §4099c(d).

Does contraceptive coverage apply only to the policyholder? 
No. Contraceptive coverage applies to a policyholder, an insured partner (through marriage or civil union), and other insured dependents. 8 V.S.A §4099c(g).

Does contraceptive coverage apply to services beyond the tangible prescription, procedure, or device? 
Yes. In addition to contraceptive prescriptions, devices, voluntary sterilization, and outpatient contraceptive services, insurance companies must cover parallel services including counseling and clinical services associated with access to and use of contraceptive devices, 8 V.S.A §4099c(c)(1), as well as related follow-up services, including management of side effects, counseling for continued adherence, and device insertion and removal. 8 V.S.A §4099c(e).

With regard to contraceptive coverage, is there anything Vermont law covers that the Affordable Care Act does not? 
Yes, the Affordable Care Act only covers female prescription contraception and sterilization procedures, Vermont law is gender inclusive and requires insurance companies to provide coverage for voluntary male sterilization procedures (vasectomy). 8 V.S.A §4099c(d), 45 C.F.R §147.130(a)(I)(iv), https://www.healthcare.gov/coverage/birth-control-benefits/.

ACCESS TO ABORTION 

Are there restrictions to abortion in Vermont? 
No. In 1972, in Beecham v. Leahy, the Vermont Supreme Court clarified a woman’s right to obtain a legal abortion in Vermont. Prior to Beecham, it was legal for a woman to get an abortion, but it was illegal for a doctor to provide an abortion. In Beecham, the Court struck down the criminal statute and held “that the legislature, having affirmed the right of a woman to abort, cannot simultaneously,… prohibit its safe exercise.” 130 Vt.164, 170 (1972).
Abortion is legal in Vermont without restriction, including no waiting periods, or mandated parental involvement. (For more information on minors and abortion can be found in the Minor Consent section of this guide.) Vermont is one of few states that provide state Medicaid funding for medically necessary abortion care. For more information visit: https://www.guttmacher.org/fact-sheet/state-facts-about-abortion-vermont.

What if my patient wants an abortion procedure or pill and I believe their choice is morally wrong?
Unlike most states, Vermont law does not protect physicians or other medical providers’ right to refuse to provide reproductive health care services because of moral or other beliefs. For more information on Conscience Clause legislation visit: https://www.guttmacher.org/gpr/2004/08/new-refusal-clauses-shatter-balance-between-provider-conscience-patient-needs; http://www.thehastingscenter.org/briefingbook/conscience-clauses-health-care-providers-and-parents/.

If I provide a patient with an abortion, do I need to notify anyone?
Yes. Hospitals, physicians, and funeral directors must report fetal death to the Commissioner of the department of public health within 7 days for all therapeutic or induced abortions legally preformed. Fetal death reports are used only for statistical purposes, are not public record, and information is destroyed after 5 years. 18 V.S.A §5222.

Do private insurance companies cover the cost of abortion services?
If a patient has private insurance, coverage of abortion services is dependent on their individual insurance plan. Unlike coverage of contraception, there is no state mandate for insurance companies to cover abortion services.

Does Medicaid cover the cost of abortion?
Federal Medicaid dollars are prohibited from funding an elective abortion through the Hyde Amendment. However, Vermont has no such prohibition on elective abortion coverage, therefore, a patient covered by Medicaid does qualify for Medicaid funded abortion services.

NON-TRADITIONAL LABOR AND DELIVERY CARE

Do my patients have options with regard to assistance for labor and delivery?
Yes. Health insurance plans that provide maternity benefits are required to provide coverage for a licensed midwife at no more cost to a consumer than similar labor and delivery benefits. 8 V.S.A §4099d.

Under this provision, if my patient wishes to give birth at home, can she?
Yes. Midwifery services can be provided at a hospital, health care facility, or in the home. 8 V.S.A §4099d.

COVERAGE FOR LGBTQ PATIENTS
If I have a patient seeking sex reassignment medical services, are there barriers to treatment?
Possibly. Insurance companies cannot unfairly discriminate against individuals because of their gender identity. Vermont law prohibits insurance companies from excluding coverage for medically necessary treatment, including gender reassignment for gender dysphoria or related health conditions. 8 V.S.A §4724

ASSISTED REPRODUCTIVE TECHNOLOGY

The law regarding assisted reproductive technologies (ART) is still in a relatively early stage of development. In some states there is some, although little, case law. Some state legislatures have passed laws covering some aspects of ART; many states have virtually no law on the subject. Vermont has little if any law on any aspect of ART. Therefore, the information contained in this section is a general overview of the law throughout the United States. This information is not to be construed as legal advice and is provided for purposes of information only.

What is artificial insemination and what is the resulting legal status of each of the parties?
Artificial insemination occurs when the sperm of a known or unknown donor is introduced into the reproductive tract of a female for the purposes of conception. Sperm may be fresh or frozen, but is usually frozen in the case of anonymous donors. To ensure that donated sperm does not pose a risk of HIV, a six-month waiting period to permit appropriate testing will be required for contemporaneous donors. A sperm donor is generally held NOT to be the father of any resulting children, with no parental rights or responsibilities (such as child support) toward the child. It is important, however, for the insemination to be done through a medical professional, not simply between the parties. If the latter, it is harder for the sperm donor to retain his status as that of a donor only, and not as the legal father of the child, even if the parties agree or contract otherwise. If the woman is married at the time of the insemination procedure, her husband is deemed to be the legal father of the child.

What is egg donation and what is the resulting legal status of each of the parties?
In egg donation, a female egg donor receives medical treatment to stimulate her ovaries to produce multiple eggs, which are then harvested for use. These eggs are used in in-vitro fertilization procedures, often with the sperm of the intending father, or if not, with donor sperm. Egg donors, like sperm donors, are generally held NOT to be the legal mother of any resulting children.

Can sperm and egg donors be paid for their donations?
Paying sperm donors has been the practice over the last several decades and continues, although amounts per donation are minimal. Because of the risk and complexity of the medical procedures regarding egg donation, payments have tended to be higher than that for sperm donation, sometimes in the thousands of dollars. Professional organizations like the American Society of Reproductive Medicine made recommendations limiting the amount of money that should be paid to gamete donors, particularly egg donors; there have been legal challenges to these actions,
calling them “price-fixing.” Advertisements for “designer” gametes, i.e., gametes from persons at Ivy League schools, or persons with high IQ scores or some other highly-valued attribute can be found on the Internet and in many national and college newspapers. These “designer” gametes are offered at inflated prices, often starting at $50,000 and often more.

**What is surrogacy and what is the resulting legal status of the parties?**

There are two kinds of surrogacy: traditional and gestational. In a tradition surrogacy, the surrogate mother contributes her own ovum, either naturally through artificial insemination with intending father’s sperm, or through in vitro fertilization. She thus has a genetic connection to the child. A traditional surrogate has a good chance of being declared the legal mother of any resulting child should there be a disagreement between the parties during the pregnancy or after the child’s birth. As such, she may be awarded custodial or visitation rights with the child. Some states give traditional surrogates the right to change their minds about their surrogate-only status following the birth of the child. This is the kind of surrogacy that existed in the famous Mary Beth Whitehead case. Traditional surrogacy, because it may result in challenges regarding maternity similar to those in the Whitehead case, is not favored and less and less frequently used.

In contrast to traditional surrogacy, a gestational surrogate is a woman who carries the child to term but who has no genetic tie to the child because the intending mother’s (or a separate donor’s) egg had been used. A gestational surrogate is less likely to be able to claim successfully that she is the child’s legal mother.

New Hampshire was one of the first states to have comprehensive legislation on the subject of surrogacy. It recently jettisoned this early effort and passed a new law in 2014 (NH-ST, Chapter 168-B). It demands that all parties have an agreement that must be submitted to and approved by the court before any medical reproductive procedures begin.

**Can surrogates be paid for their services?**

In most states surrogates can be reimbursed for medical and related expenses, needed supplies related to the pregnancy, and loss of wages if applicable. If payment is excessive, the arrangement may run afoul of prohibitions against the selling of children. One should proceed with caution in this area only after checking the laws of any states involved. This is a controversial issue; in some states and countries, only altruistic (no payment) surrogacy is permitted. Many question why women who perform this services should not be justly compensated, reasoning that the medical personnel, lawyers, and other associated with the procedure are paid and therefore the surrogate should be paid as well. Finally, surrogacy often involves an intending couple seeking a surrogate in a foreign country, where surrogates can be paid but often relatively little. Some countries have encouraged this kind of business; some—like India—after much experience with so-called “baby farms,” i.e., large communal living arrangements for would-be and pregnant surrogates, have now banned the practice for foreign potential parents.

**What is in vitro fertilization and what is the resulting legal status of each of the parties?**

In vitro fertilization is simply an outside-the-womb method of conception. Sperm and egg are placed together in culture and fertilization is allowed to occur. A special kind of in-vitro, known as ICSI, can be used in which a medical professional injects a single sperm into an egg for the
purpose of fertilization. The fertilized egg is then planted into the uterus of the mother. In either case, the egg and sperm could be those of the intending couple or one or both of the gametes could be donated. Regardless of the source of the gametes, the couple who caused the process to occur — often referred to as “the intending parents” — are most often deemed to be the legal mother and father.

What are cryopreserved (frozen) embryos and what legal issues does this process raise?
The in-vitro fertilization process results in excess embryos, i.e., more embryos than can be used in a single cycle for conception. These excess embryos are frozen for further use. Embryos can remain frozen for many years and perhaps indefinitely. Frozen embryos are stored in clinics and hospitals or in off-site storage centers (for an annual fee) for use in additional fertility cycles, either because the first cycles were not successful, or to accomplish subsequent pregnancies. Frozen embryos can also be donated to another couple or woman for implantation.

Are embryos available for research?
Yes. The individuals to whom the embryos belong may donate them for research. These frozen embryos are in fact a prime source for embryonic stem cell research. Whether or not federal funding should be used in this kind of research has been controversial and has changed over time according to presidential executive orders under the Bush and Obama administrations; the United States Supreme Court in Sherley v. Sibelius (2013) ended attempts to block federal funding.

What happens to the embryos if a couple divorces?
Prior to beginning fertility treatment, couples ought to enter into contracts that deal with what should happen to embryos in the event of the couple’s divorce. Despite these contractual provisions, one or both of the parties may change their minds during the divorce. One party may wish to have the embryos for the purpose of procreation and the other seeks to avoid becoming a parent under these circumstances. There are a handful of reported cases in which divorce courts have had to decide what will happen with these embryos. At least one state has decided to uphold the parties’ agreement. The other states have decided that contracts such as these are against public policy, and have sided with the party who does not want to procreate. Some states are requiring contemporaneous consent; each party must consent anew to each new procedure, regardless of what a prior contract had provided.

What is posthumous reproduction and what legal issues does it raise?
Posthumous reproduction occurs when the gamete of one party is used after his or her death to procreate. This can happen in a variety of ways. While still alive and capable of consent, parties can choose to have their gametes frozen and stored for a later use. Sperm can be frozen easily and for lengthy periods of time; freezing eggs is a much newer phenomenon and is more difficult to do successfully. This procedure is done most often by persons who may be about to undergo chemotherapy or other medical treatment that could cause sterility, and to a lesser extent, by some armed services members prior to their leaving to go to war. Should the individual then die, the gametes are available, usually to a spouse or partner, for use in ART procedures.

Another form of posthumous reproduction is more controversial, and involves the harvesting of sperm from an already deceased man, usually at the request of a partner or spouse, for the purpose of using the sperm to procreate. Some hospitals will perform this procedure, which must
be done within a short time after death; medical literature suggests 72 hours. Many hospitals refuse to do this, citing the inability of the deceased to consent to the procedure and to the subsequent reproduction. There have been some cases in which someone other than a spouse, e.g., parents of the deceased, have made the request.

Posthumous reproduction involving the removal of eggs from a deceased woman is uncommonly rare since harvesting immature eggs would be medically useless. However, advances in medical capabilities may allow the harvesting of ovarian tissue, which under certain medical circumstances, could produce mature eggs for use in reproductive procedures.

Children who are born as a result of these forms of posthumous reproduction must sometimes fight to be recognized as the legitimate offspring of the deceased for purposes of government benefits such as Social Security, life insurance, and estate purposes.

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RISK MANAGEMENT

Topics Covered in this Chapter:

What is risk management?
Risk Management Guidance
- Medical Records: Documentation of Patient Care
- Emergency Care Preparation in Your Office Practice: Adult Care
- Emergency Care Preparation in Your Office Practice: Pediatric Care
- Biopsy Specimen Send Outs
- Closing Your Practice
- Communication with Consulting Physicians
- Diagnostic Test Tracking Systems
- Informed Consent
- Noncompliant/Non-adherent Patient Management
- Termination of the Physician-Patient Relationship

About the Authors

By: Cheryl Peaslee, Nancy Brandow, Stacey Doten, Lou Anne McLeod
Medical Mutual Insurance Company of Maine Risk Management

WHAT IS RISK MANAGEMENT?

In general terms, risk management (RM) is the process of identifying and mitigating risk. Activities can be proactive (prevent the risk of an adverse event) or reactive (minimize the loss or damage after an adverse event). The risk management process includes five steps:

1. **Risk Identification**: What are the risks? What is the root cause?
2. **Risk Assessment**: What is the likelihood of the risk occurring? What is the impact or consequence should an event occur?
3. **Risk Plan**: What can be done to lessen the impact and reduce the likelihood?
   
   Consideration is given to:
   - transference (pass the risk to someone else, e.g., insurance)
   - avoidance (make major changes to eliminate the risk)
   - mitigation (take action to reduce the negative impact of the risk)
   - retention (retain the risk; take no further action)
4. **Risk Control and Plan Implementation**: Who is responsible for the plan? How will it be implemented? How will it be communicated?
5. **Risk Monitoring**: Is the plan effective? Did the action have the intended impact?

In healthcare, clinical risk management is an approach to improving the quality and safe delivery of care by identifying circumstances that put patients at risk of harm and implementing solutions to prevent the risk from occurring. In addition, when adverse events occur, a root cause analysis is often performed and change implemented to prevent reoccurrence. Establishing an organizational culture of safety results in the delivery of safer patient care and minimizes the financial risk of an adverse event.
Enterprise risk management (ERM) is a broad-based interdisciplinary process through which an organization identifies, analyzes, prioritizes, and addresses risks and opportunities (including value) that affect its achievement of strategic objectives. ERM was first implemented in the financial sector and has spread through all industries, slowly making its way into healthcare organizations. ERM encompasses clinical risk management and expands the program to encompass all aspects of the organization. Categories of risk are often referred to as domains. For example, the American Society for Healthcare Risk Management notes eight domains: Operational, Financial, Human Capital, Strategic, Legal/Regulatory, Technology, Hazard and Clinical/Patient Safety. In evaluating a potential or actual risk, the impact on all domains is assessed.

**RISK MANAGEMENT GUIDANCE**

Healthcare providers address risk everyday as they carry out their mission to provide quality and safe patient care. At times consulting with an expert on determining the best approach or a solution to a risk situation may be in order. For example, assistance may be needed in addressing a patient complaint, documenting refusal of care, or developing an approach to a non-compliant patient. When a risk concern arises or if assistance is needed in conducting a proactive risk assessment, contact your medical professional liability insurance company (MPLI). Most MPLI carriers offer risk management consultative support and can help identify and offer solutions to enhance patient care and mitigate risk.

Medical Mutual Insurance Company of Maine provides direct risk management support to its insureds. The following guidance addresses frequent topics addressed by MMIC risk management consultants. To access Medical Mutual Insurance Company of Maine’s complete online library of Practice Tips go to: [https://www.medicalmutual.com/risk/practice-tips](https://www.medicalmutual.com/risk/practice-tips).

*Medical Records: Documentation of Patient Care in the Legal Health Record*

The legal health record is generally the information used by the patient care team to make decisions about the treatment of a patient. The elements that constitute an organization's legal health record vary depending on how the organization defines it but must explicitly identify the sources, medium, and location of the individually identifiable data that it includes.

The importance of excellent patient care documentation in the legal health record cannot be understated as this record:
- Supports the decisions made in a patient's care.
- Supports the revenue sought from third-party payers.
- Documents the services provided as legal testimony regarding the patient's illness or injury, response to treatment, and caregiver decisions.
- Serves as the organization's business and legal record, meeting all statutory, regulatory and professional requirements for clinical and business purposes.
- Serves as a communication tool and is critical to patient safety and continuity of care.
- Is typically used when responding to formal requests for information for evidentiary purposes.
- Provides a witness to the care provided in the case of medical malpractice litigation.
The Guiding Principles for Documentation of Patient Care can be found at:

Emergency Care Preparation in Your Office Practice: Part I – Adult Care
Many providers are misinformed that the availability of emergency equipment and medication in the practice increases liability exposure. In actuality, failure to plan and lack of adequate preparation to provide emergency care may lead to increased liability. To determine how to establish proper procedures in your practice, please access:

Emergency Care Preparation in Your Office Practice: Part II – Pediatric Emergency Preparedness
Family practice and pediatric physicians experience emergencies among their patient populations more often than some physicians recognize. Status asthmaticus, upper airway obstructions, trauma and shock are reported as the most common events, as are severe respiratory distress, high fevers and seizures. Physicians need to ensure appropriate management and stabilization of emergency events encountered in their office. Perceived rarity of office emergencies should not be a reason for a lack of preparedness.

False complacency will find you unprepared to manage common, serious incidents. Parents expect that necessary equipment is available at the practice to respond to an emergency event involving their child.

Many providers are misinformed that the availability of emergency equipment and medication in the practice increases liability exposure. In actuality, failure to plan and lack of adequate preparation to provide emergency care may lead to increased liability. To determine how to establish proper procedures in your practice, please access further information at:

Biopsy Specimen Send Outs: Risk Management Recommendations to Avoid Delays in Diagnosis
Delay in cancer diagnosis is one of the leading causes for professional liability claims. Mislabeling of specimens, failure to track receipt of the results and follow-up on the results are contributing factors in these claims. A seven step process is provided at:

Closing Your Practice: Retirement - Relocation - Selling your practice
Circumstances may lead a physician to end his/her current practice arrangement. Providing notice in a timely manner promotes continuity of patient care, avoids allegations of abandonment, and fulfills contractual and regulatory obligations. When feasible, begin planning your departure years in advance. Key areas to be addressed can be found at:
Communication between the Referring and Consulting Physician

Issues arise when clear expectations are not defined between the referring and consulting physician. Poor coordination of care can be the result of undefined roles leaving the patient at risk and the physician's exposure to liability heightened. The referring physician should clearly define the expectations of the consultation to both the patient and consulting physician. For example:

- Referral is for consultation only. The consultant's assessment and recommendations will be forwarded to the referring physician for evaluation and consideration.
- Referral is a transfer of care for a specified condition. The consultant is to assess the patient and implement the recommended treatment.

To learn more about the closed loop referral process go to: https://www.medicalmutual.com/risk/practice-tips/tip/communication-between-the-referring-and-consulting-physician/45.

Diagnostic Test Tracking Systems

Failure to diagnose is one of the most frequent allegations in malpractice claims. A direct relationship exists between this allegation and the lack of systems or breakdown in systems that support:

- Tracking of results of ordered labs, diagnostic tests and consultations.
- Follow-up with the patient if results are not received.
- Physician review of results and development of the treatment plan.
- Notifying the patient of the results and communication of the treatment plan.
- Determining the patient's understanding of results and the treatment plan.
- Tracking patient completion of the treatment plan.

A patient-focused practice culture that includes effective communication and operational systems facilitates the delivery of safe patient care. A comprehensive, reliable test/consult tracking system must be established, adhered to, and monitored routinely for system failures.

Test/Consult Tracking System Development is key to assuring safe patient care. Find out the steps in the process at: https://www.medicalmutual.com/risk/practice-tips/tip/diagnostic-test-tracking-systems/53.

Informed Consent: A Process for Building Patient Confidence

Obtaining informed consent is the provider's legal responsibility. Granting informed consent is the patient's exclusive right. According to the National Quality Foundation (NQF): "Although providers often believe that a patient’s signature on a form is sufficient to ensure that he or she provided informed consent, studies show that even after agreeing to or receiving care, 18 to 45 percent of patients are unable to recall the major risks of surgery, many cannot answer basic questions about the services or procedures they agreed to receive, 44 percent do not know the exact nature of their operation, and most do not read (60 to 69 percent) or understand (60 percent) the information contained in informed consent forms, despite signing them. These patients are not truly informed about the choices they make." A fully comprehended informed consent improves patient safety and quality of care and reduces professional liability risk for providers and hospitals.
Noncompliant/Non-adherent Patient Management

Physicians are challenged and frustrated by patients who fail to adhere to and to follow a treatment regimen knowing that serious consequences may result in permanent injury and death. The reasons patients deviate from their care plan of keeping follow-up appointments, participating in ordered tests, and taking medications as prescribed may include:

- Denial, depression, dependence, and dementia.
- Lack of agreement or understanding with the physician's assessment, recommendation, or instruction.
- Lack an understanding of the seriousness of the illness/condition.
- Language, cultural, or mental barrier which hinder a patient's ability to understand or follow the care plan.
- Receiving conflicting advice from multiple caregivers with resulting confusion.
- Cost of treatment, including lack of transportation, interfering with patient's ability to comply with the care plan.

Noncompliance is often a communication breakdown or misunderstanding. Physicians need to appreciate the patient's understanding of their problems. Facilitating a face-to-face conversation with the patient allows the provider to develop effective strategies to manage and deal with the "noncompliance." Physicians must give clear explanations of health problems and recommended solutions to establish mutual cooperation.

The following tip offers steps to follow when addressing noncompliance:

Termination of the Physician/Patient Relationship

A physician's improper termination of the physician-patient relationship may put the physician at risk for a claim of abandonment. Consideration should be given to the following:

- Evaluate whether all options have been exercised to salvage the relationship. Don't act hastily in making a decision.
  - For "patient noncompliance", facilitate a face-to-face conversation with the patient to clearly communicate expectations. Allow the patient to voice their understanding and expectations. Clarify any misunderstandings or misperceptions. Facilitate a mutual agreement to a plan. Provide the patient with a copy of the written agreement.
- Review the documentation in the patient record to determine if the documentation supports the decision to terminate the relationship.
- Review managed care contracts to determine if the relationship with the patient can be terminated.
• If the patient is in a protected class or disabled, consult an attorney to determine if the termination is prohibited.


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WORKERS’ COMPENSATION

Topics Covered in this Chapter:
Navigating Workers’ Compensation in Vermont
The Workers’ Compensation Act; General Considerations for Medical Providers
Workers’ Compensation Issues When Treating a Patient Injured at Work
Special Rules Relating to Opioid Medication
Medical Marijuana
Resources
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NAVIGATING WORKERS’ COMPENSATION IN VERMONT

Vermont first adopted a no-fault workers’ compensation insurance program in 1915, with the passage of Act No. 164, “An act relating to compensation to employees for personal injury.” At that time, quarries, factories, and other physical labor comprised the majority of the state’s workforce and the legislation was viewed as a “grand compromise between labor and management.” The new law provided injured workers with access to prompt medical attention and wage replacement benefits for work-related injuries and, in exchange, granted employers immunity from personal injury civil lawsuits stemming from those workplace injuries. More than 100 years later, the core principles of the Workers’ Compensation Act remain the same.

Once an employee is injured or claims they were injured at work, whether by a discrete incident or a gradual onset type injury, and the employer receives notice of the injury, the employer or insurance carrier must notify the Department of Labor’s Workers’ Compensation Division of the injury within 72 hours. This sets in motion a series of deadlines for the employer and the employer’s insurance carrier, and a set of rules, which must be met in the adjusting of the claim. Many of these deadlines and rules may intersect with the function of medical providers who find themselves treating an injured worker.

For the casual practitioner in the realm of workers’ compensation, it can be a complicated system to navigate. This Chapter seeks to give medical providers treating patients with work injuries a basic understanding of the Act and important rules, case law, and deadlines that apply to treating providers.

THE WORKERS’ COMPENSATION ACT: GENERAL CONSIDERATIONS FOR MEDICAL PROVIDERS

The first step in any Vermont workers’ compensation matter is the filing of a First Report of Injury (Form 1) by the employer or insurance carrier/third-party administration (collectively hereinafter “carrier”). This form, which provides basic information about the injury, employee, and employer, is filed even if the employer or carrier disputes the facts of the injury or the relationship to the injured worker’s work.
The employer and carrier can then either accept the claim of injury or exercise their right to a 21-day investigatory period to determine whether they will accept or deny the claim. This determination is typically made on the basis of the injured worker’s statement, employer’s and/or carrier’s investigation of the alleged injury and the workplace in which it occurred, and medical evidence. Notably, when an injury occurs, the employer/carrier is entitled to all medical records relating not just to the injury, but also all prior medical records relating to the same body part or condition and will contact medical providers to obtain these records. 21 V.S.A.§ 655a (b). Note that Title 21 VSA §655a requires providers to utilize and comply with a Vermont Worker’s Compensation Medical Authorization (Form 7) when seeking or providing medical information relative to a workers’ compensation claim. Workers Compensation claims are expressly exempted from the terms and provisions of the Health Insurance Portability and Accountability Act of 1996 (HIPAA), 45 CFR 164.512(1). A copy of the Medical Authorization can be found here. See also Rule 3.2100.

If the claim is denied the carrier will not pay either indemnity benefits or medical expenses for the injury, but the injured employee can appeal this decision with the Workers’ Compensation Division, which will adjudicate the dispute. If the Department finds the injury compensable following such an appeal, it will order the carrier to retroactively begin paying benefits to the injured worker.

The Vermont Department of Labor also allows payment of a workers’ compensation claim on a “without prejudice” basis. Rule 3.2300. However, the employer or carrier must notify the Department of Labor and the injured worker of its election to pay any portion of the claim, including specific medical treatment, on a without prejudice basis. If an employer or carrier elects to pay a claim, or any portion thereof, on a without prejudice basis, the employer or carrier must deny the claim within 90 days of making a without prejudice payment. If the claim is not denied within 90 days of making a without prejudice payment, the claim will be deemed accepted by the employer or carrier/administrator.

When presented with a new claim of injury, the employer or carrier has the right to designate a health care facility and/or provider to initially treat an injured worker for a claimed work-related injury. 21 V.S.A. §640(b). At any time after the initial treatment, the injured worker can transfer their care to another health care provider by filing a Notice of Intent to Change Health Care Provider (Form 8), which includes the reason the worker is seeking to change providers.

Medical providers make some of the most important determinations in workers’ compensation claims, including whether an injured worker has a work capacity. Injured workers tend to recover faster from work injuries if they remain engaged in the workplace and feel that the employer is invested in having them return to the workplace. If an injured worker has a partial work capacity, they may return to work on a light or reduced duty basis. Most employers are willing to work to find or create a job that meets an injured worker’s physical and temporal limitations during the period of recovery in order to keep the employee engaged. Medical providers should keep this in mind and, rather than thinking strictly about the injured worker’s typical job assignment, should complete work capability forms (Form 20) that indicate whether
there are other physical activities the injured worker could still perform during the period of recovery.

Independent medical examinations figure heavily into workers’ compensation claims. While an injured worker is receiving benefits under the Workers’ Compensation Act, the employer/carrier has the right to schedule the injured worker for independent medical examinations at reasonable intervals and with due regard for the injured worker’s schedule and ability to travel. 21 V.S.A. § 655; Rule 6.0000. Independent medical examinations must be scheduled with a medical provider within a two-hour driving radius of the injured worker’s home. However, the Commissioner of the Department of Labor has the discretion to order an independent medical examination outside the two-hour driving radius if the injured worker consents and/or if the injured worker’s condition warrants the specialized expertise of a more remotely located provider. In practice, the Commissioner rarely grants permission to send an injured worker to a provider more than two hours away unless the injured worker consents. An employer/carrier must give the injured employee and their attorney at least seven days’ notice prior to the date of the scheduled examination. If an injured worker does not plan to attend the independent medical examination, they must notify the employer or carrier of their plan not to attend at least three days prior to the scheduled examination. The injured worker also has a right to make a video or audio recording of an independent medical examination, as long as they provide notice to the carrier at least three days prior to the examination. The carrier will in turn notify the examiner that the injured worker is exercising this right. The examiner can refuse to conduct the examination due to video recording. The examiner cannot create their own video recording of the examination except the injured worker’s prior written consent, but can create an audio recording as long as the examiner notifies the injured worker before the examination that they will be doing so.

WORKERS’ COMPENSATION ISSUES WHEN TREATING A PATIENT INJURED AT WORK

Types of Injuries
There is a basic definition in the Vermont Workers’ Compensation and Occupational Disease Rules (hereinafter “the Rules”) of what is considered by the Department to be an injury. Under Rule 2.2600, “injury” means any harmful work-related change in the body, whether occurring instantaneously or gradually, and includes a claimed or apparent injury or disease. The term also includes damage to and the cost of replacement of prosthetic devices, hearing aids and eyeglasses when the damage or need for replacement arises out of and in the course of employment. 21 V.S.A. §601(7). Depending on the circumstances, the term “injury” also includes “aggravation,” “flare-up” or “recurrence” as those terms are defined in Rules 2.1200, 2.2300 and 2.3900.

The Rules discuss four basic types of injuries in detail, although not every injury will fall into one of these special groupings. The four groupings are first aid only injuries, recurrences, aggravations, and flare ups. Whether an injury is an aggravation, recurrence, or flare-up is a commonly litigated issue. The outcome of an aggravation/recurrence/flare-up analysis, in a case involving multiple employers, will determine who will be held responsible for benefit payments. In cases of causation not involving multiple employers it will determine whether there is a responsible employer or whether there is a non-work related cause.
The first type of injury is the “first aid only” injury, defined in Rule 2.2200 as “an injury for which the injured worker loses no time from work (except for the time, not exceeding one day of work, related to medical treatment and recovery), and which requires only one treatment that generates a bill for less than $750.00.” See also 21 V.S.A. §640(e). Even though these injuries require limited medical treatment at the time, they can evolve into much greater injuries over time and therefore employers are required to report these injuries fully by completion of a First Report of Injury.

The determination of whether an injury is an aggravation vs. recurrence vs. flare-up is one of the more complex and heavily litigated analyses in workers’ compensation claims due to the fact that it is often determinative of which among several employers or carriers is liable for paying the claim. A recurrence is defined in Rule 2.3900 as “the return of symptoms following a temporary remission.” An aggravation is defined in Rule 2.1200 as “an acceleration of exacerbation of a pre-existing condition caused by some intervening event or events.” A flare-up is defined by Rule 2.2300 as “a temporary worsening of a pre-existing condition caused by a new injury for which a new employer or insurance carrier is responsible, but only until the condition returns to baseline and not thereafter.”

The Department will consider five factors in order to determine under which category an injury falls: whether a subsequent incident or work condition has destabilized a previously stable condition; whether the injured worker had stopped treating medically the earlier condition; whether the injured worker had successfully returned to work for the earlier condition; whether the injured worker had reached medical end result for the earlier condition; and whether the subsequent work contributed independently to the final disability. Rule 2.3910. If all of these factors are met, then the Department will most likely consider the injury to be an aggravation.

**Reasonable Medical Care**

Another aspect of workers’ compensation claims that is frequently litigated is whether a particular treatment is appropriate or compensable. Treatment that an injured worker receives for a work injury will only be a compensable part of the workers’ compensation claim if it is “reasonable medical treatment.” Reasonable medical treatment is defined under Rule 2.3800 as “treatment that is both medically necessary and offered for a condition that is causally related to a compensable work injury.” A determination of whether a treatment is reasonable will be based primarily on evidence establishing “the likelihood that it will improve the patient’s condition, either by relieving symptoms and/or by maintaining or increasing functional abilities.” See also 21 V.S.A. §601(27), defining “medically necessary care.”

**Reimbursement for Services**

In accordance with 21 V.S.A. §640(a), the employer/carrier is responsible for payment of reasonable and necessary medical treatment for a work-related injury. Billing for treatment of a work-related injury must be directed to the employer/carrier, not the worker, generally within 6 months after the date the health care provider had actual knowledge that the services were related to a workers’ compensation claim. Rule 40.000 establishes the Workers’ Compensation Fee Schedule. It states that the liability of an employer to pay for medical, surgical, hospital and nursing services provided to an injured employee shall not exceed the maximum fee for a particular service as provided for in the rule. Generally speaking, the maximum allowable
payment is the fee for a procedure listed in Appendix I of the rule or the health care facility's or health care provider's charge, whichever is less. For those procedures having no code listed in Appendix I of the fee schedule, the maximum allowable payment shall not exceed 83% of the charge for the service. 21 V.S.A. §640a and Rule 40.000 also contain additional information regarding types of providers that can be reimbursed, documentation required with claims and other billing and timely filing requirements. 21 VSA §§ 640a (a)-(c) outline the timeline and process by which an employer/carrier must pay or deny the bill.

Preauthorization Request Guidelines

Often when there is doubt about whether a carrier will cover a particular treatment, a medical provider or the injured worker’s attorney, if represented, will complete a preauthorization request prior to moving forward with treatment. See 21 V.S.A. §640b; Rule 7.0000. This is optional and not a requirement under the Workers’ Compensation Act, as the statute states that a provider or injured worker “may submit a request to an employer or insurance carrier that a proposed medical treatment or diagnostic procedure to be preauthorized.” 21 V.S.A. §640b. The request must be in writing and must be accompanied by documentation supporting both the medical necessity of the proposed treatment or procedure and its causal relationship to the injured worker’s compensable injury or condition. This documentation should include relevant medical records. The request also must clearly delineate the extent of any treatment or diagnostic procedure proposed, in terms of amount, duration, and frequency.

When a preauthorization request is received, the employer or carrier will have 14 days from the receipt of both the request for preauthorization and the supporting medical documentation within which to respond. 21 V.S.A. §640b(a). The employer/carrier can respond in one of three ways: 1) By authorizing the proposed treatment or diagnostic procedure. Once authorized, the employer or insurance carrier shall be obligated to pay all appropriately billed charges related to the proposed treatment or diagnostic procedure in accordance with Rule 40.000. 2) By denying the proposed treatment or diagnostic procedure on one or more of the following grounds: (a) that the preauthorization request was not accompanied by the required supporting documentation; (b) that compensability of the injury or condition for which the treatment or diagnostic procedure is sought is disputed, on either legal or factual grounds (this defense shall not be available to an employer or carrier against whom an interim order to pay benefits has been issued); or (c) that the proposed treatment or diagnostic procedure is not medically necessary and/or is not causally related to the injured worker’s compensable injury or condition. 3) By ordering a medical record review and/or scheduling an independent medical examination for the purpose of determining whether the proposed treatment or diagnostic procedure is medically necessary and causally related to the injured worker’s compensable injury or condition. 21 V.S.A. §640b(a).

When an employer/carrier responds to a preauthorization request by ordering a medical records review and/or scheduling an independent medical examination, it has 45 days following the receipt of the preauthorization and supporting medical documentation in which to either approve or deny the preauthorization request. This means that the employer/carrier must receive the records review or IME report within that 45-day window in order to make their determination. In limited instances, the employer/carrier may be able to receive an extension of up to 10 days from the Commissioner, but only upon a showing of extremely unusual and/or emergency circumstances. 21 V.S.A. § 640b (a)(3). Alternatively, if both parties agree in writing an
extension of ten or more days may be granted for any reason, provided the time period within which to respond is clearly and specifically stated.

Should the employer/carrier fail to respond to the preauthorization request either within the initial 14-day window or within the 45-day period of records reviews or IMEs, either the injured worker or the treating medical provider may request that the Commissioner issue an interim order authorizing the treatment or diagnostic procedure by operation of law. 21 V.S.A. § 640b(b).

There are several matters that do not qualify for preauthorization requests: 1) A demand that the charges for a treatment or diagnostic procedure already undertaken, including prescription medications already purchased, be paid; 2) A request that the charges for medical supplies, including special clothing, footwear or equipment, but excluding prescription medications proposed as a course of treatment, be paid or reimbursement.

**Medical End Result**

Much in workers’ compensation claims turns upon the injured worker reaching “medical end result,” “end medical result,” or “maximum medical improvement” - terms which are often used interchangeably. End medical result is defined as “the point at which a person has reached a substantial plateau in the medical recovery process, such that significant further improvement is not expected, regardless of treatment.” Rule 2.2000. A finding of medical end result triggers other activity in a workers’ compensation claim, including a permanent impairment rating, a discontinuance of temporary indemnity benefits, and sometimes a return to work or vocational rehabilitation, if a work capacity/work release was not already provided during the recovery period. Medical end result will also often lead to a discontinuance of medical treatment (or at least the carrier’s liability for medical treatment) other than ongoing palliative care. Under Rule 2.3400, palliative care is defined as “medical services rendered to reduce or moderate temporarily the intensity of an otherwise stable medical condition” but not “medical services rendered to diagnose, heal or permanently alleviate or eliminate a medical condition."

**Medical Case Management**

Carriers are allowed to assign medical case managers under Rule 2.2900. Medical case management is defined as “the planning and coordination of health care services appropriate to achieve the goal of medical rehabilitation. Medical case management may include medical case assessment, including personal interview with the injured worker, assistance in developing, implementing and coordinating a medical care plan with health care providers in consultation with the injured worker and his or her family, and evaluation of treatment results.”

**Permanent Partial Impairment – How It Is Calculated**

Following a finding of medical end result, the treating provider may be asked to rate the injured worker’s permanent impairment. If the treating provider does not perform ratings, the injured worker and carrier will discuss finding an alternative medical provider to examine the injured worker and perform a rating. For non-psychological injuries, the Workers’ Compensation Act requires the physician to use the American Medical Association Guides to the Evaluation of Permanent Impairment, 5th edition to rate any permanent partial impairment. 21 V.S.A. 648(b), Rule 10.1300. For psychological injuries, the physician is required to use Chapter 14 of the

The impairment rating is then used by the carrier to calculate the amount of permanent impairment benefits due to the injured worker. Any non-spine rating is multiplied by 405 to determine the number of weeks of benefits to which an injured worker is entitled for that injury; this figure is then multiplied by the injured worker’s weekly compensation rate. The same formula is used for spine ratings, the only difference being that the multiplier used for spine ratings is 550 rather than 405.

**SPECIAL RULES RELATING TO OPIOID MEDICATION**

Many states, including Vermont, have responded to concerns regarding long-term prescribing of opioids for chronic pain and the risk of prescription drug abuse by putting forth rules that specifically address opioid prescribing. 21 V.S.A. § 640c. At the same time, medical marijuana has become a hot topic around the country. Vermont has legalized medical marijuana, but also explicitly stated that workers’ compensation carriers are not obligated to pay for medical marijuana, even if it found to be reasonable medical treatment. The pertinent statutory language addressing this exception for workers’ compensation carriers follows in the next section.

Vermont’s new workers’ compensation rules addressing the prescribing of opioids went into effect on November 1, 2016. These track the Vermont Department of Health’s Rule Governing the Prescribing of Opioids for Chronic Pain that was promulgated by the Department of Health in 2015. Failure of a medical provider treating an injured worker to comply with the Department of Health Rule creates a rebuttable presumption on behalf of the carrier, allowing the carrier to deny or discontinue payment for opioid medications prescribed to treat an injured worker’s chronic pain. These new workers’ compensation rules can be found at Rule 11.1400 and 12.1700, and follow in their entirety:

11.1400 - Denying payment for opioid medications. A medical provider who prescribes opioid medications to an injured worker for chronic pain resulting from a compensable work-related injury must comply in all respects with the Rule Governing the Prescribing of Opioids for Chronic Pain, as currently promulgated at 4A Code of Vermont Rules 13 140 076 (2015) and as amended from time to time by the Vermont Department of Health. If credible evidence establishes that he or she has failed to do so, a rebuttable presumption shall arise that the medications, as prescribed, do not constitute reasonable medical treatment. If the employer or insurance carrier seeks to deny payment on those grounds, it shall file a Denial of Workers’ Compensation Benefits (Form 2) with the Commissioner and the injured worker, and shall comply in all respects with the requirements of this Rule 11.0000. In addition, it shall notify the prescribing provider of the specific basis for its determination that he or she has failed to comply with the above-referenced Vermont Department of Health rule. Thereafter, the injured worker shall have the burden of proving that the treatment is reasonable, notwithstanding the prescribing provider’s failure to comply. In any event, the Commissioner shall not approve a proposed discontinuance under this Rule unless credible medical evidence establishes that the effective date thereof comports with a safe taper plan. 21 V.S.A. §640c.
12.1700 - Discontinuing medical benefits:

12.1720 - If the proposed discontinuance pertains to narcotic or other medications for which a safe taper plan is medically necessary, the employer or insurance carrier shall provide credible medical evidence establishing that the date of its proposed discontinuance comports with such a plan.

12.1730 - A medical provider who prescribes opioid medications to an injured worker for chronic pain resulting from a compensable work-related injury must comply in all respects with the Rule Governing the Prescribing of Opioids for Chronic Pain, as currently promulgated at 4A Code of Vermont Rules 13 140 076 (2015) and as amended from time to time by the Vermont Department of Health. If credible evidence establishes that he or she has failed to do so, a rebuttable presumption shall arise that the medications, as prescribed, do not constitute reasonable medical treatment. If the employer or insurance carrier proposes to discontinue payment on those grounds, it shall file an Employer’s Notice of Intention to Discontinue Payments (Form 27) with the Commissioner and the injured worker, and shall comply in all respects with the requirements of this Rule 12.0000. In addition, it shall notify the prescribing provider of the specific basis for its determination that he or she has failed to comply with the above-referenced Department of Health rule. Thereafter, the injured worker shall have the burden of proving that the treatment is reasonable, notwithstanding the prescribing provider’s failure to comply. In any event, the Commissioner shall not approve a proposed discontinuance under this Rule unless credible medical evidence establishes that the effective date thereof comports with a safe taper plan as required by Rule 12.1720. 21 V.S.A. §640c

The Department of Health’s Rule Governing the Prescribing of Opioids for Chronic Pain can be found on the rules section of the Vermont Department of Health’s website. Note that updated rules for chronic pain will go into effect July 1, 2017 and will be integrated with rules for acute pain into one Rule Governing the Prescribing of Opioids for Pain.

MEDICAL MARIJUANA

According to Vermont Statute Title18, Section 4474c, Marijuana For Medical Symptom Use By Persons With Severe Illness, a workers’ compensation carrier is not obligated to pay for medical marijuana even if it is considered to be reasonable medical treatment. The provision reads in relevant part:

(b) This chapter shall not be construed to require that coverage or reimbursement for the use of marijuana for symptom relief be provided by:

(1) a health insurer as defined by section 9402 of this title, or any insurance company regulated under Title 8;

(2) Medicaid or any other public health care assistance program;

(3) an employer; or
(4) for purposes of workers’ compensation, an employer as defined in 21 V.S.A. § 601(3).

There are arguments being advanced that this should not preclude an injured worker from receiving, or the workers’ compensation carrier from paying for, medical marijuana. However, at the time of this writing there have been no decisions from the Department addressing this issue.

**RESOURCES**

Workers’ Compensation [Rules](#)
Workers’ Compensation [Statute](#)
Workers’ Compensation [Medical Fee Schedule](#) (Rule 40)
Workers’ Compensation [Forms](#)

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**Anne Cramer**, (Consent, Privacy and Medical Records; Disclosure and Reporting Requirements) a shareholder in the law firm of Primmer, Piper, Eggleston & Cramer, P.C., serves as counsel to hospitals, nursing homes, community mental health agencies, physician groups and other private health care interests in Vermont. Anne and the Primmer firm have long
served as counsel to the Vermont Association of Hospitals and Health Systems and also provide counsel to the Vermont Health Care Association, and the Council for Developmental and Mental Health Services. She is a member of the American Health Lawyers Association, and the Health Law Section of the American Bar Association. In her health law practice, Ms. Cramer emphasizes compliance with federal and state regulatory requirements, including fraud and abuse prevention, the HIPAA privacy and security rules, antitrust compliance and employment law. Ms. Cramer lectures frequently on health law topics generally, and she has been listed in New England Super Lawyers and in the Best Lawyers in America for her health law related work for over twenty-five years.

Stacey Doten (Risk Management) is a Senior Risk Manager with Medical Mutual Insurance Company of Maine. She has Associate Degrees in both Laboratory Science from the University of Maine at Augusta and Nursing from Northern Maine Community College. She has more than 20 years of healthcare experience, including healthcare quality and risk management. She is a Certified Professional in Healthcare Quality (CPHQ) and a Certified Professional in Healthcare Accreditation (HACP). She is a member of the Northern New England Society for Healthcare Risk Management (NNESHRM), National Association of Healthcare Quality (NAHQ), and the Maine Medical Group Management Association (MEMGMA).

Eileen Elliott (Fraud and Abuse Compliance) is an attorney focusing on health care and human services legal issues with the Burlington law firm of Dunkiel Saunders Elliott Raubvogel & Hand, PLLC. She was the deputy secretary of Vermont’s Agency of Human Services from 2003-2005 and the commissioner of Vermont’s Social Welfare Department from 1999-2003. From 1993 until 1999, she worked for the Office of the Attorney General, serving as chief of its Human Services Division, and before that as counsel to the Agriculture Department. She spent the first decade of her professional career in private and corporate practice. Eileen graduated with distinction from the University of Colorado with a B.A. in Environmental Conservation and she has a J.D. from the University of Denver.

Erin J. Gilmore (Workers’ Compensation) is an attorney with Ryan Smith & Carbine, Ltd. She has represented many insurers on issues before the Department of Labor. Ms. Gilmore frequently provides training and guidance for insurance adjusters in handling workers' compensation claims in the State of Vermont. Ms. Gilmore graduated with a J.D. from the University of North Carolina, Chapel Hill and a B.A. from the University of Vermont.

Shireen Hart (Consent, Privacy and Medical Records; Disclosure and Reporting Requirements), a shareholder of Primmer, Piper, Eggleston & Cramer, P.C.’s Healthcare Group, draws upon her more than 15 years of experience in the field to provide pragmatic counsel to hospitals, long term care facilities, community mental health agencies, pharmacies, laboratories, individual practitioners and healthcare related vendors on a broad range of matters. Her strong statewide connections and effectiveness working with other attorneys and state agency or department personnel creates a favorable environment in which to deliver positive results for clients.

Drew Kervick (Fraud and Abuse Compliance) is a health care and business law attorney with Dunkiel Saunders Elliott Raubvogel & Hand, PLLC. Before joining the firm in 2014, Drew worked in private practice both in Burlington, Vermont and in Boston Massachusetts. He also
clerked for Justice John Dooley of the Vermont Supreme Court. Drew graduated summa cum laude from Boston College Law School, and holds a master’s degree from University of North Carolina at Chapel Hill and a bachelor’s degree from Stanford University, both in Political Science.

Lauren Layman (Consent, Privacy and Medical Records; Disclosure and Reporting Requirements) focuses her practice on the areas of health care law and legislative relations. She brings a uniquely effective perspective to her work through her prior experiences as policy advisor for the Centers of Disease Control in Washington, D.C. and a health care staff member for Senator Jim Jeffords. A Vermont native, Lauren is knowledgeable not only about the policies and legal climate affecting the state’s hospitals, but also the patients and communities they work with on a daily basis. This mix of professional and personal experience enables Lauren to handle cases with a multi-faceted, balanced approach.

Lou Anne McLeod (Risk Management) is a Senior Risk Manager at Medical Mutual Insurance Company of Maine. She has been involved in Quality Improvement and Risk Management since 1986. Lou Anne graduated from the University of New Hampshire with a BS in Health Administration and Planning and a Master of Health Administration. She is a Certified Professional in Healthcare Risk Management (CPHRM), a Fellow of the American Society for Healthcare Risk Management (FASHRM) and a Certified Professional in Healthcare Accreditation (HACP). Lou Anne is a member ASHRM, the New Hampshire Medical Group Management Association (NHMGMA) and the Massachusetts Society for Healthcare Risk Management (MASSHRM).

Glenn S. Morgan (Workers’ Compensation), a leading expert in the field of workers' compensation law, is Chairperson of the seven attorney Workers’ Compensation Practice Group at Ryan Smith & Carbine, Ltd in Rutland, Vt. Mr. Morgan has practiced in the area of workers' compensation for over 40 years, representing national, regional and local employers and insurance carriers before the Vermont Department of Labor, the Vermont Superior Court and the Vermont Supreme Court. Mr. Morgan is a Fellow of the College of Workers' Compensation Lawyers. Mr. Morgan has been recognized as a Vermont Top Lawyer by VT Business Magazine, Best Lawyers® in America, and “Super Lawyer” in the Workers' Compensation category and Top Attorneys in New England. He has received Martindale-Hubbell, Inc.’s highest "AV" rating. Mr. Morgan graduated with a J.D. degree from Suffolk University School of Law and a B.A. from the University of Vermont.

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**David Pocius** (Regulation of Physicians) is a partner at Paul Frank + Collins P.C. He represents clients in a variety of civil lawsuits, including commercial, environmental, premises liability, product liability, construction, and professional liability claims. He regularly defends physicians and institutions in medical malpractice cases, as well as physicians before the Board of Medical Practice, to which he brings his experience as a biologist and researcher prior to attending law school. David is an experienced litigator with trial experience stretching from state and federal courts in Vermont and New York to Colorado, the Northern District of Illinois, and the Southern District of Florida.

**Stephanie P. Romeo** (Workers’ Compensation) is an attorney with Ryan Smith & Carbine, Ltd. The majority of her practice consists of representing institutions and employers in defense of workers’ compensation matters. Ms. Romeo graduated with a J.D. from Vermont Law School and a B.A. from Northeastern University.

**Jon Rose** (Fraud and Abuse Compliance) is a regulatory and civil litigation attorney at Dunkiel Saunders Elliott Raubvogel & Hand, PLLC. Prior to joining Dunkiel Saunders, Jon served as an Assistant Attorney General in the Civil Division of the Vermont Attorney General’s Office, and before that, practiced at another prominent Burlington law firm advising a wide range of institutional and corporate clients on labor and employment, health care, and regulatory issues. Jon graduated first in his class from Vermont Law School, and holds a B.S. in Mechanical Engineering from Penn State University.

**Corina N. Schaffner-Fergard** (Workers’ Compensation) is an attorney with Ryan Smith & Carbine, Ltd. representing employers and their insurance carriers with respect to workers’ compensation, liability, and environmental claims. In her twenty years of practice, she has successfully tried numerous cases before the Department of Labor. Ms. Schaffner-Fergard frequently provides training and guidance to insurance adjusters in handling workers’ compensation claims in the State of Vermont. Ms. Schaffner-Fergard graduated with a J.D. from Vermont Law School and a B.A. from Cornell University.

**Jason Turner** (Pharmaceutical Issues – Drug Diversion Section) is an Assistant Attorney General in Vermont’s Office of the Attorney General.

**Lindsey Wells**, B.S. (Therapeutic Use of Marijuana), received her Bachelor of Science Degree in Business Administration in 2005 from Norwich University. She has an extensive work history with state and federal regulations. She has been employed by the Department of Public Safety as the Marijuana Program Administrator for the Vermont Marijuana Registry since 2012. Her duties include overseeing and performing assessments of Vermont’s four registered dispensaries, overseeing the Registry’s applicant process, participating in the legislative process, and
amending the rules governing the program to facilitate the implementation of various statutory changes.